There are many "100 things to do before you die" lists on the web. Implementing Union-Find should be one of those things in a computer scientist's list.

3

2

Mr. Skeptic: "I believe what I see".

Mrs. Cynic: "He doesn't see what he doesn't believe"

6

15

17

19

21

23

24

26

- On a logical equivalence that ain't

Lecture XIII DISJOINT SETS

10 ¶1. Let U be a set of items, and let \equiv be an equivalence relation on U. We want to make
11 two kinds of requests on this equivalence relation: First, given $x, y \in U$, we want to know if x12 and y are equivalent, is $x \equiv y$? Second we want to modify the equivalence relation by declaring
13 that, from now on, x and y will be equivalent. Of course, this new \equiv relationship must be
14 propagated to other pairs to preserve transitivity.

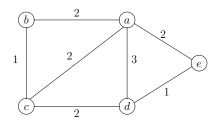
This equivalence maintenance problem is called the **union-find problem** because the two requests can be reduced the Find and Union operations. Other names for this problem include **disjoint sets** (as in our chapter title), **set union** or **set equivalence**. Underlying the best solutions to this problem is a data structure called compressed trees. Algorithms for this data structure are simple and easy to implement. Nevertheless, the ultimate complexity analysis of these algorithms is highly non-trivial, requiring a sophisticated form of amortization argument. It is remarkable that the analysis (Tarjan 1975) led to the inverse Ackermann function, an extremely slow growing function that is for all computational purposes is a small constant.

There are many practical applications of union find. We shall look at three: the implementation of Kruskal's algorithm for minimum spanning tree, a problem of reducing numerical expressions, and the problem of computing Betti numbers in Computational Topology.

§1. Union Find Problem

¶2. Implementing Kruskal's Algorithm. Consider this kind of queries on a given bigraph G = (V, E): given $u, v \in V$, are u and v in the same connected component? If G is static, this is easily solved by preprocessing the graph G. But in our problem, G is semi-dynamic in the sense that we can add edges to E. So we also need to process requests of the form: add the edge u-v to G.

Let us now use such a data structure to implement Kruskal's algorithm for the Minimum Spanning Tree ($\P V.39$). Let us simulate Kruskal's algorithm on the graph in Table . In



Edge	Weight	Accept?
b-c	1	✓
d-e	1	✓
a-b	2	✓
а-с	2	X
а-е	2	?
c-d	2	?
a-d	3	?

Table 1: Simulation of Kruskals Algorithm

Kruskal's algorithm, we first sort the edges in order of increasing cost, as shown next to the graph. We then process each edge by either accepting (\checkmark) or rejecting (x) it. The final MST comprises of the accepted edges. How do we make these decisions? We needed to decided for each edge u-v, whether u and v belong to the same component of the subgraph of accepted edges. This subgraph is dynamically changing as the algorithm progress.

Note that each edge represents an equivalence $relationship^1$ of an underlying equivalence relation. But different edge sets can represent the same relation. E.g., $\{1-2, 2-3\}$ and $\{1-2, 1-3\}$ represent the same equivalence relation.

Here is the abstract setup for our data structure: Let U be a finite set of items (called the "universe"). We may conveniently assume

$$U = \{1, 2, \dots, n\} = [1..n].$$

A partition of U is a collection

39

41

$$P = \{S_1, \dots, S_k\}$$

of pairwise disjoint non-empty sets such that $U = \bigcup_{i=1}^k S_i$. We say $x, y \in [1..n]$ are **equivalent**, written $x \equiv y$, if they belong to the same set in P. Each $S_i \in P$ is called an **equivalence class** or a **block** of the partition. For any $x \in U$, let set(x) denote the equivalence class of x. The items in U have no particular properties except that two items can be checked for equality. In particular, there is no ordering property on the items. To facilitate computation, we assume that each S_i has a unique **representative item** $x_i \in S_i$. Let $rep(S_i)$ denote this representative item x_i . The choice of representative item is arbitrary. Because of this representative item, we can now define the **Find** operation: Find(x) returns the representative item in the equivalence class of x. Hence

$$x \equiv y \iff \operatorname{rep}(x) = \operatorname{rep}(y) \iff \operatorname{set}(x) = \operatorname{set}(y).$$

The problem of checking if $x, y \in U$ are equivalent is now reduced to computing Find(x) and Find(y) and checking whether Find(x) = Find(y).

The partition P is dynamically changing because of the Union operation. If $x, y \in U$, then the **Union** operation declares that henceforth x and y are equivalent. This reduces to the problem of replacing set(x) and set(y) in P by their union $set(x) \cup set(y)$. Thus, the number of equivalence classes can only decrease, not increase. The **Union Find problem** is the problem of processing a sequence of Find/Union requests on a set U which is initially given the trivial equivalence relation (i.e., each equivalence class is a singleton $\{x\}$).

¹The terms "relation" and "relationship" are not interchangeable: by a (binary) **relation** R on a set V, we mean that $R \subseteq V \times V$. If $a, b \in V$, we call (a, b) a **relationship** of R if $(a, b) \in R$. Here, R happens to be an equivalence relation.

55

56

57

59

61

71

72

73

¶3. Example. Let n = 4 and U = [1..4]. Initially, the partition is $P = \{\{1\}, \{2\}, \{3\}, \{4\}\}$. Consider the sequence of requests:

Union(2,3), Find(3), Union(1,4), Union(2,1), Find(3).

After processing this sequence, $P = \{\{1, 2, 3, 4\}\}$. The two Find requests return (respectively) the representatives of the sets $\{2, 3\}$ and $\{1, 2, 3, 4\}$ that exist at the moment when the Finds occur. To be specific, if we choose the smallest integer in a set to be its representative then these Find requests return 2 and 1, respectively.

Each Union (resp., Find) requests can be viewed as an **equivalence assertion** (resp., **equivalence query**). The original motivation for this problem is from FORTRAN compilers where one needs to process the EQUIVALENCE statement in the FORTRAN language. This statement asserts the equivalence of two programming variables. Another application is in finding connected components in bigraphs. For more information, see Tarjan [6, 8] and a survey from Galil and Italiano [3]. Note that it is difficult to extend the union-find technique to allow the "undoing" of unions. One step in this direction is to have a persistent version of the union-find data structure where you can can go back to previous versions of the data structure. See Conchon and Filliâtre (2007).

¶4. Representation of Sets. In most solutions to the Union Find Problem, a set $S \subseteq U$ is concretely represented by a rooted unordered tree T whose node set is S. To represent T, each item x has a **parent pointer** x.p. The tree T is completely determined by these pointers. The root is the unique node $x \in S$ with the property x.p = x, and it serves as the **representative** of S. An alternative convention for indicating that x is a root is the condition x.p = nil. We call T a **compressed tree** representation of the set S. This data structure is from Galler and Fischer (1964). It is important to realize that there are no pointers from a node of T to any of its children, and the set of children are unordered (unlike, say, in binary search trees).

Where do we exploit the x.p = x convention below?

Using compressed trees, we can now represent any collection P of disjoint sets by a forest of compressed trees. For instance, let $P = \{0, \underline{5}, 6, 8, 9\}, \{\underline{3}\}, \{1, 2, 4, \underline{7}\}$. The underlined items (i.e., 5, 3 and 7) are representatives of the respective sets. A possible compressed tree representation of P is shown in Figure 1.

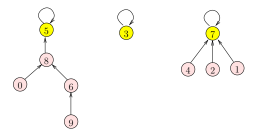


Figure 1: Forest of compressed trees representing $P = \{0, \underline{5}, 6, 8, 9\}, \{\underline{3}\}, \{1, 2, 4, \underline{7}\}.$

If x, y are roots of compressed trees, then we have the concrete or "physical" operation

as opposed to ADT operations

that links x to y. Basically, this amounts to the assignment $x.p \leftarrow y$. This amounts to making y the parent of x. We assume that x, y are roots of compressed trees. The result of linking x to y is a new compressed tree rooted at y, representing the union of the original two sets. If

91

92

97

98

100

101

102

103

104

105

106

107

108

109

x = y, then Link(x, y) is a null operation. Now we can physically implement a Union requests by two Finds and a Link:

$$Union(x,y) \equiv Link(Find(x), Find(y)).$$
 (1)

¶5. Complexity parameters. The complexity of processing a sequence of Union-Find requests will be given as a function of m and n, where n is the size of the universe U = [1..n] and m the number of requests (i.e., Union or Find operations). But the parameter m in the following discussions will be given a slightly different meaning: recall that the Union operation can be replaced by two finds and a single link as in (1). So we may replace a sequence of m Union/Find requests by an equivalent sequence of m Link/Find requests. Up to m0 order, the complexity of a sequence of m1 Link/Finds and the complexity of a sequence of m2 Union/Finds are equal. Hence, it suffices to analyze a sequence of m2 Link/Find operations. In some of our analysis, we use another parameter m1 which is the number of Find requests among the m2 operations; thus m2 m3 and m4 which is the number of Find requests among the m3 operations; thus m3 m4 m5 m5 m6 m8 and m9 operations; thus m4 m5 m9 m9 operations; thus m5 m9 operations; thus m6 m9 operations; thus m9 operations of m9 operations operations of m9 operations operations operations operations operations operatio

Although m and n are arbitrary parameters, for the purposes of lower bounds, we often assume that

$$m \ge n/2. \tag{2}$$

This inequality might be justified by insisting that every item in the universe must be referenced in either a Find or a Union request. But the "real" reason for making this assumption is that it allows us to show tight lower bounds.

¶6. Two Solutions to the Link/Find Problem.

Linked List Solution An obvious solution to the Link/Find problem is to represent each set in P by a singly linked list. We can regard a singly linked list as a special kind of compressed tree T which has only one leaf. The unique leaf of T corresponds to the **head** while the root is the **tail** of this linked list. Following links from any node will lead us to the tail of the list. This is like a traditional singly-linked list where the pointer x.p plays the role of the "next node" pointer. See Figure 2(a). Note that linking takes constant time, provided the head of each linked list stores a **tail pointer** that points to its own tail. The tail pointer is easy to maintain during a link operation. On the other hand, a Find operation requires a worst case time proportional to the length of the list. Clearly, the complexity of a sequence of m Link/Find requests is $\mathcal{O}(mn)$.

Traditionally, the tail of the linked list points to nil.

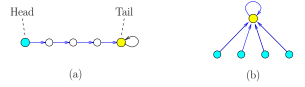


Figure 2: Two special kinds of compressed trees: (a) Linked list, (b) Anti-list.

Anti-list Solution Another solution is to represent each set in P as a compressed tree of height one: thus every non-root is a leaf that points to the root. See Figure 2(b). This data structure is the antithesis of lists: we will call such trees **anti-lists**. Clearly a

111

112

113

114

115

117

119

120

121

122

123

124

125

127

129

132

133

134

135

137

139

140

143

Find request will now take constant time. On the other hand, linking x to y takes time proportional to the size of set(x) since we need to make every element in set(x) point to y. Note that each Find has unit cost, and each Link costs O(n). Since there are at most n-1 links, we see that the cost of sequence of m operations costs

$$O(m+n^2). (3)$$

For a lower bound, we can make the *i*th operation link a set of size *i* to another set, costing *i* units. Therefore a sequence of min $\{m, n\}$ links would cost $(\min \{m, n\})^2$. Thus proves that the worst case complexity of processing m Link/Find requests is

$$\Omega(m + (\min\{m, n\})^2).$$

Assuming (2), this lower bound matches the upper bound in (3). REMARK: actually, anti-lists needs a little more structure among its nodes if we want to achieve the claimed upper bounds: assume that each node x has, besides the parent pointer x.p, also an extra pointer x.q. If x is the root of the anti-list, x.q points to one of its children. Moreover, all the children must be linked together like a singly-linked list by the q-pointers. Using both p- and q-pointers, we can carry out the Link operation in the stated time complexity.

¶7. Naive Compressed Tree Solution. The contrast between the list and anti-list representations is as sharp as can be: Linking is easy for lists but hard for anti-lists. Conversely, Find is easy using anti-lists, but hard for lists. Here, "easy" means $\Theta(1)$ and "hard" means $\Theta(n)$. Galler and Fischer (1964) shows that the relative advantages of both solutions can be exploited if we use the general compressed tree representation. But this advantage may not be immediately apparent. To see why, consider a straight forward implementation of the Find and Link operations under compressed tree representation. For Find(x), we just follow parent pointers from x until the root, which is returned. For Link(x,y), we simply set $x.p \leftarrow y$ and return y. These "naive" algorithms can lead to a degenerate tree that is a linear list of length n-1. Clearly $\Theta(mn)$ is the worst case bound for a sequence of m Link/Find Operations on n items. How can we do better?

¶8. Implementation. We briefly discuss a simple and effective way to implement all the union fine algorithms. Assume that there is an array U[1..n] that serves as the "register" for the items. Each U[i] is a reference to a node u representing item i. In our descriptions below, we often use u and i interchangeably (to no particular harm). Each node u has a member u.p that points to another node (possibly itself). Below, we need to associate a size or rank to each node. We can either have new node members such as u.size or we can have another array Size[1..n].

138 EXERCISES

Exercise 1.1: The root x of a compressed tree could also be signaled by the condition x.p =nil instead of x.p = x. What are the pros and cons of either convention? \Diamond

Exercise 1.2: Show the upper bounds for the above data structures are tight by demonstrating matching lower bounds (assume (2)).

- (a) When using compressed trees or linked lists, give an $\Omega(mn)$ lower bound.
- (b) When using anti-lists, give an $\Omega(m+n^2)$ lower bound.

147

148

149

150

151

152

153

154

155

156

157

158

159

160

161

162

163

164

Exercise 1.3: Implement Kruskal's Algorithm for Minimum Spanning Tree (Chapter V) using union-find data structure. However, we want to choose between (a) linked list and (b) anti-lists. Give their worst case complexity analysis. Which has the better complexity bound?

Exercise 1.4: Assume the anti-list representation, let us link two trees using the following heuristic: always append the smaller set to the larger set. To implement this heuristic, we can easily keep track of the sizes of anti-lists. Show that this scheme achieves $\mathcal{O}(m + n \log n)$ complexity. Prove a corresponding lower bound.

Exercise 1.5: We propose a Depth-2 Union-Find data structure: assume that the total number n of items in all the sets is known in advance. We represent each set (i.e., equivalence class) by a tree of depth exactly 2 where the set of leaves correspond bijectively to items of the set. All the leaves are at level 2; the internal nodes do not correspond to items, and are used only for book keeping. They are created at the beginning, but may be discarded as we form unions. Each node keeps track of its degree (= number of children) and maintains a parent pointer, a sibling pointer and a child pointer. The sibling pointers are linked into a circular list, and their common parent points to one child in this list. The following properties hold:

- (i) Each child of the root has degree between 1 and $2 \lg n$.
- (ii) A child of the root is **full** if it has degree at least $\lg n$. Each root has at most one **non-full** child.

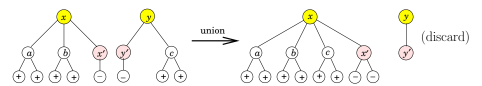


Figure 3: Illustration of Union.

Using this representation, a Find operation takes constant time. The union of two sets with roots x, y is done as follows: if $degree(x) \geq degree(y)$ then each full child of y becomes a child of x. Say x', y' are the children of x, y (respectively) which are non-full and assume that $degree(x') \geq degree(y')$. [We let the reader consider the cases where x' or y' does not exist, or when degree(x') < degree(y').] Then we make each child of y' into a child of x'. Note that since y and y' do not represent items in the sets, they can be discarded after the union.

- (a) Explain the rational for this design. Notice that the sibling and child pointers are alien to compressed trees why do we need them? This partly depends on your answer to parts (b,c).
- (b) Write pseudo-code to implement the operations of Find and Union. Also describe the initialization of the data structure from a given set of items. You must explicitly describe the assign/reassign parent/child/sibling pointers.
- (c) Prove that the complexity of this solution is

$$\mathcal{O}(m + n \log \log n)$$

where m is the total number of operations.

HINT: Devise a charging scheme so that the work in a Union operation is charged to nodes at depths 1 and 2, and the overall charges at depths 1 and 2 are (respectively) $\mathcal{O}(n)$ and $\mathcal{O}(n \log \log n)$.

165

166

167

Exercise 1.6: Extend the above Depth-2 Union-Find data structure to a Depth-3 data structure with amortized cost of $O(m + \lg \lg \lg n)$.

Exercise 1.7: Modify the above Depth-2 Union-Find data structure to the case where you do 171 not know n in advance. Specifically, you begin with n=0 and have explicit instructions 172 called New(x) to add a new singleton x, thereby incrementing n. The Find(x) and 173 Union(x,y) operations can only refer the previously introduced items. Give a complexity 174 analysis. 175

Exercise 1.8: Let (V, E) be a bigraph with weights assigned to each node. The weight of 176 a connected component of the graph is just the sum of all the weights of its nodes. 177 Suppose the edges are given "on-line" in the order e_1, e_2, \ldots, e_m . Let $G_i = (V, E_i)$ where 178 $E_i = \{e_1, \ldots, e_i\}$. After the appearance of e_i , but before we see e_{i+1} , we must output the 179 weight of the heaviest component in the graph G_i . 180

(a) Assume the weights are arbitrary (can be negative). Give an efficient implementation, and analyze its complexity.

(b) What improvements are possible for positive weights?

Exercise 1.9: Suppose we add to the Union-Find problem a third operation: break. Interpret the union operation as adding an edge to a bigraph that initially has no edges. The break operation is just the reverse of union, as it removes an edge (i.e., breaks a link). The Find(x) operation should again return a representative node of the connected component of the current bigraph that contains x. How efficiently can you solve this problem? 188

End Exercises

§2. Rank and Path Compaction Heuristics

There are several heuristics for improving the performance of the naive compressed tree algo-191 rithms. They fall under two classes, depending on whether they seek to improve the performance 192 of Links or of Finds. 193

¶9. Size and Rank heuristics. We now improve the overall performance of our data structure by restrictions on the way linking is done. For any node x, let size(x) be the number of items in the subtree rooted at x. Suppose we keep track of the size of each node, and in the union of x and y, we link the root of smaller size to the root of larger size. If the sizes are equal, the choice is arbitrary. This rule for linking is called the **size heuristic**. Thus, we modify the code for Union in (1) into the following:

> Union(x,y): $u \leftarrow Find(x); v \leftarrow Find(y)$ If (u.size < v.size)Link(u, v); $v.size \leftarrow v.size + u.size$ Else $Link(v, u); u.size \leftarrow u.size + v.size$

170

182

183

184

186

187

189

190

194

195

197

198

199 200

201

 \Diamond

§2. Two Heuristics Lecture XIII Page 8

Note that this heuristic assumes that each item has a member called *u.size*. It is easy to see that our compressed trees have depth at most $\lg n$ under this heuristic. Hence Find operations take $\mathcal{O}(\log n)$ time.

An improvement on the size heuristic was suggested by Tarjan and van Leeuwen: we keep track of a simpler number called rank(x) that can be viewed as a lower bound on $\lg(size(x))$ (see next lemma). The **rank** of x is initialized to 0 and subsequently, whenever we link x to y, we will modify the rank of y as follows:

$$rank(y) \leftarrow \max\{rank(y), rank(x) + 1\} \tag{4}$$

The **Rank Heuristic** is simply the Size Heuristic in which we use rank(x) instead of size(x).

This assignment (4) is the only way by which a rank changes. This change cannot decrease the rank of an item, and under the rank heuristic, it can increase by at most 1. Thus the rank of an item is non-decreasing over time. If compressed trees are never modified except through linking, then it is easy to see that rank(x) is simply the height of (the compressed tree rooted at) x. But in the presence of other heuristics to be specified, rank(x) is just an upper bound on the actual height of x. It turns out that the height is $O(\lg n)$ and therefore the space to store rank information is $O(\lg \lg n)$. In contrast the space to store size information is $O(\lg n)$.

¶10. Path Compaction heuristics. The size and rank heuristics apply to the Union Operation. We now consider heuristics that apply to the Find Operation. The first idea was introduced by McIlroy and Morris. When doing a Find on an item x, we traverse the path from x to the root of its tree. This is called the **find-path** of x. We specify a transformation of the compressed tree to accompany each Find operation:

Path Compression Heuristic: After performing a Find on x that returns the root u, modify the parent pointer of each node z along the find-path of x to point to u.

For example, if the find-path of x is (x, w, v, u) as in Figure 4, then after Find(x), the path compression heuristic will make x and w into children of u (note that v remains a child of u).

An alternative description of path compression is that it changes the find-path from a list into an anti-list.

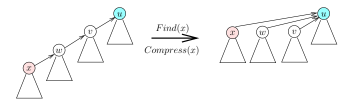


Figure 4: Path compression heuristic.

Path compression requires two passes along the find-path, first to locate u and the second time to change the parent pointer of nodes along the path. To be concrete, you might implement the Find operation with path compression as follows:

229

228

205

206

207

208

210

211

213

214

215

216

217

219

221

Lecture XIII Page 9 §2. Two Heuristics

```
Find(x):
    u \leftarrow x
     While (u.p \neq u)
         u \leftarrow u.p
     y \leftarrow x.p
     While (y \neq u)
                                \triangleleft INVARIANT: y = x.p
         x \leftarrow y; \ y \leftarrow y.p
```

Can we avoid using two while-loops? Path compression can be seen as a member of the family 232 of path compaction heuristics. Such a heuristic specifies a rule to modify the parent pointer 233 of each node z along a find-path; we require the new parent of z to be further up along the 234 find-path. Note that if z is the root or a child of the root, then the path compaction heuristic 235 has no effect on z (more precisely, on z.p).

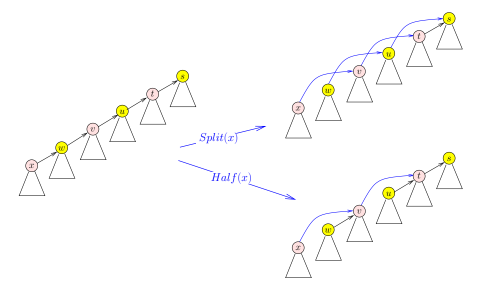


Figure 5: Path Compaction: Splitting and Halving

The "trivial" path compaction heuristic is the one that never changes z.p for any z on the path. Path compression is the other extreme, where each z.p is made to point to the ultimate ancestor, the root. We now consider two intermediate forms of path compaction, from van der Weide and van Leeuwen (1977). The path splitting heuristic says that for each node z along the find-path, we should update its parent pointer to its grandparent: $z.p \leftarrow (z.p).p$. Of course, (z.p).p is the grandparent of z, and it may also be written as

$$z.p.p$$
 or $z.p^2$.

The effect of this heuristic is to split the find-path into two paths comprising the odd and even nodes, respectively. Both the odd and even paths are about half the original length. 238 See Figure 5(a) where the find-path (x-w-v-u-t-s) is split into two paths (x-v-t-s) and 239 (w-u-s). The path splitting heuristic can now be implemented with only one pass over the 240 find-path, and in this sense is an improvement over path compression. A further improvement 241 is the following variant:

231

§2. Two Heuristics Lecture XIII Page 10

Path Halving Heuristic: After performing a Find on x, modify the parent pointer of every other node z along the find-path of x to point its grandparent.

In Figure 5(b), we see that the find-path (x-w-v-u-t-s) is halved into the path (x-v-t-s). The other node w and u join this half path at intermediate points. The halving heuristic is an improvement over path splitting in the sense that it requires only half of the number of pointer assignments.

Although the splitting and halving heuristics have the advantage of being 1-pass algorithms, they do not instantly reduce the find-path of elements along its path to length O(1) like in path compression. So it is not immediately clear these two heuristics can match the performance of path compression.

¶11. Analysis of the rank heuristic. We state some simple properties of the rank function. Notice that the rank function is defined according to equation (4); it is well-defined whether or not we use any of the heuristics for linking or path compression.

Lemma 1

243

247

249

250

251

252

253

255

256

257

259

260

261

262

263

264

265

267

269

271

The rank function (whether or not the rank heuristic is used, and even in the presence of path compression) has these properties:

- (i) A node has rank 0 iff it is a leaf.
- (ii) The rank of a node x does not change after x has been linked to another node.
- (iii) Along any find-path, the rank is strictly increasing.

If the rank heuristic is used, then the rank function has additional properties:

- (a) $rank(x) \leq degree(x)$ and $rank(x) \leq \lg(size(x))$.
- (b) No path has length more than $\lg n$.
- (c) In any sequence of Links/Find requests, for any $k \ge 1$, the number of times that the rank of any item gets promoted from k-1 to k is at most $n2^{-k}$.

Proof. Parts(i)-(iii) are immediate. Part(a) is shown by induction: it is true when x is initially a singleton, since rank(x) = degree(x) = 0 and size(x) = 1. We must look at events that causes the rank, degree or size of any node x to change. When we link y to x, the degree of x increases by one, and the size of x increased. If the rank of x did not change (this happens because rank(y) < rank(x)) then the truth of property (a) is preserved by the linking. If the rank of x changes, it must have increased by 1 and this resulted from y having the same rank as x. Let rank'(x), degree'(x) and size'(x) be the new rank, degree and size of x. Then rank'(x) = 1 + rank(x), degree'(x) = 1 + degree(x) and size'(x) = size(x) + size(y). Thus $rank'(x) \le degree'(x)$ follows from $rank(x) \le degree(x)$. We come to the key argument:

$$2^{rank'(x)} = 2^{rank(x)+1} = 2^{rank(x)} + 2^{rank(y)} \le size(x) + size(y) = size'(x).$$

Thus $rank'(x) \le \lg size'(x)$.

(b) Consider a path of length ℓ in a tree rooted at x. By part(iii), the $\ell \leq rank(x)$. By part(a), $rank(x) \leq \lg n$. Thus $\ell \leq \lg n$.

(c) Suppose x and y are two items that were promoted from rank k-1 to k at two different times. Let T_x and T_y be the subtrees rooted at x and y immediately after the respective promotions. Clearly T_x and T_y are disjoint. By $\operatorname{part}(a)$, $\operatorname{size}(x) \geq 2^{\operatorname{rank}(x)} = 2^k$, and similarly $\operatorname{size}(y) \geq 2^k$. There can be at most $n/2^k$ such trees. Q.E.D. Using the

rank heuristic alone, each Find operation takes $\mathcal{O}(\log n)$ time, by property (b). This gives an overall complexity bound of $\mathcal{O}(m \log n)$. It is also easy to see that $\Omega(m \log n)$ is a lower bound if only the rank heuristic is used.

¶12. Analysis of the path compression heuristic. We will show (see Theorem 7 below) that with the path compression heuristic alone, a charge of $\mathcal{O}(\log n)$ for each Find is sufficient. Again this leads to a complexity of $\mathcal{O}(m \log n)$.

So both the path compression heuristic alone and rank heuristic alone lead to the same complexity bound of $\mathcal{O}(m\log n)$. But closer examination shows important differences: unlike the rank heuristic, we cannot guarantee that each Find operation takes $\mathcal{O}(\log n)$ time under path compression. Thus, this $\mathcal{O}(m\log n)$ bound is a true amortization bound and not a worst case bound. On the other hand, path compression has the advantage of not requiring extra storage (the rank heuristic requires up to $\lg \lg n$ extra bits per item). Hence there is an interesting tradeoff. To prove a lower bound on path compression heuristic, we use binomial trees. A binomial tree is any tree from an infinite family

$$B_0, B_1, B_2, \ldots,$$

of trees defined recursively as follows: B_0 is trivial tree with one node, and B_{i+1} is obtained from two copies of B_i such that the root of one B_i is a child of the other root (see Figure 6). Clearly, the size of B_i is 2^i . Note that we regard B_i as an non-oriented or un-ordered tree (i.e., the children of a node is not ordered).

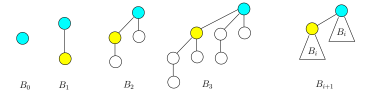


Figure 6: Binomial trees

The next lemma shows some nice properties of binomial trees (the arguments are routine and left as an Exercise).

Lemma 2

- (i) B_i has 2^i nodes.
- (ii) For $i \geq 1$, B_i can be decomposed into its root together with a sequence of subtrees of shapes

$$B_0, B_1, \ldots, B_{i-1}.$$

285 (iii) B_i has depth i. Moreover, level j (for $j=0,\ldots,i$) has $\binom{i}{j}$ nodes. In particular, B_i has a unique deepest node at level i.

Property (iii) is the reason for the name "binomial trees". To illustrate this lemma, consider B_3 in Figure 6: it has $8=2^3$ nodes as claimed by Lemma 2(i). The root has three children, each representing subtrees with shapes B_2, B_1, B_0 , as stated in Lemma 2(ii). Finally, levels 0, 1, 2, 3 (resp.) of B_3 have 1, 3, 3, 1 nodes. This is equal to $\binom{3}{0}$, $\binom{3}{1}$, $\binom{3}{2}$, $\binom{3}{3}$, as expected by Lemma 2(iii).

¶13. A Self-reproducing Property of Binomial Trees. M.J. Fischer observed an interesting property of binomial trees under path compression. Let $B_{i,k}$ denote any compressed tree which

- has size $k+2^i$,
- contains a copy of B_i , and
 - the root of this B_i coincides with the root of $B_{i,k}$.

Note that $B_{i,0}$ is just B_i . A copy of B_i that satisfies this definition of $B_{i,k}$ is called an "anchored B_i ". There may be many such anchored B_i 's in $B_{i,k}$. A node in $B_{i,k}$ is **distinguished** if it is the deepest node of some anchored B_i . The right-hand side of Figure 7 shows an instance of $B_{3,1}$.

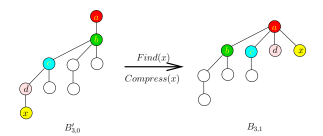


Figure 7: Self-reproducing binomial tree B_3 .

Also let $B'_{i,k}$ denote the result of linking the root of $B_{i,k}$ to a singleton. Distinguished nodes of $B'_{i,k}$ are inherited from the corresponding $B_{i,k}$. The left-hand side of Figure 7 illustrates $B'_{3,0}$.

Lemma 3 Suppose we perform a Find on any distinguished node of $B'_{i,k}$. Under the path compression heuristic, the result has shape $B_{i,k+1}$.

This lemma is illustrated in Figure 7 for $B'_{3,0}$ (the node x is the only distinguished node here). We now obtain a lower bound for path compression heuristics as follows. First perform a sequence of links to get B_i such that B_i contains between n/4 and n/2 nodes. Thus $i \ge \lg n - 2$. Then we perform a sequence of Links and Finds which reproduces B_i in the sense of the above lemma. This sequence of operations has complexity $\Omega(m \log n)$. Our construction assumes $m = \Theta(n)$ but it can be generalized to $m = \mathcal{O}(n^2)$ (Exercise).

Note that if we use path halving heuristic, the resulting tree would be a bit harder to analyze.

At this point, you have the complete algorithmic description of the (basic) Union-Find data structure and its algorithms. To understand this data structure really well, we suggest an implementation project in your favorite programming language. The rest of this chapter is concerned with the analysis of its complexity and applications of this data structure.

307

308

310

311

316 Exercises 317 Exercise 2.1: (1) Programming Project: implement the Union-Find data structure in your 318 favorite programming language. 319 (2) Use it to solve any of the three applications in the the final section of this chapter 320 (Kruskal's algorithm, Optimized Expressions, Betti Numbers). 321 **Exercise 2.2:** Suppose we start out with 5 singleton sets $\{a\}, \{b\}, \{c\}, \{d\}, \{e\},$ and we perform 322 a sequence of Link/Find operations. Assume the path compression heuristic, but not the 323 rank heuristic. The cost of a Link is 1, and the cost of a Find is the number of nodes in 324 325 (a) Construct a sequence of 10 Link/Find operations that achieves as large a cost as you can. It is possible to achieve a cost of 22. 327 (b) Suppose the cost of a Find is twice the number of nodes in the find-path (this is more 328 realistic for implementing path compression). This impacts the actual worst case cost, 329 but would your sequence of 12 operations in (a) have changed? **Exercise 2.3:** Show that when the size heuristic is used, then height of x is at most $\lg(size(x))$. 331 332 **Exercise 2.4:** We need to allocate $\mathcal{O}(\log \log n)$ extra bits to each node to implement the rank 333 heuristic. Argue that the rank heuristic only uses a **total** of $\mathcal{O}(n)$, not $\mathcal{O}(n \log \log n)$, 334 extra bits at any moment. In what sense is this fact hard to exploited in practice? [Idea: 335 to exploit this fact, we do not pre-allocate $\mathcal{O}(\log \log n)$ bits to each node.] \Diamond 336 **Exercise 2.5:** Suppose we have a full binary tree with n leaves. At each internal node u, we 337 associate an integer which is equal to the minimum of the number of leaves in the left 338 subtree at u and the number of leaves in the right subtree at u. Prove that the sum of all 339 the integers in the internal nodes is at most nH_n where H_n is the nth harmonic number. 340 NOTE: this binary tree represents a sequence of Unions Operations where the cost of the 341 union of two sets is the minimum of the sizes of the two sets. This result proves that the 342 total cost is $O(n \log n)$. ♦ 343 Exercise 2.6: Prove lemma 2 and lemma 3. \Diamond Exercise 2.7: (Chung Yung) Assuming naive linking with the path compression heuristic, 345 construct for every $i \geq 0$, a sequence $(l_1, f_1, l_2, f_2, \dots, l_{2^i}, f_{2^i})$ of alternating link and find 346 requests such that the final result is the binomial tree B_i . Here l_i links a compressed tree 347

Exercise 2.8: In the lower bound proof for the path compression heuristic, we have to make the following basic transformation:

to a singleton element and f_i is a find operation.

$$B_{i,k} \longrightarrow B'_{i,k} \longrightarrow B_{i,k+1}$$
 (5)

Beginning with $B_{i,0}$, we repeat the transformation (5) to get an arbitrarily long sequence

 $B_{i,1}, B_{i,2}, B_{i,3} \dots$ (6)

348

351 352 \Diamond

354

355

356

357

361

362

363

365

366

367

368

369

370

371

372

373

374

375

376

377

378

379

382

(a) Can you always choose your transformations so that the trees in this sequence has the form $B_{i+1,k}$ (for some $k \geq 0$)? (b) Characterize those $B_{i,k}$'s that are realizable using the lower bound construction in the text. HINT: first consider the case i = 1 and i = 2. You may restrict the transformation (5) in the sense that the node for Find is specially chosen from the possible candidates.

Exercise 2.9: For any $m \ge n$, we want a lower bound for the union-find problem when the path compression heuristic is used alone.

(a) Fix $k \ge 1$. Generalize binomial trees as follows. Let $F_k = \{T_i : i \ge 0\}$ be a family of trees where T_i is a singleton for $i = 0, \ldots, k-1$. For $i \ge k$, T_i is formed by linking a T_{i-k} to a T_{i-1} . What is the degree d(i) and height h(i) of the root of T_i ? Show that the size of T_i is at most $(1+k)^{(i/k)-1}$.

(ii) [Two Decompositions] Show that if the root T_i has degree d then T_i can be constructed by linking a sequence of trees

$$t_1, t_2, \dots, t_d \tag{7}$$

to a singleton, and each t_i belongs to F_k . Show that T_{i+1} can also be constructed from a sequence

$$s_0, s_1, \dots, s_p, \quad p = \lfloor (i+1)/k \rfloor$$
 (8)

of trees by linking s_j to s_{j-1} for $j=1,\ldots,p$, and each s_j belongs to F_k . The decompositions (7) and (8) are called the **horizontal** and **vertical decompositions** of the respective trees. Also, s_p in (8) is called the **tip** of T_{i+1} .

(iii) [Replication Property] Let T' be obtained by linking T_i to a singleton node R. Show that there are k leaves x_1, \ldots, x_k in T' such that if we do finds on x_1, \ldots, x_k (in any order), path compression would transform T' into a copy of T_i except the root has an extra child. HINT: Consider the trees $r_0, r_1, \ldots, r_{k-1}, r_k$ where $r_j = T_{i-k-j}$ (for $j = 0, \ldots, k-1$) and $r_k = T_{i-k}$. Note that T_i can be obtained by linking each of $r_0, r_1, \ldots, r_{k-1}$ to r_k . Let C be the collection of trees comprising r_k and the trees of the vertical decomposition of r_j for each $j = 0, \ldots, k-1$. What can you say about C?

(iv) Show a lower bound of $\Omega(m \log_{1+(m/n)} n)$ for the union-find problem when the path compression heuristic is used alone. HINT: $k = \lfloor m/n \rfloor$. Note that this bound is trivial when $m = \Omega(n^2)$.

Exercise 2.10: Explore the various compaction heuristics.

End Exercises

 \Diamond

§3. Multilevel Partition

The surprising result is that the combination of rank and path compression heuristics leads to a dramatic improvement on the $\mathcal{O}(m \log n)$ bound of either heuristic. The sharpest analysis of this result is from Tarjan, giving almost linear bounds. We give a simplified version of Tarjan's analysis.

¶14. Partition functions. A function

 $a: \mathbb{N} \to \mathbb{N}$

405

406

407

408

409

417

is a partition function if a(0) = 0 and a(j) < a(j+1) for all $j \ge 0$. Such a function induces a partition of the natural numbers N where each block of the partition has the form 389 [a(j)...a(j+1)-1], or more compactly denoted [a(j)...a(j+1)). For instance, if a is the identity 390 function, a(j) = j for all j, then it induces the **discrete partition** where each block has exactly 391 one number. 392

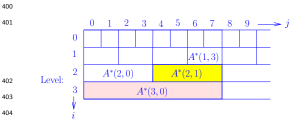
A multilevel partition function is $A : \mathbb{N} \times \mathbb{N} \to \mathbb{N}$ such that:

- (a) For each $i \geq 0$, $A(i,\cdot)$ is a partition function, which we denote by A_i . The partition on \mathbb{N} induced by A_i is called the **level** i partition. 395
- (b) The level 0 partition function A_0 is the identity function, inducing the discrete partition. Moreover, level 0 is the only level to have blocks of size 1. 397
- (c) The level i partition is a coarsening of the level i-1 partition.

Let block(i, j) = [A(i, j)...A(i, j + 1)] denote the jth **block** of the level i partition. For $i \ge 1$, let

$$b(i,j) \ge 1$$

denote the number of i-1st level blocks whose union equals block(i,j).



A simple example of a multilevel partition function is A^* defined by

$$A^*(i,j) := j2^i, \qquad (i,j \ge 0).$$
 (9)

For reference, call this the binary par**tition function**. In this case b(i, j) =2 for all $i \geq 1, j \geq 0$.

Figure 8: Binary Partition Function, $A^*(i,j)$

¶15. Basic Goal. Our goal is to analyze path compaction heuristics ($\P10$), without necessarily assuming that the rank heuristic is used. Recall the rank function defined in equation (4); we said it is meaningful even if we do not use the rank heuristic. Of course, if we do not use the rank heuristic then we need not maintain rank information in our data structure. In this analysis, we assume some fixed sequence

$$r_1, r_2, \dots, r_m \tag{10}$$

of m Links/Find request on a universe of n items.

¶16. The eventual rank of an item. The rank of an item is monotonically non-decreasing with time. Relative to the sequence (10) of requests, we may speak of the **eventual rank** of 412 an item x. E.g., if x is eventually linked to some other item, its eventual rank is its rank just 413 before it was linked. To distinguish the two notions of rank, we will write erank(x) to refer 414 to its eventual rank. For emphasis, we say "current rank of x" to refer to the time-dependent 415 concept of rank(x). 416

¶17. The level of an item. The notion of "level of an item" is defined relative to some choice of multilevel partition function A(i, j). The function A(i, j) is used in the analysis only, 418

and does not figure in the algorithms. We say x has **level** i (at any given moment) if i is the smallest integer such that the eventual ranks of x and of its current parent x.p are in the same ith level block. Note the juxtaposition of current parent with eventual rank in this definition.

For instance, assuming the binary partition function A^* of (9), if $\operatorname{erank}(x) = 2$ and $\operatorname{erank}(x.p) = 5$ then $\operatorname{level}(x) = 3$ (since 2,5 both belong in the level 3 block [0,8) but are in different level 2 blocks). Although $\operatorname{erank}(x)$ is fixed, x.p can change and consequently the level of x may change with time. Notice that x is a root if and only $\operatorname{erank}(x.p) = \operatorname{erank}(x)$. Hence x is a root iff it is at level 0.

We leave the following as an easy exercise:

Lemma 4 Assume that some form of compaction heuristic is used. The following holds, whether or not we use the rank compression heuristics:

- (i) The erank of nodes strictly increases along any find-path.
- (ii) For a fixed x, erank(x.p) is non-decreasing with time.
- 432 (iii) The level of node x is 0 iff x is a root.

430

440

441

442

444

- (iv) The level of any node is non-decreasing over time.
- (v) The levels of nodes along a find-path need not be monotonic (non-decreasing or nonincreasing), but it must finally be at level 0.

¶18. Bound on the maximum level. Let $\alpha \geq 0$ denote an upper bound on the level of any item. Of course, the definition of a level depends on the choice of multilevel partition function A(i,j). Trivially, all ranks lie in the range [0..n]. If we use the binary partition function A^* in (9) then we may choose $\alpha = \lceil \lg(n+1) \rceil$, since $[0..n] \subseteq block(k,0)$ for $k \geq \lg(n+1)$.

In general, we could choose α to be the smallest i such that A(i,1) > n since all ranks are at most n. But if the rank heuristic is used, all ranks are at most $\lg n$ and therefore, we could define α to be the smallest i such that $A(i,1) > \lceil \lg n \rceil$. In particular, if we use the binary partition function A^* we may choose

$$\alpha = \lceil \lg \lg(2n) \rceil. \tag{11}$$

¶19. Charging scheme. We charge one unit for each Link operation; this is sufficient to pay for the cost of Linking. The cost of a Find operation is proportional to the length of the corresponding find-path. To charge this operation, we introduce the idea of a level account for each item x: for each $i \ge 1$, we keep a charge account for x at level i. We charge the Find operation in two ways:

- **Upfront Charge**: we simply charge this Find $1 + \alpha$ units.
- Level Charges: For each node y along the find-path, we charge one unit to y's account at level i for each $i \in [level(y)..level(y.p)]$.

Note that if level(y.p) < level(y), no charges to y's accounts are made. If $level(y.p) \ge level(y)$, then y incurs a total debit of 1 + level(y.p) - level(y) units over all the levels.

¶20. Justification for charging scheme. To justify the charging scheme, recall that we must verify the credit-potential invariant (§VI.1).

Lemma 5 The charge for each Find operation covers the cost of the operation plus any increase in the level accounts.

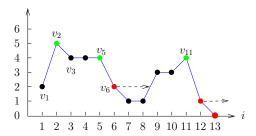


Figure 9: Plot of $level(x_i)$ against i.

This is best shown visually as follows. Let the find-path be $(x_1, x_2, ..., x_h)$ where x_h is the root. We must show that $Find(x_1)$ is charged at least h units by the above charging scheme. Plot a graph of $level(x_i)$ against i. As illustrated in Figure 9, this is a discrete set $v_1, v_2, ..., v_h$ of points where $v_i = (i, level(x_i))$. By lemma 4(v), the level of v_h is 0. The graph is a polygonal path $(v_1-v_2-\cdots-v_h)$ with h-1 edges. An edge (v_i-v_{i+1}) is called **positive**, **zero** or **negative**, according to the sign of its slope. Thus Figure 9 has 3 positive edges, 5 negative edges and 4 zero edges. Our level charges amount to charging k+1 units to a nonnegative edge whose slope is $k \geq 0$; negative edges have no charges. In Figure 9, edge (v_8-v_9) has slope 2 and so has a level charge of 3. Our figure uses the following color code for the nodes in a path:

- v_i is **black** if the slope of (v_i-v_{i+1}) is non-negative. E.g., $v_1, v_3, v_4, v_7, v_8, v_9, v_{10}$ in Figure 9.
- Else, v_i is green if the slope of $(v_{i-1}-v_i)$ is non-negative. E.g., v_2, v_5, v_{11} in Figure 9.
- Else, v_i is **red**. This means that the slopes of $(v_{i-1}-v_i)$ (if i > 1) and (v_i-v_{i+1}) (if i < h) are both negative. E.g., v_6, v_{12}, v_{13} in Figure 9.

The total cost of a Find operation can be distributed to the vertices, so each vertex has unit cost. The cost of a black vertex v_i is covered by the level charges to (v_i-v_{i+1}) ; likewise, the cost of a green vertex v_i is covered by the level charges to $(v_{i-1}-v_i)$. But how do we cover the cost of a red vertex v_i ? This is the key geometrical insight: for each red vertex v_i , we cast a horizontal ray from v_i to the right. There are two possibilities.

- (1) This ray may never hit any edge, as in the case of the rays from v_{12} and v_{13} . Note that there are most $1 + \alpha$ of these rays since $0 \le level(x_i) \le \alpha$, and there is at most one ray to infinity at each level.
- 482 (2) This ray may hit a point on a positive edge, as in the case of the ray from v_6 hitting the edge (v_8-v_9) . We will cover the cost of v_i using the level charges associated with the positive edge. To see that this is a valid method, note that a positive edge with slope $k \ge 1$ may be hit

459

460

461

462

463

464

465

466

468

470

471

472

by no more than k rays. Since it is endowed with k+1 level charges, and one of these charges is to pay for the associated black vertex, we are left with k unused charges. One subtlety is that the ray may hit a zero edge as well. For instance, if the vertex v_6 in Figure 9 were raised one unit higher, it would hit the zero edge (v_9-v_{10}) . Nevertheless, it still hits a positive edge, and so we can avoid choosing the zero edge for this accounting.

¶21. The number of level charges. Let x be an item. For each $i \ge 1$, erank(x) belongs to block(i, j(i)) for some j(i).

Lemma 6 The charges to x at level $i \geq 1$ is at most b(i, j(i)).

To show this, note that whenever x is charged at level i, then

$$level(x.p) \ge i$$

holds just preceding that Find. This means the erank's of the parent x.p and grandparent $x.p^2$ belong to different i-1st level blocks. As erank's are strictly increasing along a find-path,
after this step, the new parent of x has erank in a new i-1st level block. But the indices of the i-1st level blocks of x.p are nondecreasing over time. Thus, after at most b(i,j(i)) charges,
erank(x.p) and erank(x.p) ie in different level i blocks. This means the level of x has become
strictly greater than i, and henceforth x is no longer charged at level i. This proves our claim.

Let n(i,j) denote the number of items whose erank's lie in block(i,j). Clearly, for any i,

$$\sum_{j\geq 0} n(i,j) = n. \tag{12}$$

The total level charges, summed over all levels, is at most

$$\sum_{i=1}^{\alpha} \sum_{j>0} n(i,j)b(i,j). \tag{13}$$

¶22. Bound for pure path compression heuristic. Suppose we use the path compression heuristic but not the rank heuristic. Let us choose $A^*(i,j)$ to be the binary partition function and so we may choose $\alpha = \lg(n+1)$. Then the level charges, by equations (12)–(13), is at most

$$\sum_{i=1}^{\lg(n+1)} \sum_{j \ge 0} n(i,j) 2 = 2n \lg(n+1).$$

If there are m' Find requests, the upfront charges amount to $\mathcal{O}(m'\log n)$. This proves:

Theorem 7 Using the path compression heuristic, but not the rank heuristic, a sequence of m Link/Find Requests on a universe of n items has total cost $\mathcal{O}(m + (m' + n) \log n)$ where m' is the number of Find requests.

The number m' of Find operations is arbitrary (in particular, we do not assume $m' \geq n$). This has application in a situation where path compression is allowed but not the rank heuristic (see [7]).

499

Remark: The above definition of level is devised to work for any of the compaction heuristics noted in §2. A simpler notion of level will work for path compression heuristic (see Exercise).

EXERCISES _____

Exercise 3.1: (Chung Yung) Define the level of x to be the least i such that both x and the root r (of the tree containing x) have eranks in the same ith level block. Give a charging scheme and upper bound analysis for path compression using this definition of level. \diamondsuit

Exercise 3.2: Prove the analogues of Theorem 7 in which we replace path compression by path splitting, or by path halving.

518 _____END EXERCISES

§4. Combined Rank and Path Compression Heuristics

We are now ready to prove the main result of this chapter:

Theorem 8 When the rank and path compression heuristics are employed, a sequence of m Links/Find requests over a universe of n items has total cost

$$\mathcal{O}((m+n)\alpha(n))$$

Here $\alpha(n)$ is the inverse Ackermann function to be defined. We wish to emphasize that this is deep result about a simple data structure with simple algorithms.

Much of the groundwork has been laid. Indeed, we can glean some immediate bounds. Because of the rank heuristic, all ranks lie in the range $[0..\lg n]$. Using the binary partition function A^* , we already noted that the level of any item is $O(\lg\lg n)$ (see equation (11)). This immediately gives a bound of

$$\mathcal{O}((m+n)\lg\lg n)$$

on the problem. To get a substantially better bound, we need new multilevel partition function called the Ackermann function A(i, j). The original analysis of Tarjan uses a related function B(i, j).

¶23. Ackermann Functions. A number theoretic function $A: \mathbb{N} \times \mathbb{N} \to \mathbb{N}$ is called an **Ackermann function** if satisfies the following "Ackermann recurrence":

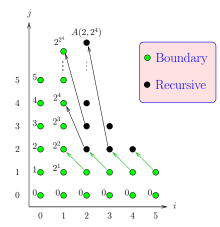


Figure 10: Ackermann's function

A(i,j) = A(i-1, A(i,j-1))(14)

524

526

527

528

529 530

510

511

Page 19

for i, j large enough. A particular instance of the Ackermann function depends on the choice of the boundary conditions. In the following, we use this boundary conditions:

$$A(i,j) = \begin{cases} 0 & \text{if } j = 0, \\ j & \text{if } i = 0, \\ 2^j & \text{if } i = 1, j \ge 1, \\ A(i-1,2) & \text{if } j = 1, i \ge 2. \end{cases}$$

$$(15)$$

This determines the values of A(i, j) when i < 2 or when j < 2. For $i \ge 2$ and $j \ge 2$, the Ackermann recurrence (14) is applicable.

Some Values of the Ackermann function. Figure 10 is a useful visual aid: the domain $\mathbb{N} \times \mathbb{N}$ is represented by an infinite array of grid points. The values of A on the green points are given by the boundary conditions (15). The values at black points are given by the recurrence (14). It follows that

$$A(2,j) = A(1, A(2, j - 1)), j \ge 2$$

$$= \begin{cases} 2^2, & j = 2\\ 2^{A(2, j - 1)}, & j > 2 \end{cases}$$

$$= \exp_2^{(j)}(2) = 2^{2^{j-1}}$$

(a stack of j 2's). Also A(3,1) = A(2,2) = A(1,4) = 16 and A(4,1) = A(3,2) = A(2,16). The last number is more than the number of atoms in the known universe (last estimated at 10^{80}). The dominant feature of an Ackermann function is its explosive growth. It grows faster than any primitive recursive function (a class of fast growing functions in complexity theory).

10⁸⁰ is taken from Wikipedia (2013).

Page 20

Lemma 9 The Ackermann function A(i,j) is a multilevel partition function.

Proof. Conditions a) and b) in the definition of a multilevel partition function are immediate. For condition c) we must show that the *i*th level partition determined by A is a coarsening of the i-1st level partition. This amounts to noting that for all $i \geq 1, j \geq 1$, we can express A(i,j) as A(i-1,j') for some j'.

Q.E.D.

Let us note some simple properties the Ackermann function.

• The 1st level partition consists of exponentially growing block sizes,

$$block(1, j) = [2^{j}..2^{j+1} - 1].$$

ullet We want to determine an upper bound lpha on the level of an item. This can be taken to be the smallest i such that

$$[0.. \lg n] \subseteq block(i, 0).$$

Clearly, this is the smallest i such that $A(i,1) > \lg n$. In fact, let us define the following inverse Ackermann function:

$$\alpha(n) := \min\{i : A(i, 1) > \lg n\}.$$

$$\alpha'(k,\ell) := \min\{i \ge 1 : A(i, \left\lfloor \frac{k}{\ell} \right\rfloor) > \lg \ell\}.$$

The one-argument function $\alpha'(\ell)$ may be defined as $\alpha'(\ell,\ell)$.

²We could also use this simpler definition for inverse Ackermann, $\alpha(n) := \min\{i : A(i,1) > n\}$. This modification has no material impact on our conclusions. Tarjan uses the following two-argument form inverse Ackermann function, $\alpha'(k,\ell)$, $k \ge \ell \ge 1$, where

It follows that the level of every item is at most $\alpha = \alpha(n)$. The dominant feature of $\alpha(n)$ is its very slow growth rate. For all practical purposes, $\alpha(n) \leq 4$ (see exercise below).

• The number of i-1st level blocks that form block(i,j) is given by the following:

$$b(i,j) = \begin{cases} 2 & \text{if } j = 0, \\ A(i,j) - A(i,j-1) & \text{if } j \ge 1. \end{cases}$$
 (16)

To see that b(i,0) = 2, we note that A(i,1) = A(i-1,2). For $j \ge 1$, we note that A(i,j+1) = A(i-1,A(i,j)).

• The number n(i,j) of items whose eranks lie in block(i,j) is bounded by:

$$n(i,j) \le \sum_{k=A(i,j)}^{A(i,j+1)-1} \frac{n}{2^k} \le \frac{2n}{2^{A(i,j)}},$$

where the first inequality comes from lemma 1(c).

• It follows that the number of charges at level $i \geq 1$ is bounded by

$$\sum_{j>0} b(i,j)n(i,j) \le b(i,0)n(i,0) + \sum_{j>1} \frac{A(i,j) \cdot 2n}{2^{A(i,j)}} \le 2n + 2n \sum_{k>2} \frac{k}{2^k} = 5n$$

where the last equality is the following sum when k=2.

$$\sum_{j=k}^{\infty} \frac{j}{2^j} = \frac{k+1}{2^{k-1}}.\tag{17}$$

To prove (17), observe that

$$\frac{j}{2^{j}} = \frac{j+1}{2^{j-1}} - \frac{j+2}{2^{j}}$$

$$\sum_{j=k}^{\infty} \frac{j}{2^{j}} = \left(\frac{k+1}{2^{k-1}} - \frac{k+2}{2^{k}}\right) + \left(\frac{k+2}{2^{k}} - \frac{k+3}{2^{k+1}}\right) + \left(\frac{k+3}{2^{k+1}} - \frac{k+4}{2^{k+2}}\right) + \cdots$$

$$= \frac{k+1}{2^{k-1}} - \left(\frac{k+2}{2^{k}} - \frac{k+2}{2^{k}}\right) - \left(\frac{k+3}{2^{k+1}} - \frac{k+3}{2^{k+1}}\right) - \cdots$$

$$= \frac{k+1}{2^{k-1}}.$$

Since there are $\alpha(n)$ levels, there is a bound of $\mathcal{O}(n\alpha(n))$ on all the level charges. Combined with the $\mathcal{O}(\alpha(n))$ Upfront charge for each Find, this proves our main theorem Theorem 8.

¶24. Amortization Framework. In Lecture VI, we presented the potential framework. We briefly review this in the setting of the Union-Find problem. We are given a sequence p_1, \ldots, p_m of requests to process. Our amortization scheme establishes a finite set of **accounts** A_1, \ldots, A_s such that the cost of each request p_i is distributed over these s accounts. More precisely, the scheme specifies a **charge** by p_i to each account A_j .

In our Link/Find analysis, we set up an account for each operation p_i (this is called the up-front charges for find requests) and for each item x at each level ℓ , we have an account $A_{x,\ell}$. Note that the charges to each account is non-negative value (these are pure debit accounts).

If $Charge(A_j, p_i)$ units is charged to account A_j by request p_i , we require

$$\sum_{i=1}^{s} Charge(A_j, p_i) \ge Cost(p_i). \tag{18}$$

556

564

If $Charged(A_j)$ is the sum of all the charges to A_j , then the amortization analysis amounts to obtaining an upper bound on $\sum_{j=1}^{s} Charged(A_j)$.

We may combine the charge account framework with the potential framework. We now allow an account to be **credited** as well as debited. To keep track of credits, we associate a **potential** $\Phi(A_j)$ to each A_j and let $\Delta\Phi(A_j, p_i)$ denote the increase in potential of A_j after request p_i . Then (18) must be generalized to

$$\sum_{j=1}^{s} (Charge(A_j, p_i) - \Delta\Phi(A_j, p_i)) \ge Cost(p_i).$$
(19)

Without loss of generality, assume that $\Phi(A_j)$ is initially 0 and let Φ_j denote the final potential of A_j . The total cost for the sequence of operations is given by

$$\sum_{j=1}^{s} (Charged(A_j) - \Phi_j).$$

If we are interested in the "amortized cost" of each type operation, this amounts to having an account for each operation p_i and charging this account an "amortized cost" corresponding to its type. We insist that $\sum_{j=1}^{s} \Phi_j \geq 0$ in order that the amortized cost is meaningful.

570 EXERCISES

Exercise 4.1: Verify the claim that $A(4,1) > 10^{80}$ (number of atoms in the known universe).

Exercise 4.2: Compute A(3,3).

Exercise 4.3: What is the smallest j such that A(2, j) is larger than a virgintillion (the number 10^{63})? Larger than a googleplex (the number $10^{10^{100}}$)?

Exercise 4.4:

577

579

581

583

584

585

- (i) Show A(i, j) is strictly increasing in each coordinate.
- (ii) When is $\alpha(m,n) \leq 1$? When is $\alpha(m,n) \leq 2$?
- (iii) Compute the smallest ℓ such that $\alpha(\ell)$ is equal to 0, 1, 2, 3, 4.

Exercise 4.5: (Tarjan) Define a two argument version of the inverse Ackermann function, $\alpha'(k,\ell), \ k \geq \ell \geq 1$, where

$$\alpha'(k,\ell) := \min\{i \ge 1 : A(i, \left\lfloor \frac{k}{\ell} \right\rfloor) > \lg \ell\}.$$

The one-argument function $\alpha'(\ell)$ may be defined as $\alpha'(\ell,\ell)$, and could be used instead of the $\alpha(\ell)$ function used in the text. Note that $\alpha'(k,\ell)$, for fixed ℓ , is actually a decreasing function in k. Improve the amortized upper bound to $\mathcal{O}(m\alpha'(m+n,n))$.

Exercise 4.6: Consider the following multilevel partition function: $A(i,j) = \lfloor \lg^{(i)}(j) \rfloor$ (*i*-fold application of \lg to j). Fix your own boundary conditions in this definition of A(i,j). Analyze the union-find heuristic using this choice of multilevel partition function. \diamondsuit

 \Diamond

Exercise 4.7: Prove similar upper bounds of the form $\mathcal{O}(m\alpha(n))$ when we replace the path compression heuristic with either the splitting or halving heuristics.

End Exercises

§5. Three Applications of Disjoint Sets

There are many applications of the disjoint set data structure. Three will be given here.
The first problem is in the implementation of Kruskal's algorithm for minimum spanning tree.
The second problem concerns restructuring of algebraic expressions, a problem which arise in optimizing compilers and in Exact Geometric Computation. The last problem concerns the computation of Betti numbers in in Computational Topology.

¶25. A. Implementing Kruskals' Algorithm for MST. This was our opening example in Table of the introduction. In Chapter V, we presented Kruskal's algorithm for minimum spanning tree (MST). We now show how to implement it efficiently using the Union Find data structure.

Recall the problem of computing the MST of an edge-costed bigraph G = (V, E; C), with edge costs $C(e), e \in E$. Kruskal's algorithm begins by sorting the edges in E in increasing order of their costs, breaking ties arbitrarily. We initialize $S = \emptyset$ and at each step, we consider the next edge e in the sorted list. We will insert e into the set S provided this does not create a cycle in S (otherwise e is discarded). Thus, inductively, S is a forest. We can stop as soon as the cardinality of S reaches |V| - 1; otherwise we stop when we have considered all the edges. The output is S, which is an MST if |S| = n - 1 and otherwise it is a minimum spanning Forest.

Implementation and complexity. We must be able to detect whether adding e to a forest S will create a cycle. This is achieved using a Union-Find structure on the set V and whose equivalence classes represent the connected components of S. If e = (u, v), then $S \cup \{e\}$ creates a cycle iff Find(u) = Find(v). Moreover, if $Find(u) \neq Find(v)$, we will next form the union of their components via Union(u, v). The algorithm makes at most 2m Finds and n Links. The amortized cost for these Link/Finds is

$$\mathcal{O}(m\alpha(n)).$$

This cost is dominated by the initial cost of $\mathcal{O}(m \log n)$ for sorting the edges. Hence the complexity of the overall algorithm is $\mathcal{O}(m \log n)$.

¶26. B. Expression Optimization Problem. Suppose you are given an arithmetic expression, represented by a directed acyclic graph (DAG) G, with operators in the internal nodes and numerical constants or variables at the leaves. For instance, G might represent the expression

$$\sqrt{2} + \sqrt{3} - \sqrt{2 + 3 - \sqrt{6}}$$

This expression is an instance of the more general expression with variables instead of constants:

 $\sqrt{x} + \sqrt{y} - \sqrt{x + y - \sqrt{xy}}. (20)$

587

588

589

595

597

599

601

603

604

Thus one possible representation of the expression (20) is given by G in Figure 11. We want to compute a "reduced expression" G' that is equivalent to G. In Figure 11, we show such a G'. What is "reduced" about G'? Notice that it has one fewer node. But in this application, the more important fact is that G' has only three square-root operators, in contrast to the four square-roots in G.

This is an important problem in the area of Exact Geometric Computation where we are interested in numerical approximations for the values val(G) of expressions. Note that even if there are variables like x or y in G, it is assumed that we know explicit constants for these variables so that val(G) is actually a number. The key problem is to decide when val(G) is really zero.

e can approximate val(G)to as many bits of accuracy as we like (using big number arithmetic). But if the expression is really zero, how do we know to stop? This problem can be solved when the expression involves only algebraic operations such as $+, -, \times, \div, \sqrt{\cdot}$. It turns out that we can compute a so-called zero **bound** $\beta(G) > 0$ for any expression G such that if $val(G) \neq 0$ then

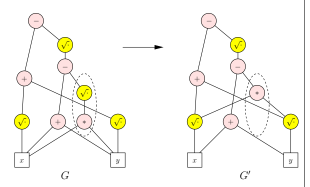


Figure 11: Reducing G to G'

 $|val(G)| > \beta(G)$. The graph G might be automatically generated by a program and can be quite large. As a result, $\beta(G)$ can be astronomically small, say $\beta(G) \simeq 2^{-10,000}$. This means that in order to decide if val(G) = 0, we need to approximate val(G) to over 10,000 bits. Suppose we have a reduced G' whose zero bound $\beta(G')$ is considerably larger (sic!) than $\beta(G)$, say $\beta(G') \simeq 2^{-100}$. The equivalence of expressions G and G' simply means that val(G) = val(G'). In this case, we only need to approximate val(G') = val(G) to about 100 bits, and thus deciding if val(G) = 0 is greatly sped up. The theory behind such zero bounds is beyond our scope.

We assume that the DAG G is represented as an adjacency list. The nodes with indegree 0 are called sources, and the nodes with outdegree 0 are called sinks. Each sink of G is labeled with a distinct variable name $(x_1, x_2, \text{ etc})$. Each non-sink node of G has outdegree 2 and are labelled with one of four arithmetic operations $(+, -, *, \div)$. See Figure 12, where the directions of edges are implicitly from top to bottom. Thus, every node of G represents an algebraic expression over the variables. E.g., node b represents the expression $x_2 - x_3$.

The two edges exiting from the '-' and '÷' nodes are distinguished (i.e., labeled as "Left" or "Right" edges) while the two edges exiting from + or * nodes are indistinguishable. This is because - and \div are non-commutative $(x-y\neq y-x,$ and $x\div y\neq y\div x$ in general) while + and * are commutative (x+y=y+x) and x*y=y*x. The Left/Right ordering of edges are implicit in our figures. In general, G is actually a multigraph because it is possible that there there is more than one edge from a node to another. E.g., the graph G' in Figure 12(b) has two edges from d to a.

Define two nodes to be **equivalent** if they are both internal nodes with the same operator label, and their "corresponding children" are identical or (recursively) equivalent. This is a

615

616

617

618

619

620

623

625

626

627

628

629

611

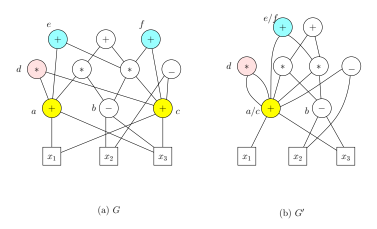


Figure 12: Two DAGs: G and G'

recursive definition. By "corresponding children" we mean that the order (left or right) of the children should be taken into account in case of - and \div , but ignored for + and * nodes. For instance, the nodes a and c in Figure 12(a) are equivalent. Recursively, this makes e and f equivalent.

The problem is to construct a **reduced DAG** G' from G in which each equivalence class of nodes in G are replaced by a single new node. For instance with G in Figure 12(a) as input, the reduced graph is G' in Figure 12(b).

The solution is as follows. The height of a node in G is the length of longest path to a sink. E.g., height of node e in Figure 12(a) is 3 (there are paths of lengths 2 and 3 from e to sinks). Note that two nodes can only be equivalent if they have the same height. Our method is therefore to checking equivalent nodes among all the nodes of a given height, assuming all nodes of smaller heights have been equivalenced. To "merge" equivalent nodes, we use a disjoint set data structure. If there are k nodes of a given height, the obvious approach takes $\Theta(k^2)$ time.

To avoid this quadratic behavior, we use a sorting technique: let A be the set of nodes of a given height. For each $u \in A$, we first find their children u_L and u_R using the adjacency lists of graph G. Then we compute $U_L = Find(u_L)$ and $U_R = Find(u_R)$. Hence U_L and U_R are the representative nodes of their respective equivalence classes. Let op(u) be the operator at node u. In case op(u) is + or *, we compare U_L and U_R (all nodes can be regarded as integers). If $U_R < U_L$, we swap their values. We now construct a 3-tuple

$$(op(u), U_L, U_R)$$

which serves as the key of u. Finally, we sort the elements of A using the keys just constructed. Since the keys are triples, we compare keys in a lexicographic order. We can assume an arbitrary order for the operators (say $'+' < ' -' < ' *' < ' \div '$). Two nodes are equivalent iff they have identical keys. After sorting, all the nodes that are equivalent will be adjacent in the sorted list for A. So, we just go through the sorted list, and for each u in the list, we check if the key of u is the same as that of the next node v in the list. If so, we perform a merge(u, v). This procedure takes time $O(k \lg k)$.

We must first compute the height of each node of G. This is easily done using depth-first search in O(m) time (see Exercise). Assume that all the nodes of the same height are put in a linked list. More precisely, if the maximum height is H, we construct an array A[1..H] such that A[i] is a linked list of all nodes with height i. Separately, we initialize a disjoint set

data structure for all the nodes, where initially all the sets are singleton sets. Whenever we discover two nodes equivalent, we form a union. We will now process the list A[i] in the order i = 1, 2, ..., H, using the sorting method described above.

Complexity: computing the height is O(m+n) (Exercise). Union Find is $m\alpha(n)$. Sorting of all the lists is $\sum_{i=1}^{H} O(k_i \log k_i) = O(n \log n)$ where $|A[i]| = k_i$. Hence the overall complexity is $O(m+n \log n)$.

Remark: We have actually solved a simple form of the general problem. For instance, the example in Figure 11 requires an additional transformation, $\sqrt{xy} \to \sqrt{x} \cdot \sqrt{y}$ before we can achieve the indicated reduction.

¶27. C. Computing Betti Numbers. Let K be a non-empty set of tetrahedra, triangles, edges and vertices, where each $s \in K$ is a closed set of \mathbb{R}^3 . Note that a tetrahedron (resp., triangle, edge, vertex) is known as a **simplex** of **dimension** 3 (resp., 2,1,0). Given two simplices s, s', we say s is a **proper face** of s' if s is contained in the closure of s', and $0 \le dim(s) < dim(s')$. For instance, a tetrahedron has 3 proper faces of dimension 2, 6 proper faces of dimension 1 and 4 proper faces of dimension 0. The empty set \emptyset is regarded as a simplex of dimension -1. Any simplex s has two **improper faces**, namely itself and \emptyset .

For K to be a **triangulation**, we require that for all $s, s' \in K$, (a) any face of s also belongs to K, and (b) $s \cap s'$ is a face of s and of s'. Thus, $\emptyset \in K$. The underlying space or support space of K is given by $|K| := \bigcup_{s \in K} s$. Delfinado and Edelsbrunner shows how we can compute $\beta_i(K)$, the i-th Betti number of K, using Union Find as follows. The Betti numbers are defined algebraically and is a topological invariant of |K|. For $|K| \subseteq \mathbb{R}^3$, these numbers have an intuitive meaning: β_0 is the number of connected components of |K|, β_1 is the number of holes of |K|. E.g., the donut (viewed as a solid) and the torus (viewed as the surface of a donut) has $\beta_1 = 1$. Finally, β_2 is the number of voids on |K|. E.g., the torus and the sphere S^2 (boundary of a solid ball) has one void $(\beta_2 = 1)$ each, but the donut has none, $\beta_2 = 0$. Let us just consider the case where K is just a graph, i.e., a set of vertices and edges only. The following basic lemma is all that we need:

Lemma 10 Let K and K' be 1-dimensional complexes, and $K' = K \cup \{s\}$ where s is an edge connecting u, v. If u, v lies in the same connected component of K, then $\beta_1(K') = \beta_1(K) + 1$; otherwise $\beta_0(K') = \beta_0(K) - 1$. The other Betti number is unchanged.

Note that if $K' = K \cup \{s\}$ and s is a vertex, then it is clear that $\beta_0(K') = \beta_0(K) + 1$ and the other Betti number is unchanged. Armed with this lemma, we can now give a simple incremental algorithm to maintain Betti numbers of K: we initialize b_0 to n, the number of vertices in K, and b_1 to 0. The n vertices are used to initialize a union-find data structure. Now we process each edge of K in turn. For each edge (u, v), if Find(u) = Find(v), then we perform Union(u, v) and update $b_1 = b_1 + 1$; otherwise we update $b_0 = b_0 - 1$. The final values of b_0, b_1 is the Betti number we seek.

The problem with extending this to 2-dimensional complexes K is that we do not have an efficient way to detect when the addition of a new triangle will create a void (although this is possible to do, by using tools of linear algebra). But by embedding K in another complex L where |L| is a tetrahedron in \mathbb{R}^3 , we can exploit duality for this purpose (Exercise).

¶28. Final Remarks. Tarjan has shown that the upper bounds obtained for disjoint sets are essentially tight under the pointer model of computation. Ben-Amram and Galil [1] proved a lower bound of $\Omega(m\alpha(m,n))$ lower bound on Union-Find problem under a suitable RAM model of computation. Gabow and Tarjan shows that the problem is linear time in a special case. There are efficient solutions that are not based on compressed trees. For instance, the exercise below shows a simple data structure in which Find requests take constant time while Union requests takes non-constant time. Another variation of this problem is to ensure that each request has a good worst case complexity. See [2, 5].

The application to expression optimization arises in the construction of optimizing compilers in general. In an application to robust geometric computation [4], such optimizations are critical for reducing the potentially huge bit complexity of the numerical computations.

_____Exercises

Exercise 5.1: Hand simulate Kruskal's algorithm on the graph in Figure 13. Specifically:

(a) First show the list of edges, sorted by non-decreasing weight. View vertices v_1, v_2, v_3 , etc as the integers 1, 2, 3, etc. We want you to break ties as follows: assume each edge has the form (i, j) where i < j. When the weights of (i, j) and (i', j') are equal, then assume the weight of (i, j) is less than that of (i', j') if (i, j) is lexicographically less than (i', j'). (b) Maintain the union-find data structure needed to answer the basic question in Kruskal's algorithm (namely, does adding this edge creates a cycle?). The algorithms for the Union and Find must use both the rank heuristic and the path compression heuristic. At each stage of Kruskal's algorithm, when we consider an edge (i, j), we want you to perform the corresponding Find(i), Find(j) and, if necessary, Union(Find(i), Find(j)). You must show the result of each of these operations on the union-find data structure.

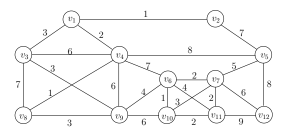


Figure 13: Another House Graph

Exercise 5.2: Let G = (V, E; C) be the usual input for MST. We consider a variant where we are also given a forest $F \subseteq E$. We want to construct a minimum cost spanning tree T of G subject to the requirement that $F \subseteq T$. Call this the **constrained MST** problem. As usual, let us just consider the cost version of this problem, where we only want to compute the minimum cost. Describe and prove the correctness of a Kruskal-like algorithm for this problem. Analyze its complexity. \diamondsuit

Exercise 5.3: Let us define a set $S \subseteq E$ to be **Kruskal-safe** if (i) S is contained in an MST and (ii) for any edge $e \in E \setminus S$, if $C(e) < \max\{C(e') : e' \in S\}$ then $S \cup \{e\}$ contains a cycle. Note that condition (i) is what we called "simply safe" in §V.3. Show that if S is Kruskal-safe and e is an edge of minimum costs among those edges that connect two connected components of S then $S \cup \{e\}$ is Kruskal safe.

 \Diamond

Exercise 5.4: Describe an algorithm to determine the height of every node in a DAG G.

Assume an adjacency list representation of G. Briefly show its correctness and analyze its running time.

- Exercise 5.5: Let K be a simplicial complex and $|K| \subseteq \mathbb{R}^3$.
 - (a) Show that we can embed K in another simplicial complex L such that |L| is a tetrahedron.
 - (b) Suppose we have an incremental algorithm to compute the Betti numbers of L. Show how we can obtain use it to compute the Betti numbers of K.
 - (c) Suppose that the triangles and tetrahedrons (or cells) of L are t_1, \ldots, t_m and c_1, \ldots, c_n . Consider the simplicial complex K_i ($i=0,\ldots,m$) such that K_i has all the vertices and edges of K but only the triangles $\{t_1,\ldots,t_i\}$. (K_i has no tetrahedrons. We want to detect the situation when a transition from K_{i-1} to K_i creates a new void. Consider the dual graph G_i ($i=0,\ldots,m$) whose vertex set is $\{c_1,\ldots,c_n\}$ and whose edges are $\{t_{i+1},\ldots,t_m\}$. NOTE that in G_i , we identify a triangle t with the edge t, t where t is t show that adding t is to t in t creates a new void in t iff the number of connected components increased by one in going from t in t i
 - (d) Show that if we construct the graph G_i 's in reverse order (starting from G_m down to G_0) then we can detect all the *i*'s such that adding t_i creates a new void. Describe this algorithm's implementation using Union-Find.
 - (e) Conclude that we can maintain the Betti numbers of L in time $O(n\alpha(n))$ where n is the number of simplices in L.

END EXERCISES

References

736

737

738

740

741

742

744

746

748

749

750

751

- 755 [1] A. M. Ben-Amram and Z. Galil. Lower bounds of data structure problems on rams. *IEEE Foundations of Comp. Sci.*, 32:622–631, 1991.
- N. Blum. On the single-operation worst-case time complexity of the disjoint set union problem. SIAM J. Computing, 15:1021–1024, 1986.
- Z. Galil and G. F. Italiano. Data structures and algorithms for disjoint set union problems.
 ACM Computing Surveys, 23(3):319–344, 1991.
- [4] C. Li and C. Yap. Recent progress in Exact Geometric Computation, 2001.
- [5] M. H. M. Smid. A datastructure for the union-find problem having a good single-operation complexity. Algorithms Review, 1(1):1–12, 1990.
- ₇₆₄ [6] R. E. Tarjan. Data Structures and Network Algorithms. SIAM, Philadelphia, PA, 1974.
- ⁷⁶⁵ [7] R. E. Tarjan. Efficiency of a good but not linear set union algorithm. *J. ACM*, 22:215–225, 1975.
- $_{767}$ [8] R. E. Tarjan and J. van Leeuwen. Worst-case analysis of set union algorithms. *J. ACM*, $_{768}$ 31:245–281, 1984.