

Triangulating a Simple Polygon in Linear Time*

Bernard Chazelle

Department of Computer Science, Princeton University
Princeton, NJ 08544, USA

Abstract

We give a deterministic algorithm for triangulating a simple polygon in linear time. The basic strategy is to build a coarse approximation of a triangulation in a bottom-up phase and then use the information computed along the way to refine the triangulation in a top-down phase. The main tools used are the polygon-cutting theorem, which provides us with a balancing scheme, and the planar separator theorem, whose role is essential in the discovery of new diagonals. Only elementary data structures are required by the algorithm. In particular, no dynamic search trees, finger trees, or point-location structures are needed. We mention a few applications of our algorithm.

1 Introduction

Triangulating a simple polygon has been one of the most outstanding open problems in two-dimensional computational geometry. It is a basic primitive in computer graphics and, generally, seems the natural preprocessing step for most nontrivial operations on simple polygons [5], [14]. Recall that to triangulate a polygon is to subdivide it into triangles without adding any new vertices. Despite its apparent simplicity, however, the triangulation problem has remained elusive. In 1978 Garey et al. [12] gave an $O(n \log n)$ -time algorithm for triangulating a simple n -gon. While it was widely believed that triangulating should be easier than sorting, no proof was to be found until 1986, when Tarjan and Van Wyk [27] discovered an $O(n \log \log n)$ -time algorithm. Following this breakthrough, Clarkson et al. [7] discovered a Las Vegas algorithm, recently simplified by Seidel [25], with $O(n \log^* n)$ expected time. In 1989 Kirkpatrick et al. [20] gave a new, conceptually simpler $O(n \log \log n)$ -time algorithm, and they also derived an $O(n \log^* n)$ bound for the case where vertices have polynomially bounded integer coordinates. Other results on the triangulation problem include linear or quasi-linear algorithms for restricted classes of polygons [6], [11], [17], [28]–[30].

Our main result is a linear-time deterministic algorithm for triangulating a simple polygon. The algorithm is elementary in that it does not require the use of any complicated data structure; in particular, it does not need dynamic search trees, finger trees, or any fancy point-location structures.

What makes fast polygon triangulation a difficult problem are the basic inadequacies of either a pure top-down or a pure bottom-up approach. To proceed top-down is to look at the whole polygon and infer global information right away. We can rely on the polygon-cutting theorem [4] which says that the polygon can be cut along a diagonal into two roughly equal-size pieces. The immediate dilemma is that to find such a diagonal appears just as difficult as triangulating the whole polygon

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to begin with. Besides, we would actually need to find such a diagonal in sublinear amortized time (say, bounded by $O(n/\log^2 n)$) to keep our hopes for an optimal triangulation algorithm alive. A bottom-up approach, on the other hand, involves computing, say, triangulations of subpieces of the polygon’s boundary. This suffers from the obvious flaw that too much information gets to be computed. Indeed, diagonals for small pieces of the boundary are not guaranteed to be diagonals of the whole polygon and might therefore be wasted. Our solution is to mix bottom-up and top-down approaches together. The basic strategy is to build a coarse approximation of a triangulation in a bottom-up phase and then use the information computed along the way to refine the triangulation in a top-down phase. The main tools used are

- (i) the polygon-cutting theorem, which provides us with a balancing scheme, and
- (ii) the planar separator theorem [21], whose role is essential in the discovery of new diagonals.

Here is a more detailed overview of the algorithm. As was observed in [6] and [11] a triangulation of a polygon can be derived in linear time from its horizontal visibility map, sometimes referred to in the literature as trapezoidal decomposition: this is the partition of the polygon obtained by drawing horizontal chords from the vertices. We can extend this notion easily and speak, more generally, of the visibility map of any simple polygonal curve. Chazelle and Incerpi [6] showed how to build the visibility map of an n -vertex curve in $O(n \log n)$ time, using divide-and-conquer. Their algorithm mimics mergesort: assuming that n is a power of 2, at the k th stage ($k = 1, 2, \dots, \log n$), the boundary of the polygon is decomposed into chains of size 2^k , whose visibility maps are computed by piecing together the maps obtained at the previous stage. Each stage can be accomplished in a linear number of operations, so computing the visibility map of the polygon takes $O(n \log n)$ time.

The new algorithm follows the same pattern: it goes through an up-phase of $\log n$ stages, each involving the merging of maps obtained at the previous stage. The novelty we bring into this process is to use only coarse samples of the visibility maps during the merges. In this way we can carry out an entire stage in sublinear time and beat the $n \log n$ barrier. The samples are uniform submaps of the visibility maps; uniform in the sense that they approximate the visibility maps anywhere equally well. Of course, in the end, we also need an efficient way to refine the submap derived for the whole polygon into its full-fledged visibility map. After this is done, it takes only linear time to compute a triangulation. To refine a submap we go down through stages in reverse (a down-phase): each transition refines the submap incrementally, until we get back to the first stage, at which point the full visibility map emerges at last. Figure 1.1 illustrates the meaning of the up and down-phases.

Perhaps now is the right time to wonder whether our approach is not inherently flawed from the start. How sound is it to mimic mergesort when our goal is to beat $n \log n$? Any attempt to speed up mergesort by using “coarse samples” of the lists to be merged is trivially doomed. So, what is so different about polygons? The difference is rooted in a notion which we call *conformality*. This is perhaps the single most important concept in our algorithm, for it is precisely where mergesort and triangulation part ways. Recall that the polygon-cutting theorem is a geometric analog of the centroid theorem for free trees: a visibility map has a tree structure and, so, can be written as a collection of “blobs” of roughly equal size, themselves interconnected in a tree pattern. These blobs are the constituents of a submap. Merging two submaps can thus be equated with “merging” two trees together. The mergesort equivalent of a submap would be a sublist (of one of the lists to be merged) obtained by picking keys at regular intervals. Notice that merging two such sublists might in the worst case produce a new sublist whose corresponding intervals are up to twice the size of the original intervals. This coarsening effect prevents us from speeding up mergesort, because repairing the damage might involve computing medians or things of that nature for which no shortcuts can

be found. To be sure, as we shall see, equally bad things can happen with submaps; repairing the damage, however, can be done by simply adding new chords to submaps, which can be made to take only sublinear time. To make this possible we must keep the coarseness of submaps under control by requiring that the tree structure of a submap be of bounded degree: in our terminology “conformal” actually means degree at most 4. Restoring conformality after merging two submaps is the linchpin of the algorithm and, as we should expect, its most delicate and subtle part as well: it can be viewed as a geometric “two-dimensional” analog of rotations in balanced dynamic search trees.

Although our algorithm is an outgrowth of the mergesort-like method of Chazelle and Incerpi, it goes far beyond it. For example, the algorithm never actually merges visibility maps but only submaps (which is done quite differently). We must also borrow ideas from a number of other sources: as we mentioned before, one of them is the polygon-cutting theorem, and, more specifically, the hierarchical polygon decomposition of Chazelle and Guibas [5]. There exist optimal algorithms for computing such decompositions [3], [14] but, as it turns out, standard suboptimal methods work just as well for our purposes. Merging submaps requires a primitive to detect new chords. In the Chazelle-Incerpi method the detection is limited to constant-size domains, so it can be done naively. But here the domains can be arbitrarily large so we need a sublinear ray-shooting method. This cannot be done by using fast planar point location, which is the approach followed in the algorithm of Kirkpatrick et al. [20]. The reason is that we need to support merging of two point-location structures, and the known methods, even the dynamic ones, are inadequate in that regard. We turn the problem around by using a weak form of divide-and-conquer based on Lipton and Tarjan’s planar separator theorem [21].

2 Visibility Maps and Submaps

Following the tradition of visibility algorithms, we begin by restricting their applicability to polygons where no two distinct vertices have the same y -coordinate. Of course, the standard excuse still works: we can easily get around this assumption by applying the symbolic perturbation techniques of [10] and [31]. Turning now to visibility maps, recall that, in the Chazelle-Incerpi method, chords are extended only toward the interior of the polygon, with a special rule to determine how far they should extend. Kirkpatrick et al. [20] use the simpler scheme of extending chords on both sides. This amounts to thinking of a polygonal curve as a very thin polygon embedded in a cylindrical plane that lets chords wrap around infinity. For the reasons we give below, we find it more convenient to embed our objects in a topological manifold, called the *spherical plane*, which is equivalent to a 2-sphere. Proofs of correctness in the polygonal merging business tend to be tricky and ridden with painful case-analyses. One reason for this is that these proofs often attempt to establish topological facts by geometric means, thus adding unnecessary complication. By sticking to topological considerations as much as possible, proofs become much simpler, provided, of course, that the ambient space has the “right” topology.

What is the right topology in this context? One problem with the cylindrical plane is that although it has a Jordan curve theorem, the two regions created by removing a simple closed curve may now have one of three types:

- (i) an open disk,
- (ii) a cylinder $S^1 \times (0, +\infty)$, or
- (iii) a perforated cylinder.

We simplify all that by defining our ambient space to be the spherical plane: this is the product space $[-\infty, +\infty]^2$ with the following identification rules:

- (i) $(-\infty, y) = (+\infty, y)$
- (ii) $(x, -\infty) = (-x, -\infty)$
- (iii) $(x, +\infty) = (-x, +\infty)$

for all $x, y \in [-\infty, +\infty]$. The spherical plane is homeomorphic to a 2-sphere, so we now have the nicest Jordan curve theorem of all: the removal of any simple closed curve creates two open disks.

The only remaining difficulty is the orientability of Jordan curves. If we fix an orientation of the spherical plane, the boundary of any region homeomorphic to a disk, being of codimension 1, has an induced orientation, called *clockwise*; the reverse orientation is called *councclockwise*. Conversely, given a Jordan curve, a clockwise tour of the curve refers to the orientation induced by one of the two homeomorphic disks that it bounds. Unless one of these disks is already understood, we choose one unambiguously by fixing a reference point somewhere in the plane and looking at the unique disk that does not contain it. Of course, this means restricting ourselves to curves that avoid that point, but this is only a minor inconvenience. From now on, we assume that the reference point is at $(0, +\infty)$. In this way, to speak of the clockwise traversal of a Jordan curve, without reference to an enclosed region, makes full sense.

2.1 The Visibility Map

Given a simple (nonclosed) polygonal curve C with vertices v_1, \dots, v_n , we define the visibility map of C , denoted $V(C)$, as the planar subdivision formed by extending two horizontal segments from each vertex v_i , one in each direction. Each segment, which we call a *chord*, is extended until it meets another point of C . If it were to go to infinity in the Cartesian plane, then it would actually wrap around in the spherical plane until it hits C again. Adding chords to C subdivides the spherical plane into regions: triangles or trapezoids with two horizontal segments and two nonhorizontal segments, all of which are possibly divided into several collinear edges (Fig. 2.1).

In order to distinguish between the two sides of an edge, we give each edge of C an infinitesimal width so as to make the curve C into a very thin simple polygon. (This is just a conceptual device, so, in particular, no actual perturbation of the polygon needs to be performed on the computer.) The boundary of that polygon is called the *double boundary* of C and is denoted ∂C . By abuse of terminology we refer to the two sides of the double boundary as one would speak of the left and right sides of a snake; of course, this is meaningful geometrically but not topologically. Each vertex of C that is not a local extremum in the y -direction gives rise to two companion vertices in ∂C , one on each side of the curve (Fig. 2.2.1). In this way, each vertex is incident upon exactly one chord (possibly the same chord for the two companions). But what about local extrema? For each such vertex of C , if it is not one of the two endpoints, we create a total of four vertices in ∂C (Fig. 2.2.2): two companion pairs of duplicate vertices; one pair on each side of C . The duplicate vertices in a given pair are next to each other along ∂C . By convention, we say that one of these pairs, the one on the “inside” of the turn, gives rise to a chord of null length. Finally, as shown in Fig. 2.2.3, for each endpoint of C we create two companion vertices; these can also be called duplicates since they are next to each other as well as on both sides of C . Figure 2.3 illustrates these definitions. Note that for simplicity we have not numbered vertices connected to null-length chords, but we have numbered all chord endpoints for later use. We have now ensured that each vertex of ∂C is incident upon exactly one chord of the visibility map. Therefore, any vertex and

actually also any point of ∂C has a unique horizontal “chord direction” (left or right) assigned to it. Roughly speaking, this direction points to the left of an observer walking clockwise around ∂C .

If a chord of $V(C)$ has p and q as endpoints, we say that p and q *see each other* with respect to C , or simply, see each other, when C is understood. Equivalently, we say that p and q are *mutually visible*. More generally, if p and q are two points (not necessarily vertices) of ∂C with the same y -coordinates, we say that p and q see each other with respect to C if one of the two (relatively open) segments joining p and q lies completely outside of C (regarded as a thin simple polygon). By extension the segment pq is also called a chord. For example, in Fig. 2.3 points labeled 6 and 7 see each other, whereas points 6 and 14 as well as points 3 and 4 and points 1 and 1' do not see each other. We close our discussion of visibility with a simple but useful fact.

Lemma 2.1. *If we remove a pair of mutually visible points from the double boundary of a simple polygonal curve, then no chord can connect the two resulting pieces.*

Proof. Let C_1 and C_2 be the two pieces of the double boundary resulting from the removal of a pair of visible points. Together with C_1 and C_2 , the chord c connecting the two points subdivides the spherical plane into three polygonal regions (i.e., regions bounded by simple polygonal curves), one of which is the infinitesimally thin polygon C itself. Any chord connecting C_1 and C_2 lies outside the “polygon” C , so it must cross c . But this is impossible because chords are horizontal. \square

The circular sequence of chord endpoints in $V(C)$ encountered during a clockwise traversal of ∂C is called the *canonical vertex enumeration* of $V(C)$: note that it contains other points besides the vertices of C . Figure 2.3 provides the sequence 1, 1', ..., 17, 18, with the primes indicating duplicated vertices. Recall that we have refrained from numbering the endpoints of null-length chords. Speaking of which, note that null-length chords create empty regions. The traversal of the double boundary leads to a canonical enumeration of the regions (with repetitions). In the case of Fig. 2.3 we have the list (including only the nonempty regions for simplicity)

(I, II, III, IV, V, VI, VII, VIII, VII, IX, VII, VI, V, IV, III, X, XI, X, III, II, I, XII, I, XIII).

It is easy to see that the dual graph of the subdivision is a tree, naturally called the *visibility tree* of C . This graph is defined by associating a distinct node with each region (empty or nonempty) of the visibility map and connecting any pair of nodes whose corresponding regions share a common chord. Note that this also includes null-length chords. Figure 2.1 shows the tree without the nodes associated with empty regions. Since any two consecutive regions in the canonical region enumeration have a common chord, the visibility tree is indeed connected. Why can it not have cycles? It suffices to show that removing any chord ab would disconnect the graph.

First, assume that a and b are not duplicates of each other (although they might be companions). Then, from Lemma 2.1, removing a and b splits the double boundary into two pieces which are closed under visibility. It follows that the boundary of any region contains segments from exactly one of the two pieces, and therefore can be classified by the piece to which these segments belong. Any chord separating a region associated with one piece from a region associated with the other piece must join two points at the juncture between the two pieces, and only ab satisfies this requirement. This concludes the first case. Assume now that a and b are duplicates of each other. If the chord ab has zero length, then removing it isolates an empty region from the rest, and our claim holds. So, suppose that ab has nonzero length. Then a is either the highest or lowest point of C , therefore removing ab would disconnect the upper or lower part of the spherical plane from the other regions. This completes the proof that the visibility tree is indeed a tree. (There is also a simple topological proof, which is given in Lemma 2.2 below.)

2.2 Visibility Submaps

An operation which we will find useful is the removal of chords from $V(C)$. Before we go any further, however, we wish to prepare for the possibility that $V(C)$ has been augmented with some additional chords (something that will often happen later). Obviously, these new chords cannot connect vertices of ∂C (since all have been used up) but rather arbitrary points on the curve. Although this clearly affects $V(C)$ as well as the visibility tree, everything we said previously regarding canonical enumerations can be trivially extended to this new situation. So, from now on, let us treat $V(C)$ either in its original state or in some augmented form. We specify which applies whenever the distinction needs to be made.

The operation we want to discuss now involves removing a given chord from $V(C)$. Since $V(C)$ may have additional chords, we need to be careful about the meaning of a removal. A chord has two endpoints; none, one, or two of which are vertices of C . So, the removal of a chord entails removing not only the chord itself but also those endpoints that are not vertices of C , and glueing back ∂C at those points. This cleanup operation is meant to prevent the presence of vertices stranded in the middle of an edge of ∂C : in other words, any vertex that is not a vertex of C must be the endpoint of a nonremoved chord.

Removing one or several chords (of zero or nonzero length) from a map produces a polygonal subdivision of the spherical plane, called a *submap* of $V(C)$ (Fig. 2.4). The boundary of a nonempty region of the submap is an oriented circular sequence of horizontal segments, called *exit chords*, alternating with pieces of C and not of C . For example, let us assume that all null-length chords have been removed in the submap of Fig. 2.4. Then the boundary of the region labeled II consists of a two-edge arc (beginning at the big dot), followed by an exit chord, a one-edge arc, an exit chord, a three-edge arc (not a four-edge arc!), an exit chord, a one-edge arc, and one final exit chord. Note that some arcs may be of zero length, as is the case in the region labeled V. As illustrated in Fig. 2.4, a clockwise traversal of ∂C induces a traversal of the boundary of each region of the submap which is counterclockwise with respect to the orientation of the region. More formally, we have the following fact.

Lemma 2.2. *Let A_1, \dots, A_k be the counterclockwise enumeration of the (oriented) arcs of a nonempty submap region (as induced from the region's orientation). Then each A_i is oriented clockwise with respect to C . Moreover, the sequence A_1, \dots, A_k also occurs clockwise around C .*

Proof. The curve ∂C is homeomorphic to a circle embedded in the spherical plane. Adding a chord is topologically equivalent to connecting two points on the circle by a simple curve lying entirely on one side of the circle. The requirement that all these curves should be mutually disjoint induces a parenthesis system which immediately reveals the tree structure of the dual graph. This is similar to the parenthesis systems in Jordan sorting [18]. As an example, Fig. 2.5 depicts the topological equivalent of the submap of Fig. 2.4. From this perspective, the lemma should be completely obvious. \square

As was the case with $V(C)$, a clockwise traversal of ∂C induces canonical vertex/region enumerations of the submap. Figure 2.4 gives the region enumeration: I, II, III, II, IV, II, V, II, I. (Recall that that particular submap is supposed to have had all its null-length chords removed and therefore has no empty regions.) Bold dots mark the points during the clockwise traversal of the double boundary where the canonically enumerated regions are first discovered. An important requirement is that a vertex enumeration of a submap should list only the endpoints of actual exit chords and thus might skip over many vertices of C . In this way, canonical enumerations of any

type take time proportional to the number of regions and not to the number of vertices (which might be much higher). We define the *weight* of a region as 0 if the region is empty, or else as the maximum number of nonnull length edges in any of its arcs. For example, regions I and II in Fig. 2.4 have weights 4 and 3, respectively.

Although weights count only edges of nonzero length, chords of zero length (if any) are taken into account inasmuch as they separate arcs. In other words, an arc never contains any chord, whether that chord be of nonzero length or not. Of course, once removed, a chord of zero length ceases to separate any arcs. Also, note the important role played by the double boundary in the definition of a region's weight. Indeed, a region may have a very small weight because its arcs are all small; but this does not prevent any one of these arcs from having a huge number of vertices on the other side of the double boundary. Of course, because vertices of C have companions those extra vertices have to be endpoints of exit chords.

Combinatorially, a region corresponds to a subtree of the visibility tree of C . The dual graph of a submap is obtained by contracting the edges of the visibility tree that correspond to the removed chords (Fig. 2.6). Being derived from the visibility tree by graph-minor operations, the dual graph of a submap is itself a tree (as was clear in the proof of Lemma 2.2), which we simply call the *tree of the submap*. Note that, conversely, contracting any edge of the visibility tree amounts to removing the corresponding chord from the visibility map. The weight of a node naturally refers to the weight of its corresponding region.

2.3 Conformality and Granularity

Since no two distinct vertices of C have the same y -coordinate, the degree of any node in a visibility tree cannot exceed 4. We should not expect this to be always true of submap trees, however, so we distinguish the trees of *conformal* submaps as those with node-degree at most 4. By analogy with the polygon-cutting theorem [4] we can decompose a conformal submap in an hierarchical manner. The idea is to pick the centroid of the submap's tree and observe that there exists at least one incident edge whose removal leaves two subtrees, each with a number of edges at most three-quarters the original number. Associating the removed edge with the root of a binary tree and recursing in this fashion with respect to the root's two children provides a tree decomposition of the submap. The tree has depth logarithmic in the number of regions (which is the number of chords plus one). The internal nodes (resp. leaves) are in bijection with the exit chords (resp. regions) of the submap. Figure 2.7 illustrates the correspondence (leaves have been omitted). By using straightforward tree-labeling techniques we can find the centroid node, and from there, the first edge to be removed, in linear time. Proceeding recursively gives us a simple $O(m \log m + 1)$ -time algorithm for computing the tree decomposition of a submap of m regions. We use this result below because it is simple and practical. Optimal methods exist but they are all fairly complicated and unnecessary for our purposes [3], [14].

One final piece of classification addresses our desire to make submaps coarse but uniform approximations of visibility maps. We say that a submap is γ -*granular* if

- (i) every node of its tree has weight at most γ and
- (ii) contracting any edge incident upon at least one node of degree less than 3 produces a new node whose weight exceeds γ .

Note that this weight might be less than the added weight of the two nodes of the contracted edge. This is either because the arcs incident upon the chord removed did not determine the weights, or more interestingly, because one or both endpoints of the chord might not be vertices of ∂C and

might thus disappear. If only condition (i) holds, then the submap is γ -semigranular. Adding condition (ii) makes the semigranularity maximal in some sense. Finally, by default, if (i) holds but the submap has no exit chord, it is still said to be γ -granular. The following result asserts that, as we would expect, a γ -granular conformal submap is more economical to encode than its full visibility map by a factor of γ .

Lemma 2.3. *If C is a polygonal curve with n vertices, any γ -granular conformal submap of the (possibly augmented) visibility map of C has $O(n/\gamma + 1)$ regions and each region is bounded by $O(\gamma)$ edges.*

Proof. We can assume that the tree of the submap has at least one edge, otherwise the lemma is trivial because of the γ -granularity. Among the edges of that tree, let E be the set of those incident upon at least one node of degree less than 3. It is trivial to show by induction on the size of the tree that E accounts for at least a fixed fraction of all the edges. Now, contracting any edge in E , or equivalently, removing a chord associated with E produces a merged region of weight greater than γ , meaning that it has an arc with more than γ edges of nonzero length. Since a vertex of C can give rise to at most four vertices of ∂C , and removed chords do not leave extra vertices behind except those of ∂C , such an arc must involve at least $\Omega(\gamma)$ distinct vertices of C . If contracting any edge of E were always to produce a disjoint merged region, then it would follow from the pigeon-hole principle that E , and hence the whole tree, has $O(n/\gamma + 1)$ edges. Unfortunately, two edges of E might produce overlapping merged regions (i.e., if they share a common node). From the conformality of the submap, however, we know that a given vertex of C can be used at most a constant number of times in this counting argument, therefore E has indeed $O(n/\gamma + 1)$ edges and the first part of the lemma is established. The second part derives from the conformality of the submap, which ensures that there is a bounded number of arcs per region and, hence, that the total number of bounding edges is at most proportional to the weight of the region. \square

2.4 Representation Issues

How do we represent visibility maps and submaps as data structures? We first describe our mode of representation, then we point out some of its idiosyncracies and explain why they are needed. Let P be the input polygonal curve (the one whose visibility map is sought) and let C be the subchain of P whose visibility map (or submap) we wish to represent. We assume that P is nonclosed; this is not restrictive since a little hole can always be punctured if it is closed to begin with. We assume that the edges of P are stored in a table (the *input table*) in the order in which they occur along the boundary of P . (A doubly linked list would also do.) Note that the notion of double boundary need not be encoded explicitly, i.e., no edges are duplicated in the table. The input table is read-only: it is never to be modified or even copied. A visibility submap of $V(C)$ is represented by its own data structure: arcs are encoded by pointing directly into the input table. More precisely, each arc is represented by a separate *arc-structure*. Null-length arcs can be represented explicitly so let us assume that the arc has nonzero length. Let e_1, \dots, e_t be the edges of an arc in clockwise order along the double boundary, where e_1 and e_t are the edges adjacent to the two chords connected by the arc. If $t = 1$, then the arc-structure consists of a single pointer into the input table to the edge e of P that contains e_1 . Since e_1 is an edge of the double boundary, we also need to indicate by a flag which side of e is to be understood. We do not need to record the endpoints of the arc because chords take care of that. If $t > 1$, we store the same information as above but now with respect to both e_1 and e_t in that order. We say that a submap (or map) is given in *normal form* if the following information is provided:

- (i) The tree of the submap (or map) is represented in standard edge/node adjacency fashion.
- (ii) Each edge of the tree contains a record describing the corresponding chord as well as pointers to the arc-structures of the two, three, or four arcs adjacent to it. Conversely, each arc-structure has a pointer to the node of the tree whose corresponding region is incident upon the arc in question.
- (iii) The arc-structures are stored in a table (the *arc-sequence table*) in the order corresponding to a canonical traversal of the double boundary ∂C . Also, the endpoints of C are identified by appropriate pointers into the input table as well as by pointers to the arc-structures whose corresponding arcs pass through the endpoints.
- (iv) If the submap is conformal, then its tree decomposition should be available.

We choose what may seem to be a contrived representation of a submap in order to use storage proportional not to the number of edges in the submap but rather to the number of regions (which is of the same order of magnitude as the number of chords and arcs). It is essential to avoid excessive duplication of information because we need to encode a collection of submaps whose number of distinct features is only $O(n)$, but whose combined size, counting redundancy, is $\Theta(n \log n)$. Note that our representation is powerful enough to let us perform canonical vertex/region enumerations in optimal time. If we wish to, we can also enumerate all the vertices of ∂C in clockwise order directly from a canonical vertex enumeration of the submap, since any arc can be reconstructed explicitly from the succinct information given by the arc-structure: it suffices to explore the input table between the locations indicated by the two pointers of the arc-structure. Note that caution must be used since an arc might wrap around both sides of C , something we call *double-backing*. This can be detected when we traverse the arc as soon as we reach an edge of P incident upon an endpoint of C .

Perhaps a less obvious task is to retrieve the arc-structure corresponding to an arc, given one of its edges. More specifically, suppose that we are given an edge e of C and a point q on it. The question is to find the arc-structures of all those arcs in the submap that pass through the point q . By passing through, we do not care whether the arc is on any particular side of the double boundary, so, for example, if q is not an endpoint of any chord in the submap, then there are at most two distinct arcs to be found. Otherwise, there are at most six of them, two of which are of zero length: this worst case occurs when q coincides with a vertex of C that is a local extremum in the y -direction. Since we know the location of the two endpoints of C in the arc-sequence table (i.e., which arcs pass through them) we can conceptually break up the circular arc sequence into two linear sequences and perform in each of them a binary search, using the name of the containing edge e as a query. Either search might take us to a unique arc-structure, in which case we are done, or else to a contiguous interval of arc-structures: this might happen if e contributes several arcs. We can disambiguate by pursuing the binary search, now using, say, the y -coordinate of q as a query. The total running time is logarithmic in the number of arcs. This operation is very useful later when we want to navigate in a submap across its arcs: we call it the *double identification* of a point of ∂C .

We have said repeatedly that a submap has a tree structure. Now let us change our perspective for a moment and look at a submap as a standard planar subdivision, without distinguishing between chords and arc edges. There are many standard representations of planar graphs [2], [15], [23] which allow us to navigate through such a subdivision along the edges in constant time per step taken. Normal-form representations are not quite that powerful. One problem arises if we attempt to cross from one side of an arc to the other along, say, a straight line. In order to find

which region we are about to enter we must perform a double identification. The difficulty here is that unlike what is commonly done in standard graph representations we do not keep adjacency information between regions and edges (except for chords). More important yet, we do not provide an explicit correspondence between the features on the two sides of an edge of C . For reasons which will become clear later, it would be a very bad idea to try to do so.

2.5 A Topological Lemma

We close this discussion of visibility submaps by proving a result on the topology of regions and chords, which we use to establish the correctness of our submap merging algorithm. Let D be a closed disk in the spherical plane, let ∂D be its boundary, and let ab denote a diametrical chord of D . Pick two distinct points c, d on ∂D such that a, c, b occur in clockwise order (with respect to D), and let A be a simple curve lying inside D and running from c to d . Consider the circular arc that runs clockwise from d to c and let B_i ($i = 1, 2$) be the closures of its intersections with the two circular arcs of $\partial D \setminus \{a, b\}$ (Fig. 2.8). Note that each B_i consists of 0, 1, or 2 circular arcs. We say that a subset α of A is *shielded* from B_j if either B_j is empty or else no point of α can be connected to any point of B_j by a curve (understood here as a closed set) that lies entirely inside D and does not intersect either ab or A . In Fig. 2.8.1, for example, the piece of A running from c to a' is shielded from B_2 since none of its points can be connected to B_2 without crossing ab . Similarly, the piece from a' to b' is shielded from B_1 since a connection to it would have to cross ab or A .

Lemma 2.4. *If $a \in B_1 \cup B_2$ (resp. $b \in B_1 \cup B_2$), let a' (resp. b') be the first (resp. last) point of $ab \cap A$ encountered when traversing the diametrical chord ab from a to b . The points a' and b' (which might not exist) subdivide A into a total of one, two, or three connected curves, each of which is shielded from some B_j (not necessarily the same j for all curves). Furthermore, an appropriate B_j can be identified simply on the basis of a, b, c, d, a', b' .*

Proof. We can assume that both B_1 and B_2 are nonempty and that A intersects ab (else the lemma is trivially correct). By attaching $B_1 \cup B_2$ to A , we obtain a simple closed curve within D , which is, therefore, the boundary of a subset R of D homeomorphic to a disk. If a (resp. b) belongs to $B_1 \cup B_2$, then the segment aa' (resp. bb') lies within R and thus, acting like a chord, subdivides R into two regions. Since aa' and bb' cannot intersect, together they subdivide R into two or three disk-like regions. The boundary of each such region intersects the boundary of D in a single connected arc and therefore avoids one B_i (outside of a and b) entirely. Figure 2.8 illustrates the two possible cases; note that the third case, where the counterclockwise traversal from c to d avoids both a and b , was eliminated earlier, since it corresponds to a situation where one of the B_i is empty. None of the curves obtained by removing a' and b' (if they exist) from A can belong to more than one of the subdividing regions, so each of them is shielded from some B_i . Which one can be determined by simple examination of the relative order of the points a, b, c, d, a', b' around the boundary of R . \square

3 Merging Two Submaps

The inner loop of the visibility algorithm involves merging two conformal submaps. Everything else in the algorithm is part of a control mechanism for deciding what gets to be merged with what, at what time, and with what desired granularity. Let C_1 and C_2 be two polygonal curves of n_1 and n_2

vertices respectively, whose union C forms a connected vertex-to-vertex piece of the input (simple and nonclosed) polygonal curve P ; we assume that $C_1 \cap C_2$ is a vertex of P . Let S_i ($i = 1, 2$) be a γ_i -granular conformal submap of $V(C_i)$, with $\gamma_1 \leq \gamma_2$. Given any integer $\gamma \geq \gamma_2$, to merge S_1 and S_2 (where γ is understood) means to compute a normal-form γ -granular conformal submap of $V(C)$.

To facilitate the exposition we assume that we have at our disposal two primitives: one is a *ray-shooting oracle*, which allows us to shoot a horizontal ray toward any subarc of S_1 or S_2 and discover which point, if any, is first hit by the ray; this gives us a way to discover new chords. The other primitive is an *arc-cutting oracle*, which enables us to cut up any subarc into a small number of pieces for which conformal submaps of the appropriate granularity have already been computed. This is to be used for restoring the conformality of merged submaps.

The merge proceeds in three stages. First we find which points of ∂C can be seen by the endpoints of the exit chords of S_i ($i = 1, 2$) and by the companion vertices resulting from the duplication of $C_1 \cap C_2$; this gives us chords which we use to create a submap S of $V(C)$ called the *fusion* of S_1 and S_2 (Section 3.1). In the second stage we ensure that the submap is conformal, which might involve adding new chords to cut up regions with more than four arcs. This is done by calling upon the arc-cutting oracle, which allows us to deal with subarcs for which conformal submaps and their tree decompositions are available. Finding new chords to cut up big regions is carried out by binary search through the appropriate tree decompositions, using the ray-shooting oracles along the way (Section 3.2). In the third stage, finally, we bring the submap S to the desired granularity by removing chords if necessary (Section 3.3). The implementation of the oracles is discussed in Section 3.4.

We need to be able to distinguish between an arc of ∂C and the piece of C from which it originates. For this purpose we introduce the notation $\bar{\alpha}$ to refer to the portion of C to which an arc α of ∂C corresponds. Recall that an arc may double-back around an endpoint of C , so $\bar{\alpha}$ may not always be as “long” as α . We assume that each S_i is given in normal form and that the following set of primitives is available. For each region arc α of S_i ($i = 1, 2$) specified by a pointer to its arc-structure:

- (i) There exists a ray-shooter which, given any point p along with a horizontal direction (left or right) and any subarc α' of α specified by its two endpoints (along with two pointers to the input table to indicate the names of the edges of P that contain these two endpoints as well as two flags indicating which side of ∂P is to be understood), reports the single point of α' (if any) that a ray of light shot from p in the given direction would hit in the absence of any obstacle except α' . In addition to the point hit, the report should also include the name of the edge of P that contains it. The report should take $O(f(\gamma_i))$ time, where f is a nondecreasing function.
- (ii) There exists an arc-cutter which, in $O(g(\gamma_i))$ time, subdivides the subarc α' into at most $g(\gamma_i)$ subarcs $\alpha_1, \alpha_2, \dots$, such that (1) each α_j is specified by its two endpoints and a pair of pointers into the input table to indicate which edges of P contain these endpoints; the pair should be ordered to reflect a clockwise traversal along ∂P and two flags should be included to indicate on which sides of ∂P these endpoints fall; (2) the relative interior of no $\bar{\alpha}_j$ should contain a point of ∂C_i that corresponds to an endpoint of C_i , that is, each subarc must lie entirely on one side of C (no double-backing); and (3) except for $\bar{\alpha}_1$ and $\bar{\alpha}_2$ (in the case where these are single-edge pieces attached to the points of C corresponding to the endpoints of α'), all the $\bar{\alpha}_j$ are vertex-to-vertex subchains of C_i (i.e., they do not stop in the middle of an edge) and, for each of them, an $h(\gamma_i)$ -granular conformal submap of $V(\bar{\alpha}_j)$ is available in normal form. Again, g and h are assumed to be nondecreasing functions.

Given these oracles we show how to merge S_1 and S_2 in

$$O((n_1/\gamma_1 + n_2/\gamma_2 + 1)f(\gamma_2)g(\gamma_2)(h(\gamma_2) + \log(n_1 + n_2)))$$

time. Ideally, we would like the extra factor $f(\gamma_2)g(\gamma_2)(h(\gamma_2) + \log(n_1 + n_2))$ to be constant. This would mean that the merge could be done in time proportional not to the total size of the submaps but to the number of chords in them. We cannot achieve this, but we can find functions f, g, h which, although nonconstant, are small enough for our purposes. Specifically, we have $f(x) = O(x^{0.74})$, $g(x) = O(\log x)$, and $h(x) = O(x^{0.20})$. This allows us to carry out a merge in time almost proportional to the total number of chords. Note that to achieve $f(x) = g(x) = x$ is trivial but, for our purposes, completely useless.

3.1 Fusion of Two Submaps

By symmetry, we may limit our discussion to the problem of fusing S_1 into S_2 , that is, determining the points of ∂C that are seen by the endpoints of the exit chords of S_1 and by the companion vertices resulting from the duplication of $C_1 \cap C_2$. The idea is then to repeat the work described below with respect to S_2 (i.e., fusing S_2 into S_1), and set up a new submap S based on the information collected. Let a_{m+1} and a_0 be the companion vertices, as they appear next to each other clockwise around ∂C_1 , resulting from the duplication of the vertex $C_1 \cap C_2$ in ∂C_1 . Let a_1, a_2, \dots, a_m be the canonical vertex enumeration of S_1 . Recall that this enumerates the exit chord endpoints in S_1 as we encounter them going clockwise around ∂C_1 . Since the sequence is circular we can assume that a_0 precedes a_1 and a_{m+1} follows a_m . Note that it could happen that a_0 and a_{m+1} are already part of the sequence, but this need not be the case because of chord removals which might have occurred during previous merges, so, for the sake of generality, we assume that they are not and therefore add them to the sequence. A clockwise tour around ∂C_1 that begins at a_0 thus ends at a_{m+1} . We compute the points of ∂C seen by a_0, a_1, \dots, a_{m+1} in that order.

We begin with a simple observation. Given any point p of ∂C_i and the arc to which it belongs (specified by its arc-structure), we can determine which point of ∂C_i it sees (with respect to C_i) in $O(f(\gamma_i))$ time. This operation is called *local shooting*. Recall that because of the double boundary the shooting direction is always uniquely defined. If p is an endpoint of an exit chord we can easily do that (even in constant time). If not, then p belongs to a unique region of S_i , which we can determine in constant time (i.e., via its node in the submap tree), and the point of ∂C_i that it sees lies on one of the region's arcs. This is because regions are closed under visibility, which is a corollary of Lemma 2.1. Using the appropriate ray-shooters, we can find that point by checking each arc in turn and finding the nearest hit. The claim on the time follows from the conformality of S_i , which ensures that at most four arcs need to be checked. Note that local shooting is still possible even if p does not lie on ∂C_i ; it can lie anywhere in the spherical plane as long as a horizontal direction (left or right) has been specified and we know in which region of S_i the point lies. We still call this operation local shooting.

To fuse S_1 into S_2 we let a variable p run through ∂C_1 in clockwise order, stopping at a_0, \dots, a_{m+1} as well as at some other places to be specified. We determine what p sees along the way, while keeping track of the current region of S_2 in which p lies. We use a start-up phase to initialize p and launch the fusion.

Start-Up. Using local shooting, we find the point of ∂C_1 that a_0 sees with respect to C_1 . Although a_0 is at worst infinitesimally close to ∂C_2 it does not always lie on it, as we shall see in the next paragraph. However, using the information about the endpoints of C_2 encoded in the

normal-form representation of S_2 (namely, pointers to incident arcs), we can find, in constant time, in which region of S_2 the point a_0 lies. This allows us to do local shooting and find the point of ∂C_2 that a_0 sees with respect to C_2 . These two pieces of information combine to give us the unique point c_0 of ∂C that a_0 sees with respect to C . We distinguish between two cases:

1. If c_0 belongs to ∂C_2 , then we set $p = a_0$ and we call the region of S_2 crossed by a_0c_0 *current*: the start-up phase is over (Figure 3.1.1).
2. If c_0 belongs to ∂C_1 from Lemma 2.1, the chord a_0c_0 splits ∂C into two curves, each closed under visibility. One of these curves, the one running from a_0 to c_0 clockwise, is a piece of ∂C_1 (Fig. 3.1.2), so the points of ∂C that its exit chord endpoints see all belong to ∂C_1 and thus are available directly from S_1 . We can therefore skip all the way to c_0 . Now, however, c_0 sees a point of ∂C_2 , namely a_0 , so we set $p = c_0$ and call the region of S_2 containing a_0 *current*.

Technically, it is not quite true that a_0 is always a point of ∂C_2 . It coincides with one most often, but when it sits at a local extremum (in the y -direction) it is not one because of duplication. What is true, however, is that when c_0 cannot see a point of ∂C_2 , an infinitesimal deformation of ∂C_2 locally around a_0 can make c_0 see one. This is a minor technicality which will not affect the remainder of the fusion algorithm, so for simplicity we still go on saying that c_0 sees a point of ∂C_2 with respect to C . Another minor problem is that a_0c_0 might lie on an exit chord of S_2 and thus there might be more than one candidate for the status of current region. We break ties by electing the region that we locally enter as we leave p in a clockwise traversal of ∂C_1 . This concludes the start-up phase. At this point we have the following situation (all visibility being understood with respect to C):

- (A) The points of ∂C that are seen by the exit chord endpoints of S_1 on the portion of ∂C_1 running clockwise from a_0 to the point p in its current position have all been determined already.
- (B) The point q of ∂C that is seen by p belongs to ∂C_2 and the chord pq lies in the region of S_2 called *current*. If p lies on a chord between two regions of S_2 , then the current region should be the one that we enter as we locally leave p clockwise around ∂C_1 .

These two conditions form our *loop invariant*, that is, they hold prior to every iteration of the process which we now describe.

Main Loop. Let A_i denote the oriented arc of S_1 running from a_{i-1} to a_i (in clockwise order around ∂C_1); by extension A_1 (resp. A_{m+1}) stands for the subarc extending from a_0 to a_1 (resp. a_m to a_{m+1}). Let A_k be the arc containing p . In the likely event that p is an endpoint of a chord of S_1 and thus belongs to two arcs, we must choose the one starting (and not ending) at p , i.e., we set the condition $p \neq a_k$. When p is set to a_{m+1} however, the algorithm simply terminates and no A_k need be defined. Let R denote the current region prior to entering the following loop: iterate through $j = k, k + 1, \dots$ until

- (i) a_j lies in R and the point of ∂C that a_j sees belongs to ∂C_2 (Figure 3.2.1), or
- (ii) the previous condition (i) does not hold, but R has at least one exit chord such that the point of ∂C seen by one of its endpoints belongs to A_j but strictly follows p (Fig. 3.2.2), or
- (iii) $j = m + 2$.

If case (i) occurs, find which point of ∂C is seen by a_j , declare that all a_i ($k \leq i < j$) see points of ∂C_1 (with respect to C), set $p = a_j$, let the current region still be R , and iterate through the loop, resetting k so as to comply with its definition. If case (ii) occurs, then of all the candidate endpoints, i.e., those chord endpoints satisfying (ii), determine the one which sees the point p' that is the last one encountered as we traverse ∂C_1 clockwise starting from p . In Fig. 3.2.2, for example, p' is the point labeled p_3 and the chosen endpoint is labeled q_3 . Next, declare that all a_i ($k \leq i < j$) see points of ∂C_1 , set $p = p'$, make current the region of S_2 which we enter as we locally cross the exit chord at p' along ∂C_1 , and iterate after updating k and R according to their definitions. In case (iii) we stop and, unless $k = m + 2$, we declare that all a_i ($k \leq i \leq m + 1$) see points of ∂C_1 . We have made several claims and skipped over important implementation issues in order to get the main idea of the algorithm across. Next, we fill in the missing parts and substantiate our claims.

Lemma 3.1. *It is possible to compute the fusion S of S_1 and S_2 in*

$$O((n_1/\gamma_1 + n_2/\gamma_2 + 1)(f(\gamma_2) + \log(n_1 + n_2)))$$

time.

Proof. We restrict our attention to the task of fusing S_1 into S_2 , the other case being similar. We have already shown that the start-up phase leads to a situation which satisfies the loop invariant, so it suffices to establish the correctness of the inner loop past a_c . In case (i) we know that a_j lies in R (actually in its interior) and sees a point of ∂C_2 , so invariant (B) is satisfied. How about (A)? We made the claim that a_k, \dots, a_{j-1} all see points of ∂C_1 . But actually the negation of (i) for a_k, \dots, a_{j-1} is not strong enough to reach the necessary conclusion about what a_i, \dots, a_{j-1} must see. Any of these points (if they exist) either sees ∂C_1 or lies outside of R . Why should lying outside R imply seeing ∂C_1 ? Suppose that, for some l ($k \leq l < j$), a_l lies in region R' distinct from R (like a_{k+1} in Figure 3.2.1) but also sees ∂C_2 . We derive a contradiction. Let Δ denote the directed portion of ∂C_1 as we go from p to a_1 clockwise, and let q (resp. q') be the point of ∂C seen by p (resp. a_l). The union of Δ , the chords pq and a_lq' , and the portion of ∂C_2 running clockwise (with respect to C_2) from q' to q forms the boundary of a simple polygon (Figure 3.3). Since the dual graph of a submap is a tree, there is a unique exit chord ab of R that leads to R' (note that ab need not be an exit chord of R' , since there might be one or even several regions separating R from R'). Because Δ runs from R' to R it passes through one of the chord endpoints, say, a . Let a' be the point of ab first encountered as we go from a to b along the chord. Note that ab cuts through Δ , so a' is well defined. The points a and a' see each other with respect to ∂C , and a' lies in A_h for some h between k and l inclusive. Because, in clockwise order around ∂C_1 , the point a' is leaving R locally, it cannot be equal to p . Therefore, the inequality $l < j$ implies that case (ii) must have already occurred when the running variable j was assigned to some integer between k and l , which is impossible.

Having shown that the loop invariant remains satisfied through case (i), we must do the same with case (ii). Let now Δ denote the directed portion of ∂C_1 as we go from p to a_j clockwise. The new assignment of p is the last point of Δ , distinct from p , that sees an endpoint of an exit chord of R . Certainly, the new assignment of the current region satisfies invariant (B). Figure 3.2.2 shows three candidate endpoints, with q_3 winning the contest. Turning now to invariant (A), we must prove our claim that the points of ∂C seen by a_k, \dots, a_{j-1} all belong to ∂C_1 . We omit the proof since it is identical to the previous one, with the role of a_j now played by p' .

What about termination? Obviously, the three cases rule out an infinite loop. Every time we fall in either of the two cases (i) or (ii) we determine more visibility information, so that all visibility relations are known from a_0 all the way to the current position of p . How about the last iteration,

the one leading to case (iii)? We claimed that all a_i ($k \leq i \leq m+1$) see points of ∂C_1 . This follows from the proof of the last paragraph, which showed that if a_i sees ∂C_2 , then either a_i lies in R (case (i)) or it does not, but then, we must fall in case (ii) after leaving p but upon or prior to getting to a_i . The proof of correctness is now complete.

Let us now analyze the complexity of the algorithm. To test whether a_j lies in R can be done in $O(f(\gamma_2))$ time by using the ray-shooters for each arc that bounds R : first we find which point of an arc is hit by a ray of light shot from a_j in its assigned chord direction. If there is no hit on any arc, a_j is not in R . Else, let s be the first point hit by the ray over all the arcs of R . Whether a_j lies in R or not can be directly inferred from the local orientation of the hit at s and which side of the double boundary is hit. This is because, as we should recall, arc-structures encode on which side(s) of the double boundary the arcs lie. If a_j lies in R , then s is the point of ∂C_2 seen by a_j with respect to C_2 . Next we use local shooting within S_1 to determine the point t of ∂C_1 hit by a ray of light shot from a_j in its assigned direction. Note that most often (i.e., when $0 < j \leq m$) a_j is the endpoint of a chord of S_1 so t is just the other endpoint of the chord. Now that we know which points of ∂C_1 and ∂C_2 the point a_j can see, we can immediately derive the point of ∂C that it sees and, from there, decide whether we are in case (i). The test takes $O(f(\gamma_1) + f(\gamma_2)) = O(f(\gamma_2))$ time (since f is nondecreasing and $\gamma_1 \leq \gamma_2$). This cost also includes the start-up phase.

Regarding (ii), we must assess how fast we can find the point of ∂C that is seen by an endpoint a of a given exit chord ab of R . Actually, we must find that point only if it belongs to A_j . So, we can shoot a ray of light from a toward A_j in the appropriate direction and see what happens, which takes $O(f(\gamma_1))$ time. If we do not get a hit, or if the hit does not lie on ab , or if it occurs before p along A_j , or if it does not have the proper orientation which lets a see A_j without the other side of the double boundary of A_j interfering (a constant-time test), then the endpoint can be disqualified. Otherwise, we must find whether the point s of A_j hit by the ray of light can see a with respect to C : the problem here is that other arcs A_i ($i \neq j$) might get in the way. Using local shooting in S_1 , however, we can shoot a ray of light from s toward a . We aim in the natural shooting direction from s since we have already ruled out that the “companion” point of s should prevent it from seeing a . The point t hit by the ray is found in $O(f(\gamma_1))$ time. If shooting from s to t passes through a , then s and a see each other with respect to C and we fall in case (ii), else we know that case (ii) cannot occur with respect to A_j and the endpoint a of the chord ab (although it might occur with respect to other endpoints of exit chords in R). This shows that testing case (ii) takes $O(f(\gamma_1))$ time.

We thus have shown that every elementary test (i), (ii) can be performed in $O(f(\gamma_2))$ time. At each such test we advance through the list of A_i or we report a pair of visible points in ∂C , one of which is an endpoint of an exit chord of S_2 . These reports are never duplicated because we always move forward among the A_i 's. Therefore, to discover all the chords of the fusion S of S_1 and S_2 takes time $O(mf(\gamma_2))$ time, where m is the total number of arcs and exit chords in S_1 and S_2 . By Lemma 2.3, m is $O(n_1/\gamma_1 + n_2/\gamma_2 + 1)$. Note that among the chords to be included in the fusion S , we not only have the newly discovered chords that connect ∂C_1 and ∂C_2 as well as the old chords of S_1 and S_2 that still form visible pairs of points with respect to C , but we also have all the null-length chords of S_1 and S_2 as well as the chords incident upon the vertices of ∂C resulting from the duplication of the vertex $C_1 \cap C_2$; this last category is where null-length chords originate.

After fusing S_1 (resp. S_2) into S_2 (resp. S_1), we have all the chords of the submap S and we can set it up in normal form quite easily. In order to allow for canonical vertex enumerations, let us sort the endpoints of these chords along ∂C , which is done by sorting the names of the edges of P on which these arcs abut, and then sorting the endpoints falling within the same edges by considering y -coordinates. This allows us to set up the required arc-sequence table. Note that merging can also be used instead of sorting but this step is not the dominant cost, anyway. With

this information it is now straightforward to set up the tree of the submap S , along with all the necessary arc-structures and their relevant pointers. Since the submap might not be conformal we dispense with the tree decomposition at this point. Very conservatively, all this work can be done in time

$$O((n_1/\gamma_1 + n_2/\gamma_2 + 1) \log(n_1 + n_2)).$$

Putting everything together, we derive the upper bound of

$$O((n_1/\gamma_1 + n_2/\gamma_2 + 1)(f(\gamma_2) + \log(n_1 + n_2)))$$

on the time needed to complete the fusion of S_1 and S_2 . \square

Remarks. 1. Since the chords of S reflect the visibility of the chord endpoints of S_1 and S_2 they need not be incident upon any vertex of ∂C , hence the notion of augmented maps and submaps.

2. It is possible to simplify the fusion procedure in various ways, albeit at the expense of a slightly more complicated proof of correctness. For one thing, the start-up phase is not strictly required and with a bit of care can be integrated within the main loop. Also, the two fusing passes can be unified into a single one (toward that end, we might want to catch the first occurrence of case (ii) and not the last one, for example).

3.2 Restoring Conformality

As we said earlier, there is no reason to believe that the fusion S should be conformal. Things can never be too bad, however. Indeed, let A_1, A_2, \dots be the arcs of a region R of S in counterclockwise order. It is clear that each A_l belongs to ∂C_1 or ∂C_2 but not both. So, we can partition the sequence of arcs into runs, B_1, B_2, \dots , meaning that $B_j = A_{i_j}, A_{i_j+1}, \dots, A_{i_{j+1}-1}$ is a maximal subsequence of arcs from either ∂C_1 or ∂C_2 (but not both). In the definition of maximal, we regard A_1, A_2, \dots as a circular sequence. Because any exit chord endpoint of S_i is still an endpoint of a chord in S and, with the possible exception of the chords incident upon a_0 or a_{m+1} , every chord of S that connects two points of ∂C_i is also a chord of S_i , it follows by conformality that a run associated with ∂C_i cannot have more than four exit chords in its midst, not counting the new chords incident upon a_0 or a_{m+1} . Therefore, a run cannot have more than a constant number of arcs. On the other hand, it follows from Lemma 2.2 that there are at most two runs. Why is that so? The lemma says that if we walk along ∂C clockwise we will in effect traverse, among other things, the boundary of R counterclockwise (except for the exit chords). If we begin our walk at one of the two points of ∂C corresponding to the vertex $C_1 \cap C_2$ we first exhaust, say, ∂C_1 and then ∂C_2 . Therefore, the counterclockwise traversal of the boundary of R must exhaust first the runs B_i contributed by S_1 and then the runs contributed by S_2 . Obviously, this leaves only the possibility of having at most one run of each type, and hence a total of at most two runs. The conclusion to draw is that, although not necessarily conformal, the submap S has no region with more than a bounded number of arcs. If S is not conformal we must now reduce the number of arcs per region to four or less by adding new chords into S . To discover these chords we need the ability to check whether two arcs or subarcs of the same region can “see” each other (Lemma 3.2). We also need to show that the desired chords do exist (Lemma 3.3).

Lemma 3.2. *Given two arcs A_1 and A_2 of the same region of S , assume that they have a pair of mutually visible points, one of which is a vertex of C (meaning that, say, A_1 contains a vertex v which is also a vertex of C and is such that the point of ∂C seen by v lies in A_2). Then we can find a point of A_1 (not necessarily a vertex of ∂C) that sees A_2 in time $O(f(\gamma_2)g(\gamma_2)(h(\gamma_2) + \log \gamma_2))$.*

Proof. To begin, observe that A_1 and A_2 are arcs or subarcs of either S_1 or S_2 but cannot overlap both ∂C_1 and ∂C_2 . The reason is that all chord endpoints in S_1 and S_2 are still chord endpoints in S (perhaps with different chords) and that we added chords incident upon the vertices of ∂C resulting from the duplication of the vertex $C_1 \cap C_2$. Therefore, because of the bounded number of arcs per region, it is still possible to do local shooting within any region of S . Since $\gamma_1 \leq \gamma_2$ and f is nondecreasing this takes $O(f(\gamma_2))$ time. Thus, we can efficiently check, in time $O(f(\gamma_2))$, whether a given vertex of A_1 qualifies as the point v sought. (Again, we must be careful that local shooting reports edges of P and does not tell us if the point hit is on the desired arc or is the companion of a point of the arc. We already discussed how local checking can decide which way it is in constant time, so we will not make further mention of that minor difficulty.) Of course, we should not check all the vertices of A_1 because there might just be too many of them. Instead, we need to do some kind of binary search among the vertices of A_1 .

For that purpose we invoke the arc-cutter associated with the arc of S_1 or S_2 containing A_1 , which allows us to subdivide A_1 into at most $g(\gamma_l)$ subarcs, with $l = 1$ if $A_1 \subseteq \partial C_1$ and $l = 2$ if $A_1 \subseteq \partial C_2$. Except for at most two single-edge subarcs attached to the endpoints of A_1 (which we ignore), for each subarc α we have a normal-form $h(\gamma_l)$ -granular conformal submap S_α of $V(\bar{\alpha})$. We search each subarc in turn, stopping as soon as we find a good vertex or point, if ever. Since the normal-form representation of S_α provides us with the tree decomposition T of the submap, we are able to check the candidacy of α in its entirety in $O(f(\gamma_2)(h(\gamma_2) + \log \gamma_2))$ time, provided that the following test can be performed in $O(f(\gamma_2))$ time: given a chord ab of S_α , either determine that a or b is a point of $\partial \bar{\alpha}$ and sees A_2 with respect to C , or find some $i \in \{1, 2\}$ such that $\alpha \cap \alpha_i$ is empty or has no point that sees A_2 , where α_1 and α_2 denote the two pieces of $\partial \bar{\alpha}$ between a and b . Note that $\partial \bar{\alpha}$ is a superset of α with twice the number of vertices (not fewer because the arc-cutter oracle guarantees that α does not double-back around an endpoint). First we show how such a test can be used to check the candidacy of α . Then we explain how to implement the test and why it covers all possible cases.

We begin by applying this test with respect to the chord corresponding to the root of the tree T (corresponding to the hierarchical decomposition of S_α). Then, as claimed, either we terminate with a positive answer or else we identify one of α_1 or α_2 , say, α_1 , such that $\alpha \cap \alpha_1$ is empty or has no point that sees A_2 . In that case, we find the child of the root that corresponds to α_2 and we iterate on this process from that node. This leads us to termination at some internal node of T or perhaps takes us to the bottom of the tree. Note that determining which node to branch to at each step is trivial once we have identified the α_i to be rejected. (So, we can perform the test just as stated above without having to “resize” α to reflect the current status of the ever-shrinking set of candidates.) If we reach a leaf, we examine each vertex of the region associated with it and, among those belonging to α , we check whether any of them can see A_2 . Since there are only $O(h(\gamma_2))$ vertices in the region and the depth of the tree is $O(\log \gamma_2)$ the running time of the algorithm is $O(f(\gamma_2)(h(\gamma_2) + \log \gamma_2))$, as claimed. Again we use the fact that $\gamma_1 \leq \gamma_2$ and that h is nondecreasing.

Whenever we discover a successful candidate point, the search can obviously be stopped right there. What remains to be seen is why upon reaching a leaf the corresponding region is the only one which can still provide the desired answer. Let us assume that the search ends up at a leaf. At the very beginning, let us say that each point of $\partial \bar{\alpha}$ is a candidate. Every time we branch down the tree we limit the candidacy to those points of $\partial \bar{\alpha}$ in the regions of S_α associated with the leaves of the subtree which we enter. At the end, the remaining candidates are the vertices in the region associated with the leaf where the tree search ends. This proves the correctness of the procedure. So, to summarize, if the basic test stated earlier can be performed in $O(f(\gamma_2))$ time, then we can solve the entire problem in $O(f(\gamma_2)g(\gamma_2)(h(\gamma_2) + \log \gamma_2))$ time, which proves the lemma.

We now show how to perform the test and why it is sound, i.e., covers all cases. Removing $\partial\bar{\alpha}$ from the spherical plane leaves two open regions, each polygonal and homeomorphic to a disk. One of them is infinitesimally small; let D be the closure of the other one. It is important that D should be homeomorphic to a closed disk and not to a 2-sphere, so the interior of $\partial\bar{\alpha}$, and, more generally, of C , should be understood as being very small but nonempty. Let c and d be the endpoints of α on $\partial\bar{\alpha}$. Removing c and d from ∂C leaves α and a curve A , both lying in D , so we have set the stage for Lemma 2.4. Figure 3.4 illustrates the correspondence: the snake on the left represents C ; the disk D corresponds to the outside of the portion of the snake between c and d , while it is the inside of the circle on the right. The curve A runs along the snake clockwise from d to c ; note that it runs on the boundary of D part of the way. The subarc α runs clockwise from c to d and corresponds to $B_1 \cup B_2$. Figure 3.4 shows the case where only one endpoint of ab lies in $B_1 \cup B_2$, which corresponds to Figure 2.8.2. The reader will easily draw an example matching the case of Figure 2.8.1, where both a and b lie in $\alpha = B_1 \cup B_2$.

To compute a' and b' (if defined) can be done by local shooting in the region of S incident upon A_1 and A_2 , which takes $O(f(\gamma_2))$ time. Note that no shooting is needed for a or b if the point in question does not lie in α . If ever a (resp. b) is a point of α and a' (resp. b') belongs to A_2 , then obviously we are done and successful in our search, so we can assume that neither conjunction holds. But, in that case, A_2 lies entirely within one of the connected components of the curve A after it has been cut up by removing a' and b' (whichever exists). Therefore, by Lemma 2.4, A_2 must be shielded from some B_j , which means that it cannot be connected to B_j without crossing ab or A . Furthermore, we know that B_j can be identified in constant time. The key observation now is that B_j coincides precisely with one of $\alpha \cap \alpha_1$ or $\alpha \cap \alpha_2$, say, the first one. It follows that no point of $\alpha \cap \alpha_1$ can see A_2 , and the test is completed. \square

Lemma 3.3. *Let A_1, \dots, A_k be the clockwise circular sequence of arcs around a region of S . If $k > 4$, then there exist i, j , such that*

- (i) $i - j \neq -1, 0, 1 \pmod k$ and
- (ii) A_i has a vertex of ∂C that sees A_j (with respect to C).

Proof. Recall that the region is associated with a subtree of the visibility tree of C . If S contains chords incident upon no vertices of ∂C , then we must include these chords as well and speak of the tree of the augmented visibility map of C . Let us add to the subtree in question the edges that connect it to the rest of the visibility tree. With respect to this augmented subtree, each exit chord of the region is associated with a distinct node of degree 1 (but the converse may not be true). Note that some of these exit chords might be of zero length. Consider the Steiner minimal tree of these particular degree-1 nodes (i.e., the smallest connected set of edges that join these nodes together), and for simplicity form a tree homomorphic to it by ignoring nodes of degree 2. Now embed this new tree in the plane and enclose it by a simple closed curve that connects together all its degree-1 nodes (Fig. 3.5). By using an embedding that preserves the local orientation of the edges around the nodes, the area inside that closed curve is partitioned into k faces, each corresponding to a distinct arc A_i . Because there are at least five degree-1 nodes and the maximum node-degree is 4, it is immediate that at least one edge of the tree must be incident upon two faces associated with A_i and A_j , respectively, where $i - j \neq -1, 0, 1 \pmod k$. (For a simple proof, try all possible cases with five nodes of degree 1 and observe that the property remains true with the addition of more nodes.) Of all the chords in the region only the exit chords might fail to be incident upon at least one vertex of ∂C . It follows that the edge in question corresponds to one or several chords of the original, nonaugmented $V(C)$ that connect A_i and A_j . \square

Equipped with the two previous lemmas, making S conformal is now quite easy. Recall that although S might not be conformal right after fusion, none of its regions has more than a bounded number of arcs. For any region with more than four arcs, let us apply Lemma 3.2 to every pair of nonconsecutive arcs until we find a chord which we can add to S . We iterate on this process until no region has more than four arcs. Note that although S keeps changing, Lemma 3.2 always holds since region arcs can only become smaller. Lemma 3.3 tells us that this chord addition process will not stop until S becomes conformal. Since the total number of arcs in S is $O(n_1/\gamma_1 + n_2/\gamma_2 + 1)$, we conclude:

Lemma 3.4. *The submap S can be made conformal in time*

$$O((n_1/\gamma_1 + n_2/\gamma_2 + 1)f(\gamma_2)g(\gamma_2)(h(\gamma_2) + \log \gamma_2)).$$

3.3 Maintaining Granularity

Since by making S conformal we did not remove any exit chord, it is still the case that, as observed in the proof of Lemma 3.2, no arc has more than γ_2 edges. Therefore, S is conformal and γ_2 -semigranular. We must now check whether the second criterion for γ_2 granularity holds. This criterion says that contracting any edge of the submap tree that is incident upon at least one node of degree less than 3 produces a new node whose weight exceeds γ_2 . This is very easy to enforce: if an edge does not pass the test, we just contract it by removing its corresponding exit chord (and those endpoints that are not vertices of ∂C). Note that this will not cause a violation of the first criterion, since the size of all the arcs will always remain within γ_2 . Similarly, the removal keeps the submap conformal. We process each exit chord in turn and check whether it is removable. Chords need be processed only once since the removals cannot make any chord removable if it was not already so before. Therefore, γ_2 -granularity, and more generally γ -granularity, for any $\gamma \geq \gamma_2$, can be enforced in this nondeterministic fashion in time linear in the size of the submap tree, that is, $O(n_1/\gamma_1 + n_2/\gamma_2 + 1)$. We can now put S in normal form, which includes computing its tree decomposition. As we discussed earlier, this can be done very simply in time $O((n_1/\gamma_1 + n_2/\gamma_2 + 1)\log(n_1 + n_2))$. With the inconsequential assumption that $\gamma_i = O(n_1 + n_2)$, we derive the following result from Lemmas 3.1 and 3.4:

Lemma 3.5. *Let C_1 and C_2 be two polygonal curves of n_1 and n_2 vertices, respectively, whose union forms a connected vertex-to-vertex piece of the input (simple and nonclosed) polygonal curve P . Suppose that we are given a normal-form γ_i -granular conformal submap of each $V(C_i)$, where $\gamma_1 \leq \gamma_2$, along with the ray-shooting and arc-cutting oracles necessary for merging. Then, for any $\gamma \geq \gamma_2$, it is possible to compute a normal-form γ -granular conformal submap of $V(C)$ in time $O((n_1/\gamma_1 + n_2/\gamma_2 + 1)f(\gamma_2)g(\gamma_2)(h(\gamma_2) + \log(n_1 + n_2)))$.*

3.4 Implementing the Oracles

Of the two oracles defined earlier, the ray-shooter is the more challenging to implement, the reason being that it addresses the key issue in the triangulation business, which is the discovery of new chords. The arc-cutter is implemented by using the divide-and-conquer structure of the up-phase of the visibility algorithm. Since a better understanding of the up-phase is necessary to understand how that oracle works, we postpone the discussion of its implementation a little. Turning our attention to the ray-shooting oracle, it might appear at first that fast planar point location should be the answer. But traditional methods, e.g., [9] and [19], are inadequate for several reasons, the

most crucial of which is their inability to support merging in sublinear time. We turn this problem around by exploiting further the approximation scheme provided by the concept of granularity.

Let C be a connected vertex-to-vertex piece of the input polygonal curve P and let m be its number of vertices. Let S be a normal-form γ -granular conformal submap of $V(C)$. So far, we have focused mostly on the tree structure of S . But let us now regard S as a planar graph. For this purpose, we must temporarily forget the fact that C has been given a double boundary. We define S^* to be the planar subdivision obtained by taking S and making every vertex (vertices of C and chord endpoints) a vertex on both sides of the double boundary, whose thickness is now null. As a result, the edges of S^* might be smaller than those of S but, unlike in S , no edge of S^* is of zero length (zero-length edges are now “contracted” into vertices). More important, each face of S^* coincides exactly with a distinct region of S , except for the fact that it might have many more vertices incident upon it. Indeed, a region’s only vertices are the endpoints of its own exit chords along with some vertices of ∂C , whereas the vertices of a face include all of the above plus all the chord endpoints that abut on the corresponding region from the outside. Notice that since the notion of double boundary is lost, a face might have dangling edges or edges incident upon it on both sides. There are several examples of this in Fig. 3.6, which shows the subdivision S^* corresponding to the submap of Fig. 2.6. Note also that the correspondence face/region is not surjective because empty regions have no associated faces. Besides being planar, the graph S^* has two remarkable properties:

- (i) From Lemma 2.3 we know that it has $O(m/\gamma + 1)$ faces, which is much smaller than the number of vertices (when γ is large).
- (ii) Although a given face might be very complex (i.e., incident upon many edges) its number of noncollinear edges is small, i.e., $O(\gamma)$.

These two features allow us to implement an efficient ray-shooting oracle.

Let G be the dual graph of S^* , that is, the graph obtained by associating a distinct node with each face of S^* and connecting two nodes if and only if they are distinct and their corresponding faces share a common edge. It is a classical result of graph theory that G is planar. How hard is it to compute G , say, in the form of adjacency lists? Two faces are adjacent if and only if either they share a chord or one of them has a chord endpoint that abuts on a nonnull length arc of the region associated with the other face. The first type of adjacencies can be detected immediately from S . The latter can be done by double identification, as discussed in Section 2.4, followed by sorting along ∂C , which takes $O(\mu \log m)$ time, where $\mu = O(m/\gamma + 1)$ is the number of nodes in G . It can also be done faster by merging chord endpoints along both sides of C .

If $\mu = 1$, then ray-shooting can be done trivially in $O(m)$ time, so let us assume that $\mu > 1$. We show that after $O(\mu \log m)$ preprocessing we can do ray-shooting in $O(\gamma \mu^{2/3})$ time. The planarity of G works wonders for us. The first payoff is that the number of edges is at most $3\mu - 6$. The second reward is that we can apply the linear-time algorithm of Lipton and Tarjan [21] to find a good separator. This partitions the nodes of G into three subsets A, B, D , such that

- (i) no edge joins a node of A with a node of B ,
- (ii) neither A nor B contains more than $2\mu/3$ nodes, and
- (iii) D contains at most $\sqrt{8\mu}$ nodes.

Let G_A (resp. G_B) be the graph obtained by keeping only the nodes of A (resp. B) and the edges of G that join only nodes of A (resp. B). We repeat the procedure over with respect to each of G_A

and G_B and iterate in this fashion until none of the graphs have more than μ^δ nodes, for some fixed δ ($0 < \delta < 1$). Let D^* be the set of all separators, i.e., the union of all the D 's. We easily verify that $|D^*| = O(\mu^\delta)$ provided that δ is chosen large enough; for example, $\delta = \frac{2}{3}$ [22]. In $O(\mu \log \mu)$ time we can compute D^* and partition the remaining nodes into subsets D_1, D_2 , etc., each of size at most $\mu^{2/3}$, such that no path of G can join two nodes in distinct subsets without passing through a node of D^* .

What is the utility of D^* for ray-shooting? Take a vertical line passing to the right of all the vertices of P , and intersect it with the chords of the regions in S . This breaks up the line into segments, every one of which falls entirely within some region; to split up the line and identify the regions cut by each segment can be done by traversing G and checking each chord for intersection with the vertical line. Since the regions cut correspond to nodes of G lying on a path, sorting the intersections comes for free, and all the work can be done in $O(\mu)$ time. We now claim that ray-shooting toward ∂C from any point can be done in $O(\gamma\mu^{2/3})$ time. Our first task is to shoot within each region that is dual to a node of D^* , using a naive algorithm which involves checking all the $O(\gamma)$ edges of the region (and not the edges of the face, which might be much more numerous). Assume that the ray of light hits a point among the edges of the regions dual to the nodes of D^* . Let R be the last region of S traversed before the first hit. To identify R can be done by double identification, followed by checking the local orientation of the hit. If R is a region dual to a node v of D^* , then the starting point of the ray lies in R (otherwise an earlier hit would have been detected) and we are trivially done.

So, assume now that R is dual to a node v not in D^* . Incidentally, note that the double identification needed to find R might require a binary search among a large collection of collinear edges. Let R' be the region incident upon the (region) edge containing the point of ∂C actually hit by the ray-shooting: this is the region that we are looking for (Fig. 3.7). If R and R' are not the same then the two regions can be connected by a horizontal line segment that avoids all the regions dual to D^* . It follows that the node w associated with R' can be reached by a path in G from v that avoids D^* . Consequently, v and w both lie in the same D_i . We can find w , and, from there, answer the ray-shooting query, by first finding D_i , which takes constant time since we know R , and then naively checking all the regions dual to nodes in D_i which takes $O(\gamma\mu^{2/3})$ time. Returning to our earlier case-analysis, assume now that the ray of light hits no region dual to a node in D^* . Then the ray-shooting takes place entirely within the regions dual to the nodes of a single D_i . To find out which one, we shoot toward the vertical line and find which segment of the line is hit. This takes $O(\log \mu)$ time by binary search. We can now identify the region R immediately. The remainder of the algorithm is unchanged. We conclude that ray-shooting can be done in $O(m)$ time if $\mu = 1$, and $O(\gamma\mu^{2/3})$ time if $\mu > 1$, after $O(\mu \log m)$ preprocessing. Since $\mu = O(m/\gamma + 1)$, we have:

Lemma 3.6. *Let C be a connected vertex-to-vertex piece of the input polygonal curve P and let m be its number of vertices. Let S be a normal-form γ -granular conformal submap of $V(C)$. Then it is possible to preprocess S in $O(m(\log m)/\gamma + 1)$ time so that ray-shooting within S can be done in time $O(\gamma^{1/3}m^{2/3})$.*

4 The Visibility Algorithm

Let P be a simple nonclosed polygonal curve with n vertices v_1, \dots, v_n . By padding the curve with additional vertices, if necessary, we can assume that $n = 2^p + 1$. Any subcurve of P of the form v_a, \dots, v_b , where $a - 1$ is a multiple of 2^λ and $b - a = 2^\lambda$ is called a *chain in grade λ* . Obviously,

- (i) a grade- λ chain has $2^\lambda + 1$ vertices,
- (ii) there are $2^{p-\lambda}$ chains in grade λ , and
- (iii) there are $p+1$ grades: $0, 1, \dots, p$.

We begin our work bottom-up, computing conformal submaps of granularity roughly m^β where m is the size of the underlying curve and β is some small enough positive constant; we set $\beta = \frac{1}{5}$, but to make the complexity analysis more explicit we leave β as a parameter in most of the calculations. We pursue the computation until the submap for the whole polygon has been obtained, which completes the up-phase. Then we reverse the process and work top-down until the submap has been completely refined into its full-fledged visibility map. The down-phase does not work by calling the visibility algorithm recursively on the regions of the top submap, but rather it uses data structures left behind during the top-phase (the submaps for the chains and their ray-shooting structures) to speed up the refinement process.

4.1 The Up-Phase

We begin with a piece of terminology: given a curve C consisting of m contiguous edges of P , we say that a submap of $V(C)$ is *canonical* if it is $2^{\lceil \beta \lceil \log m \rceil \rceil}$ -granular, conformal, and represented in normal form. Note that a canonical submap for a chain in grade λ is $2^{\lceil \beta \lambda \rceil}$ -granular. For $\lambda = 0, 1, \dots, p$ in that order, we process grade λ , which means:

- (i) We compute a canonical submap of $V(C)$ for each chain C in that grade.
- (ii) We preprocess each canonical submap for ray-shooting along the lines of Lemma 3.6, setting γ to the value $2^{\lceil \beta \lambda \rceil}$.

This work can be done naively for the early grades, so let us pick up the action at a grade λ larger than some appropriate constant, assuming that all grades less than λ have been processed already. We need the following result.

Lemma 4.1. *Suppose that all grades less than λ have been processed. Then, given any portion D of P of the form v_a, \dots, v_b , where $2^{\lambda-1} < b-a \leq 2^\lambda$, we can compute a canonical submap of $V(D)$ in time proportional to $\lambda^2(\log \lambda)2^{\lambda(1-\beta/3+4\beta^2/3)}$.*

Proof. In $O(\lambda)$ time we can partition D into $j \leq 2\lambda$ chains, D_1, \dots, D_j , in grades less than λ , with at most two chains per grade. This implies that, for each $i = 1, \dots, j$, a canonical submap S_i of $V(D_i)$ is available. Let γ be the granularity of a canonical submap of $V(D)$; we have $\gamma = 2^{\lceil \beta \lambda \rceil}$. Since the granularity of canonical submaps grows monotonically with the size of the underlying polygonal curve, we can trivially reset the granularity of each S_i to γ (Section 3.3). The time to do that is proportional to the total number of chords in all the S_i , which, from Lemma 2.3, is on the order of $\sum_{0 \leq k < \lambda} 2^{k-\lceil \beta k \rceil}$, that is, $O(2^{\lambda(1-\beta)})$.

Let us now merge these submaps two-by-two (D_1 with D_2 , D_3 with D_4 , etc.). More generally, we consider a perfectly balanced binary tree whose leaves are in bijection with the D_i and we merge submaps bottom-up by following the tree pattern. Application of Lemma 3.5 results in a canonical submap of $V(D)$ provided, of course, that the required oracles are available. But are they? Notice that during any merge any arc α in either of the two input submaps consists of at most γ edges. Therefore, any subarc $\alpha' \subseteq \alpha$ can be subdivided into a constant number of contiguous pieces (with no double-backing) whose corresponding portions of P consist of single line segments (at most two of them) and vertex-to-vertex pieces of P , each with at most $2^{\lceil \beta \lambda \rceil}$ edges. Each of

these pieces can be partitioned into a collection of $O(\lambda)$ chains in grades at most $\lceil \beta\lambda \rceil$. Our work at previous grades ensures that we have ray-shooting structures for the canonical submaps associated with these chains. Thus, to shoot a ray toward α' , we shoot toward each of the $O(\lambda)$ subarcs of its decomposition and determine the closest hit (if any). Shooting toward a single-edge subarc is trivial. Shooting toward any other subarc makes use of the shooting structure of a canonical submap for a chain in grade $\mu \leq \lceil \beta\lambda \rceil$. Assuming that $\lceil \beta\lambda \rceil < \lambda$ (which is true for λ large enough) all these shooting structures have been computed and therefore, by Lemma 3.6, ray-shooting can be done in time $O(2^{\lceil \beta\mu \rceil/3+2\mu/3})$, which is $O(2^{\beta^2\lambda/3+2\beta\lambda/3})$. Since there are $O(\lambda)$ subarcs, it follows that the ray-shooting oracle can be implemented so that

$$f(\gamma) = \lambda 2^{\beta^2\lambda/3+2\beta\lambda/3}.$$

As we mentioned, the subarc α' is decomposed into at most two single-edge pieces, along with $O(\lambda)$ pieces for which we have conformal submaps of granularity at most $2^{\lceil \beta\lceil \beta\lambda \rceil \rceil}$. We verify that all the requirements of the arc-cutting oracle are satisfied by this decomposition, so that we can set

$$g(\gamma) = O(\lambda)$$

and

$$h(\gamma) \leq 2^{\lceil \beta\lceil \beta\lambda \rceil \rceil}.$$

To appreciate the connection between the left and right-hand sides of these relations, recall that γ and λ are related by the identity $\gamma = 2^{\lceil \beta\lambda \rceil}$. By Lemma 3.5, if a merge takes input curves with a total of m vertices, then the time to carry it out is at most proportional to

$$\lambda^2(m/2^{\beta\lambda})2^{\beta^2\lambda/3+2\beta\lambda/3}(2^{\beta^2\lambda} + \log m).$$

There are $O(\log \lambda)$ levels of merging to be performed, each involving a total of $b - a \leq 2^\lambda$ edges, therefore the time to merge the submaps for all the D_i 's into one is at most (up to within a constant factor)

$$\lambda^2(\log \lambda)2^{\lambda-\beta\lambda/3+4\beta^2\lambda/3}.$$

Since the initial cost of resetting the granularity is only $O(2^{\lambda(1-\beta)})$ the lemma follows readily. \square

Let us now turn to the processing of grade λ . Lemma 4.1 can be called upon to compute a canonical submap of the visibility map of each chain in grade λ . Preprocessing each chain for ray-shooting is done by using Lemma 3.6. Since there are $(n-1)/2^\lambda$ chains in grade λ , we conclude that processing grade λ requires time at most proportional to

$$n\lambda^2(\log \lambda)2^{\beta\lambda(4\beta/3-1/3)} + n\lambda 2^{-\beta\lambda}.$$

From our choice of $\beta = \frac{1}{5}$ it follows that preprocessing grade λ takes $O(n2^{-\lambda/76})$ time, therefore processing all $p+1$ grades, and thereby completing the up-phase, takes linear time.

4.2 The Down-Phase

Now that we have canonical submaps for each chain in each grade, along with their oracle structures and tree decompositions, we are ready to refine the canonical submap of $V(P)$. This is done incrementally by going down the tree. The following lemma provides the key to the algorithm:

Lemma 4.2. *Let λ be any positive grade and let C be an arbitrary chain in any grade $l \geq \lambda$. If a $2^{\lceil \beta \lambda \rceil}$ -granular conformal submap of $V(C)$ is available in normal form, then it is possible to compute $V(C)$ in time at most $(c - 1/\lambda)2^l$, where c is some constant large enough.*

Proof. We proceed by induction on λ . Let S be the $2^{\lceil \beta \lambda \rceil}$ -granular conformal submap of $V(C)$. The case where λ is a constant is trivial since the regions of S have bounded size, and therefore the missing chords can be provided in constant time per region. So, let us switch directly to the inductive case, assuming that λ is large enough. Let R be a region of S . Because of conformality, the union of all the arcs of R can be partitioned into a constant number of single edges and vertex-to-vertex pieces of C with at most $2^{\lceil \beta \lambda \rceil}$ edges. Applying Lemma 4.1, we can compute a canonical submap for each connected polygonal piece in the partition in time at most proportional to

$$\lambda^2(\log \lambda)2^{\beta\lambda(1-\beta/3+4\beta^2/3)}.$$

Each of these submaps has granularity at most $2^{\lceil \beta \lceil \beta \lambda \rceil \rceil}$ so we can pursue the merging by putting together all these submaps and thus create a single normal-form $2^{\lceil \beta \lceil \beta \lambda \rceil \rceil}$ -granular conformal submap of $V(R^*)$, where R^* is the boundary of R minus a vertex (to ensure that it is nonclosed). For consistency, we should regard R^* as a standard polygonal curve and not as part of a double boundary. The operation requires a constant number of merges, so we can carry it out effectively by merging submaps two-by-two like in Lemma 4.1.

There is a small subtlety in this last round of merges, which we should explain. To take a simple example, suppose that R^* has two arcs and two exit chords: $\alpha_1, a_1b_1, \alpha_2, a_2b_2$, in cyclic order. It could be that the endpoints of α_1 or α_2 are not vertices of ∂C , so to deal with the most general case, assume that α_1 consists of $b_2b'_2, \beta_1, a'_1a_1$ and α_2 consists of $b_1b'_1, \beta_2, a'_2a_2$ where a'_1, b'_1, a'_2, b'_2 are all vertices of ∂C (Fig. 4.1). Let S_1 (resp. S_2) be the canonical submap for the vertex-to-vertex piece of P corresponding to β_1 (resp. β_2) and let T_1 (resp. T_2) be canonical submaps for the 3-edge polygonal curve $a'_1a_1b_1b'_1$ (resp. $a'_2a_2b_2b'_2$). We obtain S_1 and S_2 by application of Lemma 4.1, while T_1 and T_2 are computed directly (tilting the edges a_1b_1 and a_2b_2 symbolically to keep the merging algorithm from complaining later).

We are now ready to merge S_1 with T_1 then merge the resulting submap with S_2 , and finally merge the result with T_2 . Note that we treat the edges a_1b_1 and a_2b_2 as part of the input curve although they are not part of P . As a result, ceasing (temporarily) to be chords, these edges cannot be removed during the merges. Since we add at most a constant number of new edges to the input curve, all the oracle machinery needed for the merges is still available, i.e., the new edges create only constant-time multiplicative overhead. Although the final submap is conformal it might no longer be so if we now reinterpret a_1b_1 and a_2b_2 as chords, which we do once the last round of merges is completed. To remedy this we apply the conformality-restoring procedure of Section 3.2 to each region that might have more than four chords with this new interpretation. Again, it is immediate to see that all the required oracles are still available.

The time taken by this last round of merges is dominated by the cost of the earlier merges, so computing the $2^{\lceil \beta \lceil \beta \lambda \rceil \rceil}$ -granular conformal submap of all the $V(R^*)$'s takes time at most proportional to

$$2^l \lambda^2(\log \lambda)2^{\beta^2\lambda(4\beta/3-1/3)}.$$

We can now extract the relevant information, i.e., the exit chords falling entirely within each region R . This involves checking the exit chords of the computed submap of $V(R^*)$ and keeping only those both of whose endpoints lie on the arcs (in the double boundary sense) of the region R . This leads to a new map S^* of $V(C)$ which is a refinement of S : all its arcs originate from the previous

merges, therefore S^* is a $2^{\lceil \beta \lceil \beta \lambda \rceil \rceil}$ -semigranular conformal submap of $V(C)$. We can only speak of semigranularity because some of the chords connecting the R^* 's might be removable now. We can check each of the exit chords directly, which as we saw in Section 3.3, takes a total amount of time linear in the number of exit chords in S^* .

Now that we have a $2^{\lceil \beta \lceil \beta \lambda \rceil \rceil}$ -granular conformal submap of $V(C)$ at our disposal we observe that $\lceil \beta \lambda \rceil \leq \lambda - 1$ for λ large enough, so that we can apply the induction hypothesis and derive $V(C)$ in time at most $(c - 1/(\lambda - 1))2^l$. Putting everything together, the total running time for the construction of $V(C)$ is at most

$$a2^l \lambda^2 (\log \lambda) 2^{\beta^2 \lambda (4\beta/3 - 1/3)} + (c - \frac{1}{\lambda - 1})2^l$$

for some constant $a > 0$. With the setting $\beta = \frac{1}{5}$, this is no more than

$$a2^l \lambda^2 (\log \lambda) 2^{-\lambda/375} + (c - \frac{1}{\lambda - 1})2^l \leq (c - \frac{1}{\lambda})2^l$$

for λ large enough. \square

During the up-phase we built a normal-form $2^{\lceil \beta p \rceil}$ -granular conformal submap of $V(P)$ in linear time. By Lemma 4.2, therefore, $V(P)$ can be obtained also in linear time. As demonstrated in [6] and [11], a triangulation can be easily derived from the visibility map in linear time, so our main goal has been reached.

Theorem 4.3. *It is possible to compute the visibility map of a simple polygonal curve, and, hence, a triangulation of a simple polygon, in linear time.*

5 Some Applications and Open Problems

There are many uses for a fast triangulation algorithm. We mention only two of them and refer the reader to [1], [13], [14], [16], [20], [24], [26], and [28] for some pointers to other applications. Kirkpatrick [19] and Edelsbrunner et al. [9] have given optimal planar point-location algorithms which require linear preprocessing time provided that the planar subdivision has triangulated or monotone faces. With our algorithm the preprocessing can be made linear as long as the graph is connected.

The second application follows from the observation that the triangulation algorithm can be adapted to check whether a polygonal curve is simple. We briefly outline the method. We begin by computing the visibility map. In all likelihood the program will crash if the polygonal curve is not simple. But if it does not, then we are handed over a would-be visibility map with its corresponding visibility tree. In linear time we can verify that the map is locally sound by checking that all regions and adjacencies are topologically and geometrically satisfactory. If the curve has a self-intersection p , then two regions must strictly overlap around that point, but this can be shown to contradict the local soundness of the map. Indeed, draw a line connecting p to any point q that provably belongs to only one region (if an arbitrary choice of q does not work, then obviously the curve is not simple). By a continuity argument, the segment pq must cross an edge at a point where the number of overlapping regions differs locally around that point. But, occurring on an edge, this must be detected by the local tests. So, to summarize, a linear number of local geometric tests suffice for testing simplicity, once the output of the visibility algorithm is available.

From this we easily derive a new result: testing whether two simple polygons intersect in linear time. The reduction of this problem to simplicity-testing was observed by Dobkin et al. [8].

Slightly simplified, their method consists of taking the highest points of both polygons and shoot horizontally from the lower point p toward the other polygon in both directions. If there is no hit, then no intersection can occur. Otherwise, we can immediately infer from the local orientation of the hits whether p lies inside or outside the polygon. In the first case we conclude with a positive answer, else we connect the two polygonal boundaries into a single one by adding one of the connecting segments and duplicating it, thus reducing the problem to that of testing simplicity.

It is interesting to notice that our algorithm can be used to perform Jordan sorting in linear time. This provides a completely different alternative to Hoffman et al.'s algorithm [18]. Recall that the problem is to sort a sequence of numbers a_1, \dots, a_n , which correspond to the intersection points of a Jordan curve with the x -axis. Assuming that we do not know anything else about the curve, we construct the polygon P with vertices $A_1, B_1, A_2, B_2, \dots, A_n, B_n$, where $A_i = (a_i, 0)$ and

$$B_i = \left(\frac{a_i + a_{i+1}}{2}, (-1)^i \left| \frac{a_{i+1} - a_i}{2} \right| \right),$$

with $a_{n+1} = a_1$. It is immediate that P is simple and that its visibility map gives us the a_i in sorted order. (Recall that to have many vertices with the same y -coordinate is not a serious problem). It is instructive to compare the two methods: Hoffman et al.'s algorithm is on-line and corresponds to an asymptotically optimal search scheme for Jordan permutations. Our method works off-line and uses divide-and-conquer. It is an intriguing open question whether triangulation can be done on-line in a manner similar to [18]. Tarjan and Van Wyk's method (almost) falls in that category but it is not optimal. Can their algorithm be made optimal? More generally, is it possible to maintain the visibility map optimally under on-line insertion of new edges?. Obviously not in an explicit fashion, since a new edge can cut through a linear number of diagonals, hence creating a quadratic blowup. Even implicitly, however, it can be trivially shown that identifying the order types of the visibility maps of all the prefixes of a simple n -vertex polygonal curve requires $\Theta(n \log n)$ bits, which is bad news. Thus, something with less information content should be maintained by an optimal on-line algorithm. But what? This question might seem rhetorical in light of our linear-time algorithm, but the underlying issue is whether a fundamentally different optimal algorithm exists, in particular, one that works on-line. A close look at the up-phase of our visibility algorithm shows that constructing the computation tree in symmetric order is tantamount to inserting the edges one at a time along the boundary and maintaining a small collection of distinct visibility submaps. Since these maps provide only partial visibility information this can be looked at as a partial answer to our question.

Also, we might wonder whether there exists a simple optimal probabilistic triangulation algorithm, say, one as straightforward as Clarkson et al.'s [7] or Seidel's [25]?. Our algorithm can be modified by replacing the planar separator step by a randomized construction, but whether the resulting algorithm is really simpler is debatable. We close by mentioning what is probably the most interesting open question left on the subject of polygonal curves. This is the problem, previously posed by Tarjan and Van Wyk [27], of computing all the self-intersections of a nonsimple polygon in time linear in the sum of the input and output sizes.

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