

OPTIMAL RANDOMIZED PARALLEL ALGORITHMS FOR COMPUTATIONAL GEOMETRY I*

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Abstract

We present parallel algorithms for some fundamental problems in computational geometry which have running time of $O(\log n)$ using n processors, with very high probability (approaching 1 as $n \rightarrow \infty$). These include planar point location, triangulation and trapezoidal decomposition. We also present optimal algorithms for 3-D maxima and two-set dominance counting by an application of integer sorting. Most of these algorithms run on CREW PRAM model and have optimal processor-time product which improve on the previously best known algorithms of Atallah and Goodrich [3] for these problems. The crux of these algorithms is a useful data structure which emulates the plane sweeping paradigm used for sequential algorithms. We extend some of the techniques used by Reischuk [22] Reif and Valiant [21] for flashsort algorithm to perform divide and conquer in a plane very efficiently leading to the improved performance by our approach.

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1 Introduction

Recently there has been a growing effort towards developing efficient parallel algorithms for various fundamental problems in computational geometry. Aggarwal et al. [1] gave polylog algorithms for a various fundamental problems viz. Voronoi diagram, Planar point location, Triangulation among others. Atallah and Goodrich [3] improved these algorithms by reducing the running time of most of these algorithms to $O(\log n \cdot \log \log n)$ time using $O(n)$ processors. Hence, the previous efforts are sub-optimal by at least a factor of $\log \log n$, since most of these problems need $\Theta(n \log n)$ time in the sequential case. For an optimal processor-time product the running time for most of these algorithms should be $O(\log n)$ using $O(n)$ processors.

Randomization has been used in a wide number of applications successfully (for example see [21], [16], [24]) and seems to be an ideal choice for optimizing algorithms for various problems in computational geometry. Clarkson ([5] and [6]) used random sampling techniques to derive better upper-bounds for algorithms for the post-office problem and higher-order Voronoi diagrams as well as some combinatorial quantities like k -sets and polytope-separation. Haussler and Welzl [14] also used random sampling to solve half-space query problems. This is the first paper where random sampling methods have been applied to yield efficient parallel algorithms in computational geometry. Clarkson's methods yielded bounds for the *expected* running time of the algorithms. In contrast, we are able to bound the tail estimates of the running time with high probability. It is not clear how Clarkson's algorithms can be modified to obtain high-probability bounds without increasing the asymptotic running time of the algorithms (i.e. since the algorithms terminate in the claimed time bounds with constant probability, one needs to repeat the algorithm $O(\log n)$ times to decrease the failure probability to $1/n$). Although this may not be very important in a sequential environment, it turns out to be very crucial for bounding the resources used by a parallel algorithm. This will become clearer in the proof of Theorem 2, where expected bounds do not appear to be sufficient for deriving the claimed bounds.

The term *very high likelihood (probability)* has been used in this paper to denote probability $> 1 - n^{-\beta}$ for some $\beta > 1$ where n is the input size. Just like the big-O function serves to represent the complexity bounds of deterministic algorithms, we shall use \tilde{O} to represent complexity bounds of the randomized algorithms. We say that a randomized algorithm has resource bound $\tilde{O}(f(n))$ if there is a constant c such that the resource used by the algorithm is no more than $cf(n)$ with probability $\geq 1 - 1/n^\alpha$ for any $\alpha > 1$. (An equivalent definition will be bounding the resource by $\alpha \cdot f(n)$ with probability greater than $1 - n^{-c\alpha}$ and in the rest of the paper they will be used in an interchangeable manner).

We have used some known techniques methods like *random mate*, and extended techniques of *Flashsort* [21] to two-dimensions successfully to reduce the running time of a wide variety of fundamental problems. Random mate was introduced in [17] to discard a constant fraction of zeroes in a zero-one string in constant time, and was also later used by Gazit [12]. Random sampling methods used by Reischuk [22] and Flashsort provided the initial inspiration for divide and conquer on the plane; although the problem at hand being much harder, required several additional techniques. We note that Clarkson [7] independently derived bounds for doing divide-and-conquer efficiently for geometric problems in a more general setting.

The algorithms in this paper have optimal speed-up when compared to their corresponding sequential cases. Table 1 summarizes the results in this paper and compares them to the em

previously ¹ best known results. The organization of the paper is as follows : In section 1, we present a very simple algorithm for planar-point location which demonstrates the usefulness of randomization. In section 2, we present a new data structure which can be constructed efficiently using random-sampling, and we call it **nested-plane-sweep** tree. In the following section we apply this data structure to develop optimal algorithms for a number of problems for which only efficient deterministic algorithms were known. In the final section we reduce problems like 3-D dominance and range counting to integer sorting which can be then be optimally solved using an algorithm similar to Reif [20]. Since the currently best-known algorithm for integer sorting is optimal for word-size n^ϵ , for any $\epsilon > 0$, our algorithms are not optimal in an information-theoretic sense (i.e. the sequential algorithms use only $O(\log n)$ bits per word). However, we feel that these are extremely simple and should be of interest from a practical viewpoint.

Table 1

Summary of our results		
Problem	Previous bounds	Our bounds
Planar Point location	$O(\log n \log \log n)$	$\tilde{O}(\log n)$
Trapezoidal decomposition	$O(\log n \log \log n)$	$\tilde{O}(\log n)$
Triangulation	$O(\log n \log \log n)$	$\tilde{O}(\log n)$
3-D Maxima	$O(\log n \log \log n)$	$O(\log n)$
Two-set dominance counting	$O(\log n \log \log n)$	$O(\log n)$
Multiple Range-counting	$O(\log n \log \log n)$	$O(\log n)$
Visibility	$O(\log n \log \log n)$	$\tilde{O}(\log n)$

2 Point Location In Planar Subdivisions In Logarithmic Time With High Probability

A PSLG (planar straight line graph) is a connected graph which can be embedded on a plane using only straight line edges. Given a PSLG and a query point, the point location problem is to identify the subdivision in the plane (induced by the PSLG) which contains the query point. If a PSLG has n vertices, in the sequential (uniprocessor) case, the asymptotic optimal query time performance of $O(\log n)$ can be achieved by a variety of approaches (Kirkpatrick [15], Edelsbrunner [13] among others). The performance is achieved by binary search on a well organized data structure constructed on the PSLG during the preprocessing. The preprocessing cost for the best known sequential algorithms for point location is $\Omega(n \log n)$ operations, and thus the best parallel time for this problem cannot be better than $O(\log n)$ time using n processors. We propose a randomized method by which the preprocessing can be done in $O(\log n)$ time with very high probability using $O(n)$ processors given a PSLG which has only convex subdivisions. A similar algorithm has also been independently proposed by Dadoun and Kirkpatrick [10].

¹Optimal deterministic algorithms for these problems were obtained independently by Atallah, Cole and Goodrich [2]. Also, see conclusions for some additional comments.

2.1 The sequential algorithm

Kirkpatrick [15] proposed a triangulation refinement technique which builds a hierarchy of triangulated PSLG from the initial graph. Unfortunately, his method does not appear to be directly parallelizable. A review of his method will help the reader to understand the parallel version presented here. Starting with a triangulated PSLG (i.e. all faces including the exterior face are triangular), his algorithm removes an independent set of vertices, and retriangulates the remaining graph. In addition, it keeps track of the set of the eliminated triangles that have non-empty intersection with each of the new triangles of the retriangulated graph. This procedure is repeated successively until we are left with a constant number of triangles. Actually we can stop when the problem size is reduced to $O(\log n)$. The search proceeds in a hierarchical manner from the highest level, where the problem can now be solved in constant time ($O(\log n)$ time if we stop at $O(\log n)$ size graph). Subsequently, at each level we relocate the point with respect to the triangles which intersect the triangle in the current level, thus refining the regions of location at each level. At the lowest level the point is located with respect to the regions of the original PSLG and the search is complete. The search data structure is a directed acyclic graph (not a tree). Each of the nodes represent a triangular region, and is connected by directed arcs to all the triangular regions that it intersects in the previous level. At any node, the algorithm determines which of its children the point lies in. Notice that a node can have more than one parent and hence the graph (the underlying search structure) is not a tree. The efficiency of this approach depends on the number of levels of the triangulated PSLG, and the number of intersections of each triangle with the triangles in the next lower level. By guaranteeing that a constant fraction of the vertices, and hence the triangles, (since it is a planar graph) are eliminated at each successive level a logarithmic level search is achieved. However in doing so, one must also be careful that a triangle intersects a maximum of a constant number of triangles in the next level (i.e. each node has a bounded constant degree so that a constant time is spent at each search level). Both of the above criteria can be satisfied by identifying an independent set of vertices each having a degree $\leq d$, where d is a fixed constant. The number edges in a planar triangulated graph (V, E) is $3|V| - 6$ from Euler's formula (assuming that the exterior face is also a triangle). This adds up to a total vertex degree of $6|V| - 12$, giving us a lower bound on the number of vertices with degree less than d . Thus the number of vertices with degree less than d is at least $6|V|/d - 2$ in a triangulated graph for $d > 6$, (a typical value of d is 12). Since a constant fraction of these vertices can be eliminated at each stage by identifying a maximal independent set of vertices, the height of the search tree is at most $O(\log |V|)$. The preprocessing time is dominated by the identification of this independent set of degree $\leq d$, which costs linear in size of the PSLG at each level resulting in total time of $O(n)$. If the initial PSLG is not triangulated, the cost of triangulization dominates the pre-processing time.

In the discussion below we present a randomized method to identify a large independent set in constant time for each level using n processors with a high probability. As a result, the search structure can be constructed in $O(\log n)$ time.

2.2 Choosing a large independent set with high probability

Associate a processor with each vertex and each edge of the PSLG (V, E) . From Euler's formula we need only $O(|V|)$ processors for the PSLG. An independent set consisting of vertices of degrees

less than or equal to d can be identified in the following manner. ²

Algorithm Random-mate

Input: A PSLG in form of a doubly connected edge list (DCEL).

Output: A large independent set of vertices with degree $\leq d$.

(1) Identify the set of vertices with degree $\leq d$. This can be done in constant time using one processor for each vertex. There will be at least $6|V|/d$ (ignoring the small constant) such vertices (from the previous discussion).

(2) This has two phases :

(2a) Randomly assign a tag ‘male’ or ‘female’ with equal probability to each selected vertex from step (1). The ‘males’ constitute the candidates for an independent set. For each edge between two ‘males’ pronounce both vertices ‘dead’ by marking their corresponding cells. This is easily done by using one processor for every edge with concurrent write capability. Note that it is very easy to get rid of the concurrent-read feature since a vertex has at most degree d and hence this step can be carried out in constant number of steps (instead of exactly one step).

(2b) The ‘males’ which are not ‘dead’ form an independent set.

(3) Output the set X of ‘males’ which are not ‘dead’. The set X is an independent set of vertices having degree less than d .

It is obvious that the algorithm gives us an independent set which is possibly smaller than the maximal independent set (a null set in the worst case for a chain of males). However, we shall show that on the average this technique will produce an independent set proportional to a constant fraction of the total number of nodes (vertices of degree $\leq d$) with a very high probability.

Lemma 1 *Let $n = |V|$ and n_d denote the number of vertices with degree less than or equal to d ($n_d \geq 6n/d$ from [15]). There exist constants c, ν ($0 < \nu < 1$), such that $P(|X| < \nu n) \leq e^{-cn}$.*

Proof: Assuming equal probabilities for a vertex being assigned a ‘male’ and a ‘female’ tag, the probability of a vertex being in the independent set is greater than or equal to $(\frac{1}{2})^{d+1}$ (since it has $\leq d + 1$ neighbors). We consider an independent set $I_3(n_d)$ of vertices which are at a distance 3 (i.e. the shortest distance between any two vertex in this set is three edges). Since the graph had a bounded degree d , the selection of each vertex can prevent the selection of at most $d_{max} = (d-1)^3 + (d-1)^2 + (d-1)$ other vertices. Thus the cardinality of this set is at least $(1/d_{max})6n/d$ or $6n/D$ (D is a constant). The event of a vertex $v \in I_3(n_d)$ being in X (independent set) is *independent* of any other vertex in $I_3(n_d)$. Thus the distribution of the cardinality of the

²this is the random-mate technique mentioned earlier

independent set produced by algorithm Random-mate can be bounded from below by a binomial distribution with $p = \left(\frac{1}{2}\right)^{d+1}$, and mean $\mu = |I_3(n_d)|p \geq 6np/D$.

Using Angluin-Valiant's bounds (see equation (3) in appendix), for $0 < \epsilon < 1$,

$$Prob[|X| \leq (1 - \epsilon)\mu] \leq \exp(-\epsilon^2\mu/2)$$

Using $\nu = (1 - \epsilon)6/D$ and $c = \epsilon^2/(12D)$ we arrive at the required form. \square

2.3 A parallel algorithm

We have shown that with very high probability, the size of the independent set will be greater than a constant fraction of n , which is the key to finding a logarithmic depth search structure. We present the complete **Point-Location-Tree** algorithm below

Procedure Point-Location-Tree

(0) Let N be the number of vertices in the current stage. If $N \leq k \log n$ (for some fixed constant k), then halt. (then we can locate the query point in the $O(\log n)$ triangular faces in $O(\log n)$ time using a single processor).

(1) Assume that the given PSLG is triangulated. (If not, then we can use the procedure for triangulation described in the latter sections to triangulate a PSLG in $O(\log n)$ time using $O(n)$ processors.)

(2) Choose an independent set of vertices each of which has degree ≤ 12 using the method described above with $d = 12$. Time = $O(1)$.

(3) Triangulate the remaining PSLG (after the removal of the independent set). Note that removal of a vertex of degree d ($d \leq 12$) necessitates triangulating a simple polygon of d vertices. This can be done in constant time using 1 processor for each polygon.

(4) Determine for each of the new triangles (created in step 3), which of the old triangles it intersects. This can also be done in constant time using one processor for each of the new triangles using concurrent read.

(5) go to (0)

Theorem 1 *Algorithm Point-Location-Tree runs in $\tilde{O}(\log n)$ time using $O(n)$ processors and $O(n \log \log n)$ space in a CREW PRAM model. Furthermore, if the input PSLG is triangulated, we can reduce the processor bound to $O(n/\log n)$ without increasing the asymptotic running time of the algorithm.*

Proof : Each of the steps at any level of recursion of the algorithm can be done in constant time using $O(n)$ processors. From Lemma 1, there exist constants ν and c such that the probability of reducing the problem by a constant fraction $(1 - \nu)$ is very high. Let n_i denote the problem size (the number of vertices remaining in the graph) at stage i of recursion, where $n_0 = n$. We shall show that after $O(\log n)$ stages, the problem size is less than $O(\log n)$ with very high likelihood.

From lemma 1, $P[n_i \leq (1 - \nu)n_{i-1}] \geq 1 - n^{-cK}$ for $n_{i-1} \leq K \log n$ for some constant K . The

probability that this fails to hold in any level i ($1 \leq i \leq \log_{1/(1-\nu)} n$) is less than $n^{-cK+\delta}$ for any $\delta > 0$. We abort the algorithm if the number of vertices is greater than $K \log n$ after the last stage and re-run the entire procedure. The argument for space is similar to the sequential case. We need an extra $O(\log \log n)$ factor space for the triangulation procedure.

If the input PSLG is triangulated, the number of operations performed is linear in n (follows from the sequential algorithm) and hence by Brent's slow-down procedure and using the load-balancing scheme of Cole and Viskin [9] or the randomized scheme of Miller and Reif [17], the processor bounds can be reduced to $O(n/\log n)$ without affecting the asymptotic run-time. \square

REMARK : *Dadoun and Kirkpatrick [10] show that their algorithm runs in $O(\log n)$ expected parallel time. However they do not extend their analysis for the high-probability bounds which is also $O(\log n)$ as shown in the previous theorem.*

Corollary 1: Given a planar subdivision S , consisting of $O(n)$ vertices, we can locate $O(n)$ query points in $\tilde{O}(\log n)$ time using $O(n)$ processors.

Proof: Follows immediately from the previous discussion. \square

A direct application of this result leads to the improvement in performance of Aggarwal et al. [1] two-dimensional Voronoi diagram algorithm.

Corollary 2: The Voronoi diagram of a set of $O(n)$ points on a plane can be constructed in $O(\log^2 n)$ expected running time using $O(n)$ processors in a CREW PRAM model.

Proof : Aggarwal et al. [1] uses a divide and conquer algorithm, where in each of the $\log n$ stages of recursion we need to perform point location simultaneously for $O(n)$ vertices. Since their planar-point location needs $O(\log^2 n)$ time, the overall algorithm runs in $O(\log^3 n)$ time. Using our randomized planar-point location algorithm, the expected running time of algorithm is $O(\log^2 n)$. \square

3 Constructing The Plane-Sweep Tree In $O(\log n)$ Parallel Time Using Randomization

There are many problems in computational geometry whose optimal sequential algorithms use the plane-sweeping paradigm. Aggarwal et al. [1] had proposed a data-structure, which they call **plane-sweep-tree**. Using this, they were able to solve a number of problems efficiently (such as planar point location and triangulations among others) in parallel using a linear number of processors. This data-structure is similar to the **segment-tree** (proposed by J. Bentley and is also discussed in [18]), the intervals of which are induced by the projection of the endpoints of the input set of segments on one of the axes. This was used to emulate the plane-sweeping paradigm. The idea of using the plane sweep tree was later further developed by Atallah and Goodrich [3]. We shall review some definitions and results from their papers as they relate to our work.

3.1 Review of plane-sweep tree

Let T be a complete binary tree with its leaves corresponding to the $2e + 1$ intervals formed by projecting the $2e$ endpoints of a given set of e line segments onto the x -axis. Associated with each node v in the tree is an interval $[a_v, b_v]$ on the x axis corresponding to the union of intervals of its descendants. Let Π_v represent the vertical strip $[a_v, b_v] \times (-\infty, \infty)$. A segment s_i covers a node v

in T if its projection $X(s_i)$ on the x-axis spans the interval $[a_v, b_v]$ (corresponding to v) but does not span the interval of v 's parent node. It can be easily shown that no segment s_i covers more than 2 nodes at any level and hence no more than $O(\log n)$ nodes of the tree.

For each node v of T , we keep track of the the following properties:

$H(v) = \{s_i \mid s_i \text{ covers } v\}$,

$W(v) = \{s_i \mid s_i \text{ has at least one endpoint in } \Pi_v\}$.

$L(v) = \{s_i \mid s_i \in W(v) \text{ and } s_i \text{ intersects the left boundary of } \Pi_v\}$.

$R(v) = \{s_i \mid s_i \in W(v) \text{ and it intersects the right boundary}\}$.

$I(v) = \{s_i \mid \text{the left endpoint of } s_i \text{ is in } \Pi_{\text{leftchild}(v)} \text{ and the right endpoint is in } \Pi_{\text{rightchild}(v)}\}$.

If the given set of segments are non-intersecting (as in the case of a polygon), then these sets are completely ordered on y coordinate. The above quantities are computed while constructing the plane-sweep tree, which are used to search for segments directly above or below a given query point.

Multilocation : Given a plane-sweep tree, this procedure locates for each input point p , the segment in $H(v)$ which is directly above (or below) p for all vertices v in the tree such that p belongs to the interval Π_v .

Augmented Plane sweep tree : This is a data structure obtained from a plane sweep tree by adding pointers to every node such that given the position of an element in any node, its position in the parent node can be located in an additional constant time. (For readers familiar with fractional cascading, this data structure enables us to perform that.)

3.2 Development of the new data-structure

Fact 1 (Atallah and Goodrich [3]) Multilocation of any query point can be done in $O(\log n)$ sequential time in an augmented plane-sweep tree.

The preprocessing cost of building this data-structure was $O(\log^2 n)$ time using $O(n)$ processors and $O(n \log n)$ space. Consequently, the above-mentioned problems needed $\Omega(\log^2 n)$ time. Atallah and Goodrich [3] improved the preprocessing time for constructing the plane-sweep tree to $O(\log n \log \log n)$ using parallel merging techniques of Borodin and Hopcroft [4] while keeping the number of processors and the space requirement unchanged. The dependence of their approach on parallel merging can be summed up in the following manner :

Fact 2: The Plane-Sweep Tree of k segments can be constructed in $O(f(p,k) \log k)$ time using p processors ($p \geq k$), where $f(p,k)$ denotes the time complexity of merging two sorted lists of $O(k)$ elements using p processors.

Since $f(k,k) = \Theta(\log \log k)$ (see Valiant [23] and Borodin and Hopcroft [4]), the bound $O(\log n \log \log n)$ of Atallah and Goodrich's **Build-Up** algorithm for plane-sweep tree follows. Moreover, it enables us to state a useful lemma:

Lemma 2 *The Plane-Sweep tree of n segments can be constructed in $O(\log n)$ time with $O(n^{1+\epsilon})$ processors using Atallah and Goodrich's [3] approach.*

The proof follows from the well known result (Valiant [23] and Borodin and Hopcroft [4]) that two sorted lists of $O(n)$ elements can be merged in constant time using $n^{1+\epsilon}$ processors. Using $f(p,k) = O(1)$ in Fact 1 yields the required result.

An important common characteristic of Atallah and Goodrich's [3] algorithms is that the time complexity is dominated by the preprocessing cost. More specifically, given the precomputed data-structure, triangulation and planar-point location can be done in $O(\log n)$ time using n processors in a CREW model. Thus, it is worthwhile to try to improve the preprocessing time to $O(\log n)$ so that the overall running time of the algorithms can be reduced to $O(\log n)$. Our approach avoids merging two $O(n)$ size lists by using random sampling to build up a new data-structure called the **Nested-Plane-Sweep tree** in $O(\log n)$ time with very high probability. The overall running time of our algorithms is $\tilde{O}(\log n)$ with $O(n)$ processors.

We shall now give an outline of our method, which will be formalized thereafter. The techniques used are similar to the random sampling used in the Flashsort algorithm of Reif and Valiant [21]. The use of random-splitting is extended to two-dimensions. For this, we choose randomly a sample of \sqrt{n} segments from the given set of n segments. (We shall later see that, we choose only n^{ϵ_o} size sample, $0 < \epsilon_o < 1$ and the actual value of ϵ_o will be less than $1/13$. All the arguments in this section apply for any ϵ_o which has been set to $1/2$ for ease of presentation.) Every segment has an equal probability i.e. $\frac{1}{\sqrt{n}}$ of being selected. We build up the augmented plane-sweep tree of Atallah and Goodrich [3] on this sample of \sqrt{n} segments using $O(n)$ processors. The randomly chosen set of $O(\sqrt{n})$ segments partitions the plane into $O(\sqrt{n})$ disjoint regions (proof follows later). Each of the remaining $O(n)$ segments is located (using the multilocation algorithm of Atallah and Goodrich) in the $O(\sqrt{n})$ regions, in $O(\log n)$ time, using n processors. Though each segment may be divided into a number of smaller segments by the intervals induced by the plane-sweep tree of the sample set, the expected number of segments in each region will be shown to be $O(\sqrt{n})$ and less than $O(\sqrt{n} \log n)$ with very high probability. The algorithm can now be applied recursively to the sub-problems in each of the $O(\sqrt{n})$ sub-regions, resulting in a recurrence equation of the form $\bar{T}(n) = \bar{T}(\sqrt{n} \log n) + O(\log n)$ which has a solution $\bar{T}(n) = O(\log n)$. A step-wise description of the algorithm is given below.

Procedure Nested-Sweep-Tree

Input: Set S of non-intersecting line-segments l_1, l_2, \dots, l_n in a plane.

- (1) Choose a set of \sqrt{n} segments randomly from the original set. Each segment is selected with equal probability of $\frac{1}{\sqrt{n}}$.
- (2) Use the **Build-up** and **Augment** algorithms of Atallah and Goodrich [3] to construct the augmented plane sweep tree. Time = $O(\log n)$, Space = $O(\sqrt{n} \log n)$.
- (3) Locate the $n - \sqrt{n}$ segments in the plane sweep tree i.e. identify the region (the $O(\sqrt{n})$ regions) into which the sample set divides the plane) where each of the segments lie. Note that a segment may lie in more than one region in which case it is broken into a number of smaller segments each of which lies in only one region. This step uses Atallah and Goodrich's [3] *multilocation* algorithm. The segments which lie completely within a region can be located using the algorithm directly. However segments that cross the boundaries can be split into $O(\sqrt{n})$ broken segments (in the worst case) are located using a scheme described later.
- (4) If the number of segments in any region is more than a threshold, then apply procedure Nested-Sweep-Tree to each of these regions. (the threshold can be chosen to be $O(\log^r n)$ for some non-negative integer r).

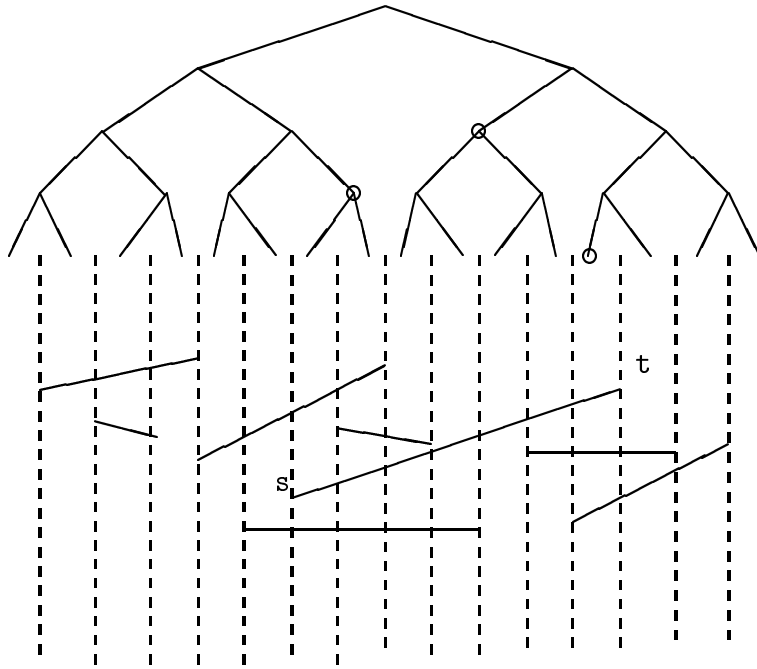


Figure 1: The skeleton of a plane-sweep tree. The circled nodes have portions of the segment $s-t$ (corresponding to the intervals covered by the nodes).

Step 3 needs further explanation. For this, we need a few probabilistic lemmas which bound the size of each subproblem and the total size of all the subproblems at any level of recursive call.

3.3 Random-sampling methods and probabilistic bounds

Lemma 3 *The Plane-Sweep tree of the random sample of \sqrt{n} segments divides the plane into less than or equal to $3\sqrt{n}$ trapezoidal regions.*

Proof : Assuming distinct endpoints there are $2\sqrt{n}$ vertical segments through the endpoints, each of which can be shared by 3 such trapezoidal regions (see Figure 1). Moreover each trapezoidal region is bounded by at most two vertical segments. Thus, there are at most $3\sqrt{n}$ such regions. \square

To bound the number of segments lying in a given trapezoidal region, we classify the segments into two groups:

- (a) Segments that have at most one of their endpoints inside the region.
- (b) Segments that lie completely within the region.

(a) The segments that lie partially in the region intersect either the left boundary, or the right boundary or both boundaries of the region. Consider an ordered list of the segments $L_U(i)$

intersecting the vertical segment passing through an end-point in the upward direction. These segments can be completely ordered w.r.t. the ordinate (on y). The probability that this ordered list exceeds m is less than $(1 - \frac{1}{\sqrt{n}})^m$ because each segment has probability $\frac{1}{\sqrt{n}}$ of being chosen in the sample. Let us denote by E_i the event that $m < k\sqrt{n}\log n$ (for some $k > 1$) for a fixed endpoint i . The probability that the event occurs for any end-point (and there are $2n$ such end-points) can be upper-bounded by $\sum_{i=1}^{2n} e^{-k \log n}$. For $k > 2$, the event that the ordered list of line segments $L_U(i)$ is less than $k\sqrt{n}\log n$ for all i (which is the complement of the previous event) holds with very high likelihood. Thus the number of segments intersecting any boundary is less than $O(\sqrt{n}\log n)$ with high likelihood (this is only a subset of the previous event).

Moreover, the expected number of line segments that intersect a boundary can be upper-bounded by \sqrt{n} (mean of geometric distribution). Note that the random variables representing the number of segments intersecting each of the regions are not independent. Since the expectation of the sum is the sum of expectation, the total number of such segments over all the regions is expected to be $< 12n$ (since a trapezoid has at most two vertical segments going up and two going down and there are $< 3\sqrt{n}$ trapezoids). It follows that, the probability that the total number of lines intersecting all the trapezoids exceeds $k_{max}n$ ($k_{max} > 24$) is less than $1/2$. This implies that by repeating the experiment of choosing \sqrt{n} segments $\log n$ times, the probability of failure (where a success is the event that the total number of intersections is less than $k_{max}n$) is less than $\frac{1}{n^c}$.

If we choose independently $p(n) = O(\log n)$ sets of samples, one of them is good with very high likelihood. However, to determine if a sample is ‘good’, we would have to carry out step 3, $O(\log n)$ times each of which requires $O(\log n)$ time (such a method is described in section 3.4). Instead, we try to estimate the the number of segments intersecting a trapezoid T_i using only a fraction of the input segments. For example, we can choose $c_0 \cdot n/\log^d n$ (for some fixed integer $d > 2$ and a constant c_0 (which will be determined from the required success probability of the algorithm) of the input segments randomly for the j th sample, R_j . Let X_i^j be the number of segments intersecting trapezoid T_i corresponding to sample R_j , $1 \leq j \leq b \log n$ (b is fixed integer greater than 0) and A_i^j be the number of segments intersecting T_i out of the $n/\log^d n$ randomly chosen input segments for the same sample. Clearly, A_i^j is a binomial random variable with parameters $c_0 \cdot n/\log^d n$, X_i^j/n . Assuming that X_i^j is greater than $\bar{c} \cdot \log^{d+1} n$, for some constant \bar{c} , we can apply Chernoff bounds (see appendix) to tightly bound the estimates within a constant multiplicative factor. Since we do it only for $1/\log^d n$ of the input segments, the total work done for the $O(\log n)$ random subsets does not change the total work done (which we shall show to be $O(n \log n)$ in the next section). (Note that $X_i^j < \bar{c} \log^d n$, it is an easy case since $\sqrt{n} \cdot p \log^d n = o(n)$).

More formally, by invoking Chernoff bounds (see Appendix equations (1) and (2)), for any $\alpha > 0$ (α is a function of c_0), there exists a c_1 , independent of n ,

$$\text{Prob}(A_i^j \leq \alpha c_1 X_i^j / \log^d n) \leq 1/n^\alpha \text{ and}$$

$$\text{Prob}(A_i^j \geq c_2 \alpha c_0 \cdot X_i^j / \log^d n) < 1/n^{c_0 \alpha} < 1/n^\alpha \text{ (for } c_0 > 1).$$

From the last two inequalities, X_i^j is bounded by $L^j = A_i^j \log^d n / c_0 c_2 \alpha$ from below, and $U^j = A_i^j \log^d n / c_1 \alpha$ from above. With appropriate changes in the constants, this condition holds with high likelihood (as defined in section 2.1) for all X_i^j simultaneously. We do the procedure

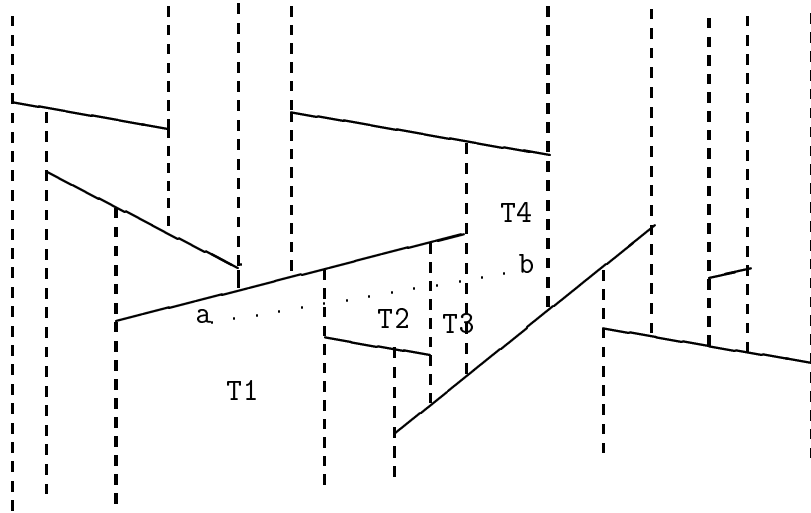


Figure 2: Segment a-b is multilocated in the trapezoidal regions labeled T1, T2, T3, T4.

(described in the next section) simultaneously for all the samples R_j and choose the sample R_{j_0} using the following simple test:

Algorithm *Sample-select*

(Let $N^j = \sum A_i^j$ and let the actual number of intersections be denoted by T^j and the upper and lower bounds obtained from N^j by U^j and L^j respectively).

If $k_{total}n > U^j$ then accept sample R^j (since $k_{total}n \geq U^j \geq T^j$), else

if $k_{total}n \leq L^j$ then the sample is 'bad' (since $k_{total}n \leq L^j \leq T^j$), else

if $L^j \leq k_{total}n \leq U^j$, then accept the sample R^{j_0} for which E^{j_0} is minimum. Since both $k_{total}n$ and T^{j_0} lie in this interval this guarantees that $T^{j_0} \leq c_3 \cdot k_{total}n$ where $c_3 = U^j/L^j$ which is a constant.

Recall, that from our earlier discussion at least one of the samples would satisfy conditions 1 or 3 with very high likelihood.

(b) The $O(\sqrt{n})$ sub-divisions are trapezoidal (see Figure 2) regions which can be specified by at most four segments. Thus, there are $O(n^4)$ potential trapezoids of which only $O(\sqrt{n})$ were chosen in the random sample (Lemma 3). We shall call a trapezoid "good" if it has less than $O(\sqrt{n} \log n)$

segments within it and "bad" otherwise. Since the probability that any chosen trapezoid containing more than m segments is less than $(1 - \frac{1}{\sqrt{n}})^m$, the bounds of (a) hold. Thus all the chosen trapezoids are "good" with very high probability (the number of potential trapezoids from \sqrt{n} segments is less than n^2). The total number of segments lying within all trapezoids is obviously less than n (from the input size).

We can summarize the above observations in the following lemma:

Lemma 4 *There exist constants α, \bar{c} such that the probability of any one of the $O(\sqrt{n})$ trapezoidal regions containing more than $O(\bar{c}\alpha\sqrt{n}\log n)$ segments (or more appropriately broken segments) is less than $n^{-\alpha}$. Moreover, the total number of such segments over all trapezoidal regions is less than $k_{max}n$ with very high likelihood.*

The success of the recursive calls at any level can be argued in an identical manner except that we allow the probability of success at level i to diminish to $1 - n^{-1/2^{i-1}\alpha}$. For example, in level i , we do the resampling only $\log n/2^i$ times.

3.4 Partitioning segments into trapezoidal regions

We shall use a *locus-based* approach to solve this problem. This approach involves considering each query as a higher dimensional point and partitioning the underlying space into regions providing the same answer. Thus the query problem is reduced to a point location problem, given sufficient preprocessing time and space. The problem at hand involves preprocessing the trapezoidal subdivisions (induced by the sample) in such a manner that given the end-points of any segment we should be able to list the regions it intersects in $O(\log n)$ time using $\lceil k/\log n \rceil$ processors where k is the number of regions that it intersects. We shall show that the preprocessing for n segments can be done in $O(\log n)$ time using $O(n^c)$ processors, where c is a fixed constant. Thus we can choose any sample of size less than $n^{1/c}$.

Since the input segments are non-intersecting, these can only extend between regions where there exists a 'clear-path' which does not intersect any other sample segment. It must also be clear that there can exist more than such 'clear-path' between two regions. Our objective is to partition the plane into regions (equivalent classes), such that given any fixed pair of such regions (where the end-points of a segment lies), the regions that the segment intersects can be pre-determined. The boundaries of these 'clear-paths' are straight-lines joining vertices of the trapezoidal subdivisions and the equivalent regions are intersections of the half-spaces defined by these boundaries. Since we need a fast preprocessing procedure, we shall settle for a finer partition of space (i.e. more than one pair of partitioned region may intersect exactly the same set of trapezoids). If n is the size of the trapezoidal regions, there are $O(n)$ vertices which gives rise to at most $O(n^2)$ boundary conditions which are straight lines joining every pair of vertices. Some of them may intersect sampled segments and hence are not boundaries of 'clear-paths' but it doesn't affect our procedure since they would only make the partition more fine (See Figure 3 for an illustration). These $O(n^2)$ lines can intersect in $O(n^4)$ vertices, giving rise to a $O(n^4)$ regions in the partition. This implies that there are $O(n^8)$ possible pairs of regions where end-points of a segment could lie and one can pre-compute for each such pair which trapezoids the corresponding segment intersects using $O(n^9)$ processors in $O(\log n)$ time.

For the point location problem, we use a pre-processing scheme similar to Dobkin and Lip-ton [11]. The following result is a straightforward consequence of their paper

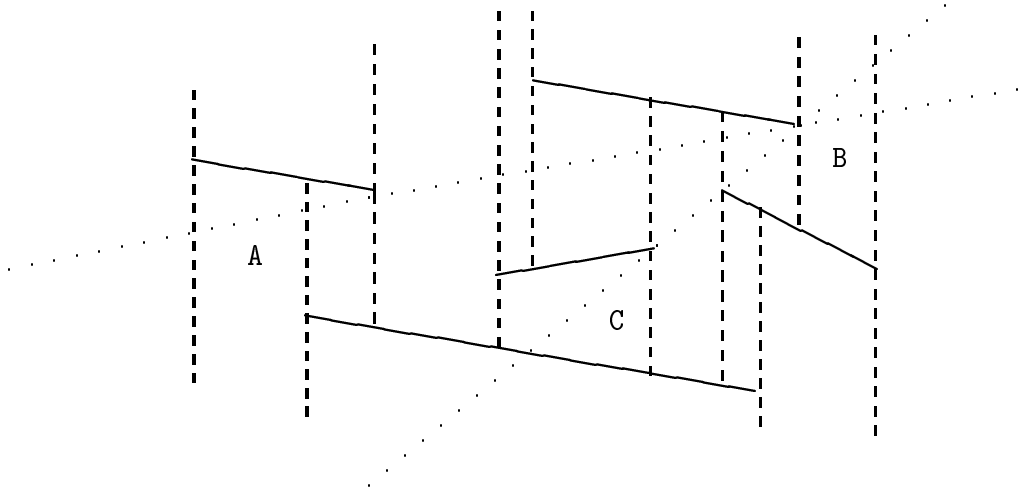


Figure 3: There is a clear path between regions A and B where as there cannot be a segment with end-points in regions C and B. This redundant partitioning does not affect the asymptotic complexity of the algorithm.

Lemma 5 *Given a set of m line segments in a plane, it is possible to preprocess them in $O(\log m)$ time time using $O(m^3)$ processors such that point-location for any query point can be done in $O(\log m)$ time. The space needed is $O(m^3)$.*

Proof: We merely mimic the sequential algorithm. After computing all possible pairs of intersections in $O(1)$ time using $O(m^2)$ processors, we project the intersection on the x-axis and for each interval we compute the total ordering of the lines in $O(\log m)$ time using m processors for each of the intervals. Thus the space required is $O(m^3)$. \square

Because of this pre-processing there is an increase in the number of regions from what we mentioned in the previous paragraph. Each region is now a trapezoid (can also be a triangle) and there are $O(n^6)$ regions. Accordingly we require $O(n^{13})$ processors to pre-compute the possible intersections. Since each region is a trapezoid, we choose a sample point in each trapezoid to do the pre-computing part. We make a table corresponding to each pair of regions containing the trapezoids the corresponding segment intersects. For any input segment, we first do the point location for its end-points and then perform another search on this ordered pair of regions. Clearly the whole procedure is a constant number of binary searches and takes $O(\log n)$ time. We can also store the number of intersecting trapezoids for each entry, so that we can allocate the required number of processors corresponding to each segment to list the intersecting trapezoids (using a prefix sum).

Thus we can state the main result of this section as follows :

Theorem 2 *Algorithm Nested-Plane-Sweep-tree runs in $\tilde{O}(\log n)$ time and $O(n \log \log n)$ space using*

$O(n)$ processors in a CREW PRAM model.

Proof : Since steps 1, 2 and 3 run in $O(\log n)$ time, we have the following recurrence equation for $\bar{T}(n)$, the expected time complexity of the algorithm :

$$\bar{T}(n) = O(\log n) + (1 - n^{-\alpha})\bar{T}(n^{1-\epsilon} \log n) + n^{-\alpha}\bar{T}(n - n^\epsilon)$$

The solution of this recurrence equation gives $\bar{T}(n) = O(\log n) + \text{low order terms}$. From our previous discussion $\epsilon \geq 1/13$.

At this point we have shown that with sufficient number of processors, the algorithm executes in expected $O(\log n)$ time. Since the number of (broken) segments can increase by a constant factor (k_{max}), this could increase the input size by a polylogarithmic factor over $O(\log \log n)$ levels. Consequently, the processor bounds would have to increase by the same factor (from being linear) to achieve the time bound of $O(\log n)$. To avoid this, we make the following modification in the algorithm. After partitioning the segments into the trapezoidal regions (using the procedure in section 3.4), we group the segments into two categories:

- (a) Segments (part-segments) that have at least one end-point in the region
- (b) Segments that span the region (horizontally)

Notice that number of segments of type (a) is less than $2n$ and the segments of type (b) in a region i , S^i can be completely ordered (with respect to y -coordinate). While performing *multilocation*, a straight-forward binary search suffices for ordered segments in S_i . Consequently, we do not further pre-process the segments in S_i , i.e. we can leave them out from further recursive calls. Thus, the total size of the subproblems at any level of the recursive call is no more than $2n$. Processor allocation is achieved by simply setting processors in a region equal to the number of end-points lying in it. This shows that the algorithm does not need more than $O(n)$ processors.

To analyze the cumulative effects of the nested calls on the worst case total runtime of the algorithm, we need to consider only one such sequence of nested calls. This gives the probability that a leaf-node process fails to complete within $O(\log n)$ time. Since the success of the algorithm depends on completion of all such processes (at the leaf level), the probability that the algorithm fails to halt in $O(\log n)$ time is less than $n \cdot \text{Prob}[\text{one such process fails}]$. Note that this part of the analysis is very similar to *Flashsort*

From lemma 4, we know that for any given level i , the time T_i can be bounded by $P[T_i \geq c_o \log n (\epsilon_o)^i] \leq 2^{-(\epsilon_o)^i \log n \alpha c_o}$ where $\epsilon_o < 12/13$, since we choose only $n^{1/13}$ sample segments. Setting $t_i = k(\epsilon_o)^i \log n \alpha (c - c_o)$, where c_o is some constant, we obtain

$$\text{Prob}[T_i \geq k\alpha c \log n (\epsilon_o)^i + t_i] \leq 2^{-(\epsilon_o)^i \alpha \log n} \leq 2^{-t_i/k}$$

If T is the total time for this worst case chain of nested calls and $m = 1/(1-\epsilon_o)$, the probability that it takes more than $mk\alpha \log n c_o + t$, is less than the sum of the probability of events where $\sum_i t_i = t + j$, $0 \leq j \leq \mu$. Here $\mu = mk\alpha \log n c_o$. We shall compute the probability that $\sum_i t_i = t$ and multiply by μ .

$$\prod_{\sum t_i = t} 2^{-t_i} \leq \sum 2^{-t/k} \text{ over } t^{O(\log \log n)} \text{ tuples.}$$

$$\text{Thus Prob}[T > km\alpha \log n c_o + t] < \mu 2^{-t/k + O(\log t \log \log^2 n)}$$

Using $t \geq km\alpha(c - c_o) \log n$, for large values of n and $m > 1$, we can rewrite the above expression as

$$\text{Prob}[T > km\alpha c \log n] < \mu 2^{-\alpha m(c - c_o) \log n}$$

For $c > 4c_o$, i.e. $c - c_o > 3/4c$, we have the following required bound,

$$\text{Prob}[T > a \log n] \leq \mu 2^{-(3/4)c\alpha \log n} \leq n^{-c_1\alpha}.$$

assuming that k , m and c are larger than 1. \square

4 Applications Of Nested-Plane-Sweep Tree

In this section, we exploit the data structure that we constructed in the previous section to improve the running time of various algorithms like triangulation, intersection detection, 3-D maxima, 2-D dominance counting, and visibility from a point.

Lemma 6 *The nested-plane-sweep-tree constructed in the previous section can be used to multilocate any query point p in $\tilde{O}(\log n)$ sequential time.*

Proof: From Fact 1 multilocation in an $O(n_i)$ node plane-sweep tree can be performed in $O(\log n_i)$ serial time. Thus, the expected time for multilocation in the nested-plane sweep tree obeys the following recurrence relation:

$$\bar{T}(n) = O(\log n) + (1 - n^{-\beta})\bar{T}(\sqrt{n} \log n) + (n^{-\beta})\bar{T}(n)$$

giving $\bar{T}(n) = O(\log n)$ which is the multilocation time for each query point p using one processor. Equivalently, multilocation for $O(n)$ query points can be performed in $O(\log n)$ expected time using $O(n)$ processors. The worst case time bounds can be derived using techniques similar to the proof of Theorem 2. \square

4.1 Trapezoidal decomposition and triangulation

Definition *it Trapezoidal Decomposition* Let $P = \{v_1, v_2, \dots, v_n\}$ be a simple polygon, where v_i 's denote the vertices of P , and are listed so that the interior of P is to the left of the walk $v_1 v_2 \dots v_n$. For any vertex v_i of P a trapezoidal edge for v_i is an edge of P , which is directly above or below v_i , such that the vertical line segment from v_i to this edge is interior to P . A vertex can have 0,1 or 2 such edges. Trapezoidal decomposition of a polygon P is to find the trapezoidal edges for each vertex of P .

Lemma 7 *Given a simple polygon P , a trapezoidal decomposition of P can be constructed in $\tilde{O}(\log n)$ time using $O(n)$ processors and $O(n \log n)$ space in a CREW PRAM model.*

Proof : The trapezoidal decomposition can be done in two distinct stages (see Atallah and Goodrich [3] for details). In the preprocessing part, a nested plane-sweep tree is constructed on the edges of the simple polygon P . From theorem 2 this runs in $O(\log n)$ time with very high probability. Subsequently, all the vertices of the polygon are multilocated simultaneously using

$O(n)$ processors which takes $O(\log n)$ time with very high likelihood (from lemma 6). For each point, it takes a constant time to determine if the vertical line intersecting the trapezoidal edge is within the polygon P using one processor per point. \square

Fact 3 (Atallah and Goodrich [3]): Given a one-sided monotone polygon with n vertices, P can be triangulated in $O(\log n)$ time and $O(n)$ space using $O(n)$ processors in a CREW PRAM model. Our next theorem is a direct consequence of this fact and Lemma 7.

Theorem 3 *Given a simple polygon with n vertices, P can be triangulated in $\tilde{O}(\log n)$ time and $O(n \log n)$ space using $O(n)$ processors in a CREW PRAM model.*

Proof: The algorithm consists of three phases:

(1) We first build the Nested Plane-Sweep-Tree on the edges of the polygon. With very high probability this can be done in $O(\log n)$ time and $O(n \log n)$ space (from theorem 2).

(2) The simple polygon is decomposed into one-sided monotone polygons by using trapezoidal decomposition (see Atallah and Goodrich [3] for details). Thus, from lemma 7 the expected running time for this phase is $O(\log n)$.

(3) Each of the one-sided monotone polygons can be triangulated in $O(\log n)$ time and $O(n)$ space from fact 3.

The proof of the theorem follows directly from the above steps. \square

4.2 Visibility from a point

Definition *Visibility from a point* Given a set of line segments $S = \{s_1, s_2 \dots s_n\}$, which do not intersect, except possibly at end-points, and a point p , determine the part of the plane which is visible from p when every segment s_i is opaque.

We shall assume the point p to be at negative infinity below all segments. The algorithm that we present below can appropriately modified for any general point. Notice that now (with the assumption) the problem is to determine the projections of segments on the x -axis, such that in case of overlap the segment closer to x axis has precedence (see Figure 4).

The solution to this problem consists of a sequence of points $\{p_1, p_2 \dots p_{2n}\}$ such that two consecutive points p_i, p_{i+1} define a segment visible in the interval $[x(p_i), x(p_{i+1})]$. An important property of this problem is that the visibility is a continuous function between the endpoints of the segments, i.e. for all points between two endpoints (projection of the endpoints on the x axis), the same segment will be visible. We can refer to the endpoints as *critical* points where the visibility function defined as $\text{Vis}(x)$ may change its value. This property enables us to solve the problem efficiently. Since $\text{Vis}(x)$ is constant between any two consecutive endpoints, we choose a point p which lies in this interval below all the segments and draw a vertical line through p . The segment which intersects the vertical line first (from below) is the visible segment in the corresponding interval. Formally, the algorithm can be described as follows:

Algorithm *Visibility*

(1) Given a set S of non-intersecting segments, sort their endpoints on the x -coordinate. This can be done in $O(\log n)$ time using Cole's [8] parallel mergesort algorithm which does not have a

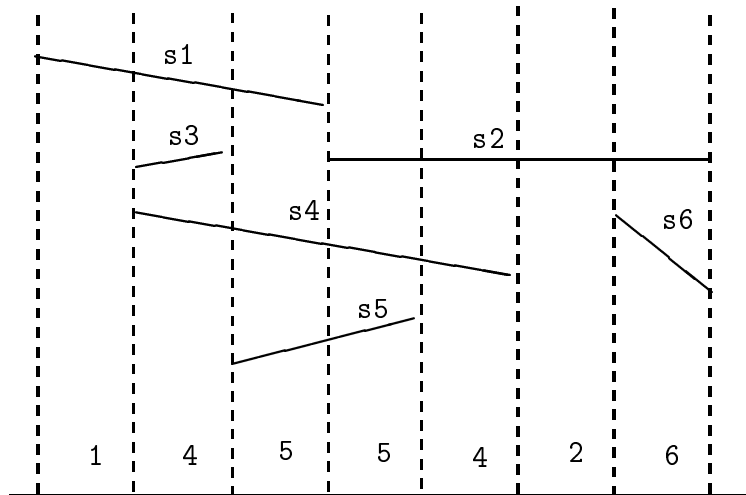


Figure 4: The intervals on the x-axis are labeled by the segment visible in that interval.

large constant as does the AKS algorithm.

(2) For each of the $2n-1$ bounded intervals, choose a point within the interval. A possible choice is the midpoint of an interval. Let us denote them by $\{p_1, p_2, \dots, p_{2n-1}\}$.

(3) Construct a Nested Plane-Sweep tree on the segments of set S .

(4) Multilocate the $2n-1$ midpoints on this tree giving us ordered pairs of the form (p_i, s_j) which carries the required visibility information for the interval $[x_i, x_{i+1}]$.

The following theorem is a direct consequence of our previous discussion:

Theorem 4 *Given a set S of segments and a point p , the visibility of the plane induced by this set of segments from this point can be computed in $\tilde{O}(\log n)$ time using $O(n)$ processors in a CREW PRAM.*

5 Dominance Problems

The problems tackled in the previous section used trapezoidal decomposition which can be efficiently carried out with the nested-plane sweep tree. A plane sweep tree was constructed on a random sample, and was used to divide and conquer on the plane which was divided into trapezoidal regions. An important property which was not mentioned explicitly was that by dividing up the plane in such a manner we did not require a merging phase. For trapezoidal decomposition, locating a point within a particular trapezium at any phase eliminated interaction with the other regions since the trapezoidal edge could not be exterior to this region. The problems dealt in this section are of somewhat different nature, and there may arise a need to determine the relative location of a query point with respect to more than one trapezoidal regions. This distinction can be brought out more clearly by the following example : Given a set of non-intersecting segments and a query point, p ,

we wish to determine the following:

- (1) Which segment is immediately above p ?
- (2) Among all the segments above p which one is the longest?

The first problem can be solved by trapezoidal decomposition which can be done efficiently using the Nested-Plane-Sweep tree. However, the second problem appears more difficult and indeed it is because it cannot be solved by simply finding the segment immediately above the point p within one region. It will be shown that the latter problem can be solved easily using the normal plane-sweep tree but the construction of this tree is the bottleneck i.e. it takes $O(\log n \log \log n)$ time using $O(n)$ processors. To avoid building the full data structure we shall argue that we do not need all the attributes (mentioned in section 3.1) in the plane-sweep tree and also avoid the need for fractional cascading in the algorithms presented in this section.

5.1 Maximal set in three-dimensions

The **Maxima** problem is defined as follows : Given a set S of points in d -dimensions, the maxima of S is the set of all points p_i , such that $p_i \in S$, and is not dominated on all dimensions by any other point in set S .

Efficient sequential algorithms are known for only two and three dimensions ($O(n \log n)$ time algorithm is optimal). For two dimensions, an $O(\log n)$ algorithm using $O(n)$ processors is easily obtainable by using the $O(\log n)$ AKS sorting network or Cole's [8] parallel mergesort (which has a much lower constant). The best known result for 3-D maxima is $O(\log n \log \log n)$ from Atallah and Goodrich's algorithm ([3]) in which they use parallel merging techniques. We present a method to reduce the 3-D maxima problem to integer sorting so that we can solve it in $O(\log n)$ time using $O(n)$ processors.

Given a set S of points in three dimensions, consider their projection on the x - y plane. For any point (x_i, y_i) draw the segment s_i parallel to x -axis joining $(0, y_i)$ and (x_i, y_i) . Note that any point (x_j, y_j) is dominated by another point (x_i, y_i) iff the point lies below s_i and also $z_j \leq z_i$. This means that the point is dominated by p_i on all the axes. Instead of comparing the z coordinates of all s_i which are above p_j , it is enough to compare z_j with the maximum of the z coordinate for all the segments s_i which dominate p_j (Figure 5). Note that identifying the segments which are above p_j is similar to the multilocation of p_j . However, this time we are not only interested in the segment which lie directly above p_j but also which of them has the maximum z -coordinate and this is unique for any interval. It is possible to precompute the maximum z coordinate for each interval so that the necessary comparisons can be performed in constant additional time during each step of multilocation. This preprocessing requires a parallel-prefix type of computation. The following well known result can be used to do this efficiently:

Fact 4: Parallel prefix computation for n element sequence can be done in $O(\log n)$ time using $O(n/\log n)$ processors (for example see Reif [20]).

The sequence of elements in each node are segments "covering" the corresponding interval of the plane-sweep-tree sorted on y axes and the operation needed is **MAX**. (Recall the definition of 'covering' from section 3.1)

However, as mentioned previously, we cannot afford to construct the full data structure of [3]

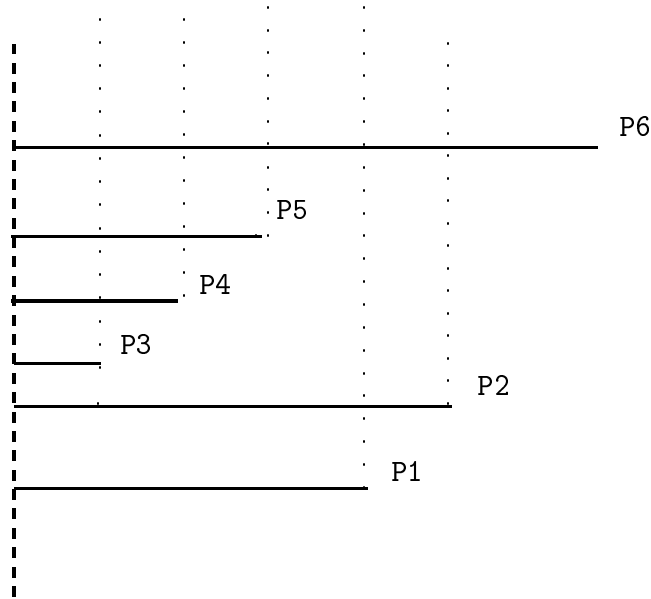


Figure 5: Point P1 is not a maxima iff it is dominated by P2 or P6 in the z-coordinate.

and therefore the following observations can be made :

Observation 1: We do not require fractional cascading to perform multilocation of the input points. We can instead use their ranks in the list $H(v)$. Note that fractional cascading gives us more information for multilocating those points that are not in the input set (used for building the tree).

Observation 2: We can build the sets $H(v)$ in $O(\log n)$ time using the following result:

Fact 5: (Rajasekaran [19]) Integer sort of n keys in the range $[1, n^{O(1)}]$ can be performed in time $O(\log n)$ using $n/\log n$ processors in a CREW PRAM model given word size of n^ϵ bits for any $\epsilon > 0$.

We first build the skeleton of the plane sweep tree on the intervals induced by the end-points of the segments (corresponding to the input points). Note that the segments can be totally ordered on the y coordinates and after an initial sorting on the y-coordinate, we can make use of their ranks (which is an integer in the interval $[1, n]$) instead of the actual y coordinate. Using one processor per segment and concurrent read, we can find the allocation nodes for each segment - there can be at most $2 \cdot \log n$ for each. Due to the special nature of the line segments (all have the same left x coordinate), it can be seen that the allocation node cannot be a right child. We also make a slight modification - we not only allocate segments to the appropriate intervals but also to all

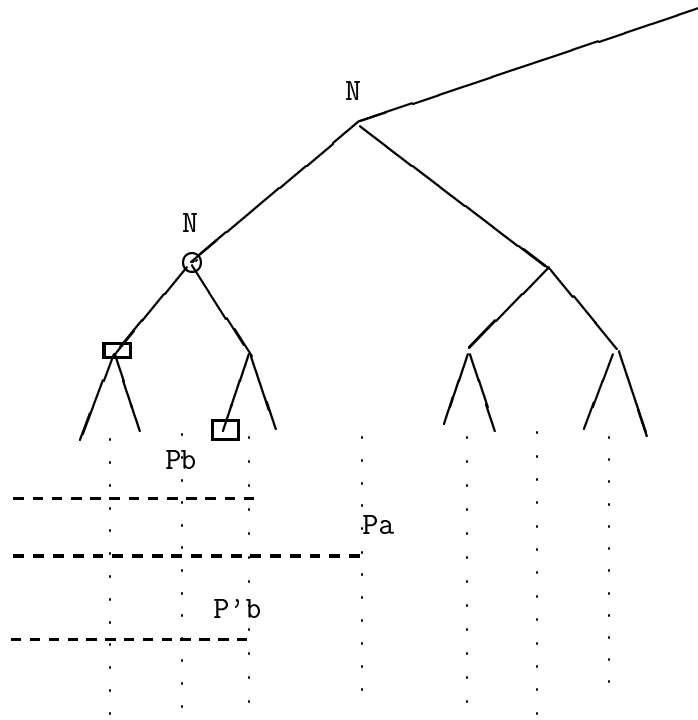


Figure 6: P_a is allocated to the circled node and the square nodes store P_b and P'_b . Nodes marked 'N' are the special nodes of allocation for P_b which are the left-children on the allocation path of P_b .

the left childs in the **allocation path** (see Figure 6 for an illustration of such nodes). These are special allocation nodes of a segment so we distinguish them by some special markings. Note that there can be at most $\log n$ such special allocation nodes for each segment. We can next build the unordered lists for $H(v)$ ($H(v)$ is ordered on y axis) for each node v of the plane-sweep tree using lexicographic sorting and prefix computation on the ordered pairs (v, s_i) . Though there can be $O(n \log n)$ such pairs, this can still be done in $O(\log n)$ time using n processors using Facts 4 and 5. Another application of sorting (on the ranks of the y coordinate) and prefix computation gives us the $H(v)$ in the same time bounds.

The full algorithm is presented below :

Procedure 3-D Maxima

it Input n points in 3-dimensions, where each point p_i is defined by its 3-coordinates (x_i, y_i, z_i) .
 (1) Build a plane-sweep tree on the segments joining (x_i, y_i) and $(0, y_i)$, and compute the $H(v)$'s using the scheme described in observation 2. At the end of this step, we shall have for each segment, its rank in all the $H(v)$'s (at most $2 \log n$) which will be used for multilocation.

(2) For each node n_i , calculate $\text{Max}(s_{k,n_i}) = \underset{z\text{-coord}}{\text{max}} \{s_{1,n_i}, s_{2,n_i}, \dots, s_{k,n_i}\}$ for $k \leq |H(n_i)|$,

where $s_{1,n_i}, s_{2,n_i}, \dots, s_{|H(n_i)|,n_i}$ are segments in node n_i ordered on y coordinate (which is the set $H(n_i)$ from its definition.) During this step, the marked entries (which are to be used for multilo-
cation) are assigned y-coordinates of $-\infty$ so that they do not affect the computation.

(3) At each node, in the path followed by a point p_i from the root to the right most allocation node of s_i , compare $z(p_i)$ with the $\text{Max}(s_{k,n_i})$, where p_i lies immediately below s_{k,n_i} using the rank of s_i in the corresponding $H(v)$. If at any time during the search $z(p_i)$ is less than this value, p_i is not a Maxima of S . (Note that this step can be done in $O(\log n)$ time by using one processor per point.)

end

Theorem 5 *Algorithm 3-D Maxima finds the maximal points correctly in $O(\log n)$ time and $O(n \log n)$ space using $O(n)$ processors in a CREW PRAM model.*

Proof : The proof of the time complexity is based on the previous discussion. To see that it identifies the maximal points correctly, we have to explain significance of the "marked" nodes in more detail. Given points p_a and p_b , where p_a dominates p_b , we have to ensure that they are allocated to at least one common node, so that in step 3, we are able to eliminate p_b from the set of maximal elements. This may not happen in the normal allocation scheme (as shown in Figure 6), hence there is a need for special nodes. The assignment of $-\infty$ to the special nodes in step 2, prevents p_b from dominating p_a if $y(p_b) > y(p_a)$ and $z(p_b) > z(p_a)$, but $x(p_b) < x(p_a)$. \square

5.2 Two-set dominance on a plane

The **two-set dominance counting** problem is defined as follows: Given a set $V = \{p_1, p_2, \dots, p_l\}$ and a set $U = \{q_1, q_2, \dots, q_m\}$ of points in the plane, we wish to determine the number of points in V which are dominated (on two coordinates) by each point q_i in U . This seems to be closely related to the two-dimensional dominance problem but is harder since not only we have to identify points that are not maximum but also find *all* points which dominate a given point. However this problem can also be solved using a scheme similar to the previous problem.

Given the set U and V , we transform each point $q_i \in U$, with coordinates (x_i, y_i) , into a segment s_i joining (x_i, y_i) and $(0, y_i)$ (similar to the previous problem) and build a plane-sweep tree on these segments. Then, in parallel, allocate each point p_i of set V on this tree to the appropriate intervals (all the nodes in the path which are left children in the tree). These are marked as special elements to be used for dominance counting. Next, we construct the sets $H(v)$'s using the original segments as well as the points of V . We mark the elements corresponding to p_i and q_i with 1 and 0 respectively. In each node, the number of special points in $H(v)$ (the points of V) lying under a segment s_i is the number of points dominated by s_i in that interval. Thus, we can carry out a prefix sum computation using the reverse order of the $H(v)$'s. To find the total number of points dominated by q_i , we simply add up sequentially the numbers over all the intervals (at most $\log n$) in which the segment is allocated. A formal description of the algorithm is given below

Two-set dominance counting

- (1) Build a Plane-Sweep tree on the segments joining (x_i, y_i) and $(0, y_i)$ where (x_i, y_i) correspond to coordinates of a point q_i in set U .
- (2) Locate each point p_i of set V simultaneously on the plane-sweep tree constructed in step 1 and allocate it to all the nodes (intervals) which are the left children in the path taken by the point in the tree. (Note that p_i is not stored as a segment but just as a point.)
- (3) Mark the points of V and construct $H(v)$ by lexicographic sorting on the pairs $(v, y(p_i))$ using the integer sorting algorithm in [19].
- (4) For each node v of the tree perform a parallel prefix sum on the the set $H(v)$ to count the number of points of V lying below a q_i . This gives us the total number of points dominated by a segment (and hence the corresponding point) in a certain interval.
- (5) For any segment find the total number of points dominated by all the intervals of the segment. This can also be done using one processor per q_i , since there can be at most $\log n$ intervals for each segment s_i .

Theorem 6 *Algorithm Two-set dominance counting runs correctly in $O(\log n)$ time using $O(n)$ processors in a CREW model where $n = |U| + |V|$.*

Proof : The proof of correctness of the algorithm is similar to 3-D dominance where we use some special allocation nodes. To ensure a correct count of the number of points in set V dominated by q_k , we have to count each point p_i (dominated by q_k) exactly once. Thus q_k and p_i should share exactly one node of allocation. The special allocation nodes guarantee that there will be at least one shared node. The argument that there is at most one such node follows from the way that p_i and q_i are stored in the tree. All the nodes where p_i 's are allocated are in a path (and hence are related as ancestors), where as the q_i are stored (as transformed segments) in nodes which cannot be ancestors of each other. The justification for time complexity follows easily from the preceding discussion since each of the steps can be completed in $O(\log n)$ time. \square

Two-set dominance counting can be used to solve a number of problems efficiently, among which the multiple range-counting problem is very important. Multiple range counting problem can be described as following : Given a set V of l points and a set R of m rectangles (ranges), the multiple range-counting problem is to compute the number of points interior to each rectangle. Note that this is equivalent to a data base query where the ranges are defined by two different keys. The next result follows immediately.

Corollary 3 Given a set V of l points in a plane, and a set R of m isothetic rectangles, we can solve the multiple range-counting problem for V and R in $O(\log n)$ time with very high probability using $O(n)$ processors where $n = l + m$.

Proof : We can transform this problem into two-dimensional dominance counting problem as follows: Given a rectangle r defined by its four corners (p_1, p_2, p_3, p_4) the number of points enclosed by a rectangle is equal to $d(p_1) - d(p_2) - d(p_3) + d(p_4)$. Here p_1 is the upper right corner of the rectangle, p_4 the lower right corner and $d(v)$ denotes the number of points in V two-dominated (dominated on both coordinates) by v . The result follows. \square

6 Concluding remarks

In this paper we have designed parallel algorithms for solving a variety of problems in computational geometry which have optimal running time with high probability. Some of these algorithms use a new data structure, which we call Nested-Plane-Sweep tree. This appears to be an useful modification of the plane-sweep tree described in Aggarwal et al. [1] and Atallah and Goodrich [3]. On the other hand, plane-sweep tree itself can be used to solve a number of problems optimally if it can be constructed optimally. Random splitting was used very effectively to divide and conquer on the plane, raising hopes about extending these techniques efficiently to problems with higher dimensions like the three-dimensional convex hulls. A challenging exercise will be mapping these algorithms to fixed interconnection networks like hypercube and maintain the same performance.

As mentioned in the introduction, Atallah, Cole and Goodrich [2] have independently discovered optimal deterministic algorithms for all these problems. Some of their algorithms use the less powerful EREW model and their algorithms and they do not use n^ϵ size words that we need for integer-sorting. Since they are based on an approach used by Cole for the optimal parallel mergesort algorithm (see [8]), there is little hope of extending them to the fixed interconnection models while maintaining the optimal performance achieved in the PRAM models.

Moreover, as demonstrated by Clarkson [7], randomized methods have very wide applications in computational geometry in terms of simplifying deterministic algorithms or speeding up sub-optimal algorithms. Though his methods were restricted to a sequential setting, this paper provides ample evidence that they can be extended to the parallel setting with some additional techniques (as seen in section 3.4). In addition, the algorithms can be made to run in a certain time (and number of processors) with very high likelihood instead of just a constant probability. There are still a number of important problems in computational geometry which do not have optimal parallel algorithms and we feel that this paper has merely scratched the surface.

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A Appendix

We say a random variable X upper-bounds another random variable Y (equivalently Y lower bounds X) if for all x such that $0 \leq x \leq 1$, $\text{Prob}(X \leq x) \leq \text{Prob}(Y \leq x)$.

A Bernoulli trial is an experiment with two possible outcomes viz. success and failure. The probability of success is p.

A binomial variable X with parameters (n,p) is the number of successes in n independent Bernoulli trials, the probability of success in each trial being p. The *probability mass function* of X can be easily seen to be

$$\text{Prob}(X \leq x) = \sum_{k=0}^x \binom{n}{k} p^k (1-p)^{n-k}$$

The tail end of the Binomial distribution can be bounded by *Chernoff* bounds. In particular the following approximations due to Angluin and Valiant are frequently used:

$$\text{Prob}(X \geq m) \leq \left(\frac{np}{m}\right)^m e^{m-np} \quad (1)$$

$$\text{Prob}(X \leq m) \leq \left(\frac{np}{m}\right)^m e^{-np+m} \quad (2)$$

$$\text{Prob}(X \leq (1-\epsilon)np) \leq \exp(-\epsilon^2 np/2) \quad (3)$$

$$\text{Prob}(X \geq (1+\epsilon)np) \leq \exp(-\epsilon^2 np/3) \quad (4)$$

for all $0 < \epsilon < 1$. The last two bounds actually follow from the Chernoff bounds which (for a discrete distribution) can be stated as

$$\text{Prob}[A \geq x] \leq z^{-x} G_A(z) \text{ where } G_A(z) \text{ is the probability generating function.}$$

To minimize the bound we substitute $z = z_0$ that minimizes the right side expression.

The probability mass function of a modified geometric random variable X is specified by $p_X(i) = p(1-p)^i$ for $i = 0, 1, 2, \dots$. The probability generating function of a modified geometric random variable X is given by $G_Y(z) = \frac{p}{1-(1-p)z}$ where p is the probability of success in each individual trial. Recall that, Y is the number of trials before we have a success. Thus the generating function of sum of logn independent identical modified geometric random variable is given by

$$G_Y(z) \left(\frac{p}{1-(1-p)z}\right)^{\log n} G_Y(z) \left(\frac{p}{1-(1-p)z}\right)^{\log n}$$

Using Chernoff bounds and minimizing the bounds by differentiating we obtain

$$\text{Prob}[Y \geq c \log n] \leq n^{-c \log_2 \frac{2c}{c+1}} \text{ for } p = 1/2$$

For $c > 2$, we can write it as $\text{Prob}[Y \geq c \log n] \leq n^{-\alpha c}$ where $\alpha > 0$. (5)