CMSC 754
Computational Geometry

David M. Mount
Department of Computer Science
University of Maryland
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1 Introduction to Computational Geometry

What is Computational Geometry? Computational geometry is a term claimed by a number of different groups. The term was coined perhaps first by Marvin Minsky in his book “Perceptrons”, which was about pattern recognition, and it has also been used often to describe algorithms for manipulating curves and surfaces in solid modeling. Its most widely recognized use, however, is to describe the subfield of algorithm theory that involves the design and analysis of efficient algorithms for problems involving geometric input and output.

The field of computational geometry developed rapidly in the late 70’s and through the 80’s and 90’s, and it still continues to develop. Historically, computational geometry developed as a generalization of the study of algorithms for sorting and searching in 1-dimensional space to problems involving multi-dimensional inputs. Because of its history, the field of computational geometry has focused mostly on problems in 2-dimensional space and to a lesser extent in 3-dimensional space. When problems are considered in multi-dimensional spaces, it is usually assumed that the dimension of the space is a small constant (say, 10 or lower). Nonetheless, recent work in this area has considered a limited set of problems in very high dimensional spaces.

Because the field was developed by researchers whose training was in discrete algorithms (as opposed to numerical analysis) the field has also focused more on the discrete nature of geometric problems, as opposed to continuous issues. The field primarily deals with straight or flat objects (lines, line segments, polygons, planes, and polyhedra) or simple curved objects such as circles. This is in contrast, say, to fields such as solid modeling, which focus on issues involving curves and surfaces and their representations.

A Typical Problem in Computational Geometry: Here is an example of a typical problem, called the shortest path problem. Given a set polygonal obstacles in the plane, find the shortest obstacle-avoiding path from some given start point to a given goal point. Although it is possible to reduce this to a shortest path problem on a graph (called the visibility graph, which we will discuss later this semester), and then apply a nongeometric algorithm such as Dijkstra’s algorithm, it seems that by solving the problem in its geometric domain it should be possible to devise more efficient solutions. This is one of the main reasons for the growth of interest in geometric algorithms.

The measure of the quality of an algorithm in computational geometry has traditionally been its asymptotic worst-case running time. Thus, an algorithm running in $O(n)$ time is better than one running in $O(n \log n)$ time which is better than one running in $O(n^2)$ time. (This particular problem can be solved in $O(n^2 \log n)$ time by a fairly simple algorithm, in $O(n \log n)$ by a relatively complex algorithm, and it can be approximated quite well by an algorithm whose running time is $O(n \log n)$.) In some cases average case running time is considered instead. However, for many types of geometric
inputs (this one for example) it is difficult to define input distributions that are both easy to analyze and representative of typical inputs.

There are many fields of computer science that deal with solving problems of a geometric nature. These include computer graphics, computer vision and image processing, robotics, computer-aided design and manufacturing, computational fluid-dynamics, and geographic information systems, to name a few. One of the goals of computational geometry is to provide the basic geometric tools needed from which application areas can then build their programs. There has been significant progress made towards this goal, but it is still far from being fully realized.

**Strengths Computational Geometry:**

**Development of Geometric Tools:** Prior to computational geometry, there were many ad hoc solutions to geometric computational problems, some efficient, some inefficient, and some simply incorrect. Because of its emphasis of mathematical rigor, computational geometry has made great strides in establishing correct, provably efficient algorithmic solutions to many of these problems.

**Emphasis on Provable Efficiency:** Prior to the development of computational geometry little was understood about the computational complexity of many geometric computations. For example, given an encoding of all the zip code regions in the USA, and given a latitude and longitude from a GPS device, how long should it take to compute the zip code associated with the location? How should the computation time depend on the amount of preprocessing time and space available? Computational geometry put such questions on the firm grounding of asymptotic complexity, and in some cases it has been possible to prove that algorithms discovered in this area are optimal solutions.

**Emphasis on Correctness/Robustness:** Prior to the development of computational geometry, many of the software systems that were developed were troubled by bugs arising from the confluence of the continuous nature of geometry and the discrete nature of computation. For example, given two line segments in the plane, do they intersect? This problem is remarkably tricky to solve since two line segments may arise from many different configurations: lying on parallel lines, lying on the same line, touching end-to-end, touching as in a T-junction. Software that is based on discrete decisions involving millions of such intersection tests may very well fail if any one of these tests is computed erroneously. Computational geometry research has put the robust and correct computing of geometric primitives on a solid mathematical foundations.

**Linkage to Discrete Combinatorial Geometry:** The study of new solutions to computational problems has given rise to many new problems in the mathematical field of discrete combinatorial geometry. For example, consider a polygon bounded by \( n \) sides in the plane. Such a polygon might be thought of as the top-down view of the walls in an art gallery. As a function of \( n \), how many “guarding points” suffice so that every point within the polygon can be seen by at least one of these guards. Such combinatorial questions can have profound implications on the complexity of algorithms.

**Limitations of Computational Geometry:**

**Emphasis on discrete geometry:** There are some fairly natural reasons why computational geometry may never fully address the needs of all these applications areas, and these limitations
should be understood before undertaking this course. One is the discrete nature of computational geometry. There are many applications in which objects are of a very continuous nature: computational physics, computational fluid dynamics, motion planning.

**Emphasis on flat objects:** Another limitation is the fact that computational geometry deals primarily with straight or flat objects. To a large extent, this is a consequence of CG’ers interest in discrete geometric complexity, as opposed to continuous mathematics. Another issues is that proving the correctness and efficiency of an algorithm is only possible when all the computations are well defined. Many computations on continuous objects (e.g., solving differential and integral equations) cannot guarantee that their results are correct nor that they converge in specified amount of time. Note that it is possible to approximate curved objects with piecewise planar polygons or polyhedra. This assumption has freed computational geometry to deal with the combinatorial elements of most of the problems, as opposed to dealing with numerical issues.

**Emphasis on low-dimensional spaces:** One more limitation is that computational geometry has focused primarily on 2-dimensional problems, and 3-dimensional problems to a limited extent. The nice thing about 2-dimensional problems is that they are easy to visualize and easy to understand. But many of the daunting applications problems reside in 3-dimensional and higher dimensional spaces. Furthermore, issues related to topology are much cleaner in 2- and 3-dimensional spaces than in higher dimensional spaces.

**Overview of the Semester:** Here are some of the topics that we will discuss this semester.

**Convex Hulls:** Convexity is a very important geometric property. A geometric set is convex if for every two points in the set, the line segment joining them is also in the set. One of the first problems identified in the field of computational geometry is that of computing the smallest convex shape, called the convex hull, that encloses a set of points.

![Convex hull](image1.png) ![Polygon triangulation](image2.png)

**Intersections:** One of the most basic geometric problems is that of determining when two sets of objects intersect one another. Determining whether complex objects intersect often reduces to determining which individual pairs of primitive entities (e.g., line segments) intersect. We will discuss efficient algorithms for computing the intersections of a set of line segments.

**Triangulation and Partitioning:** Triangulation is a catchword for the more general problem of subdividing a complex domain into a disjoint collection of “simple” objects. The simplest region into which one can decompose a planar object is a triangle (a tetrahedron in 3-d and simplex in general). We will discuss how to subdivide a polygon into triangles and later in the semester discuss more general subdivisions into trapezoids.
Low-dimensional Linear Programming: Many optimization problems in computational geometry can be stated in the form of a linear programming problem, namely, find the extreme points (e.g. highest or lowest) that satisfies a collection of linear inequalities. Linear programming is an important problem in the combinatorial optimization, and people often need to solve such problems in hundred to perhaps thousand dimensional spaces. However there are many interesting problems (e.g. find the smallest disc enclosing a set of points) that can be posed as low dimensional linear programming problems. In low-dimensional spaces, very simple efficient solutions exist.

Voronoi Diagrams and Delaunay Triangulations: Given a set $S$ of points in space, one of the most important problems is the nearest neighbor problem. Given a point that is not in $S$ which point of $S$ is closest to it? One of the techniques used for solving this problem is to subdivide space into regions, according to which point is closest. This gives rise to a geometric partition of space called a Voronoi diagram. This geometric structure arises in many applications of geometry. The dual structure, called a Delaunay triangulation also has many interesting properties.

Line Arrangements and Duality: Perhaps one of the most important mathematical structures in computational geometry is that of an arrangement of lines (or generally the arrangement of curves and surfaces). Given $n$ lines in the plane, an arrangement is just the graph formed by considering the intersection points as vertices and line segments joining them as edges. We will show that such a structure can be constructed in $O(n^2)$ time.

The reason that this structure is so important is that many problems involving points can be transformed into problems involving lines by a method of duality. For example, suppose that you want to determine whether any three points of a planar point set are collinear. This could be determines in $O(n^3)$ time by brute-force checking of each triple. However, if the points are dualized into lines, then (as we will see later this semester) this reduces to the question of whether there is a vertex of degree greater than 4 in the arrangement.
Search:  Geometric search problems are of the following general form. Given a data set (e.g. points, lines, polygons) which will not change, preprocess this data set into a data structure so that some type of query can be answered as efficiently as possible. For example, a nearest neighbor search query is: determine the point of the data set that is closest to a given query point. A range query is: determine the set of points (or count the number of points) from the data set that lie within a given region. The region may be a rectangle, disc, or polygonal shape, like a triangle.

Approximation:  In many real-world applications geometric inputs are subject to measurement error. In such cases it may not be necessary to compute results exactly, since the input data itself is not exact. Often the ability to produce an approximately correct solution leads to much simpler and faster algorithmic solutions. Consider for example the problem of computing the diameter (that is, the maximum pairwise distance) among a set of $n$ points in space. In the plane efficient solutions are known for this problem. In higher dimensions it is quite hard to solve this problem exactly in much less than the brute-force time of $O(n^2)$. It is easy to construct input instances in which many pairs of points are very close to the diametrical distance. Suppose however that you are willing to settle for an approximation, say a pair of points at distance at least $(1 - \varepsilon)\Delta$, where $\Delta$ is the diameter and $\varepsilon > 0$ is an approximation parameter set by the user. There exist algorithms whose running time is nearly linear in $n$, assuming that $\varepsilon$ is a fixed constant. As $\varepsilon$ approaches zero, the running time increases.
2 Geometric Basics

This material will not be covered in lecture, but is presented here just in case you are interested in refreshing your knowledge on how basic geometric entities are represented and manipulated.

Geometry Basics: As we go through the semester, we will introduce much of the geometric facts and computational primitives that we will be needing. For the most part, we will assume that any geometric primitive involving a constant number of elements of constant complexity can be computed in $O(1)$ time, and we will not concern ourselves with how this computation is done. (For example, given three non-collinear points in the plane, compute the unique circle passing through these points.) Nonetheless, for a bit of completeness, let us begin with a quick review of the basic elements of affine and Euclidean geometry.

There are a number of different geometric systems that can be used to express geometric algorithms: affine geometry, Euclidean geometry, and projective geometry, for example. This semester we will be working almost exclusively with affine and Euclidean geometry. Before getting to Euclidean geometry we will first define a somewhat more basic geometry called affine geometry. Later we will add one operation, called an inner product, which extends affine geometry to Euclidean geometry.

Affine Geometry: An affine geometry consists of a set of scalars (the real numbers), a set of points, and a set of free vectors (or simply vectors). Points are used to specify position. Free vectors are used to specify direction and magnitude, but have no fixed position in space. (This is in contrast to linear algebra where there is no real distinction between points and vectors. However this distinction is useful, since the two are conceptually quite different.)

The following are the operations that can be performed on scalars, points, and vectors. Vector operations are just the familiar ones from linear algebra. It is possible to subtract two points. The difference $p - q$ of two points results in a free vector directed from $q$ to $p$. It is also possible to add a point to a vector. In point-vector addition $p + v$ results in the point which is translated by $v$ from $p$. Letting $S$ denote an generic scalar, $V$ a generic vector and $P$ a generic point, the following are the legal operations in affine geometry:

$$
\begin{align*}
S \cdot V & \rightarrow V \quad \text{scalar-vector multiplication} \\
V + V & \rightarrow V \quad \text{vector addition} \\
P - P & \rightarrow V \quad \text{point subtraction} \\
P + V & \rightarrow P \quad \text{point-vector addition}
\end{align*}
$$

Figure 2.1: Affine operations.
A number of operations can be derived from these. For example, we can define the subtraction of two vectors \( \vec{u} - \vec{v} \) as \( \vec{u} + (-1) \cdot \vec{v} \) or scalar-vector division \( \vec{v}/\alpha \) as \( (1/\alpha) \cdot \vec{v} \) provided \( \alpha \neq 0 \). There is one special vector, called the zero vector, \( \vec{0} \), which has no magnitude, such that \( \vec{v} + \vec{0} = \vec{v} \).

Note that it is not possible to multiply a point times a scalar or to add two points together. However there is a special operation that combines these two elements, called an affine combination. Given two points \( p_0 \) and \( p_1 \) and two scalars \( \alpha_0 \) and \( \alpha_1 \), such that \( \alpha_0 + \alpha_1 = 1 \), we define the affine combination

\[
\text{aff}(p_0, p_1; \alpha_0, \alpha_1) = \alpha_0 p_0 + \alpha_1 p_1 = p_0 + \alpha_1 (p_1 - p_0).
\]

Note that the middle term of the above equation is not legal given our list of operations. But this is how the affine combination is typically expressed, namely as the weighted average of two points. The right-hand side (which is easily seen to be algebraically equivalent) is legal. An important observation is that, if \( p_0 \neq p_1 \), then the point \( \text{aff}(p_0, p_1; \alpha_0, \alpha_1) \) lies on the line joining \( p_0 \) and \( p_1 \). As \( \alpha_1 \) varies from \(-\infty\) to \(+\infty\) it traces out all the points on this line.

\[
\begin{align*}
\text{aff}(p, q; 3/2, -1/2) \\
\text{aff}(p, q; 1/2, 1/2) \\
\text{aff}(p, q; 1, 0) \\
\text{aff}(p, q; 0, 1)
\end{align*}
\]

Figure 2.2: Affine combination.

In the special case where \( 0 \leq \alpha_0, \alpha_1 \leq 1 \), \( \text{aff}(p_0, p_1; \alpha_0, \alpha_1) \) is a point that subdivides the line segment \( \overline{p_0p_1} \) into two subsegments of relative sizes \( \alpha_1 \) to \( \alpha_0 \). The resulting operation is called a convex combination, and the set of all convex combinations traces out the line segment \( \overline{p_0p_1} \).

It is easy to extend both types of combinations to more than two points, by adding the condition that the sum \( \alpha_0 + \alpha_1 + \alpha_2 = 1 \).

\[
\text{aff}(p_0, p_1, p_2; \alpha_0, \alpha_1, \alpha_2) = \alpha_0 p_0 + \alpha_1 p_1 + \alpha_2 p_2 = p_0 + \alpha_1 (p_1 - p_0) + \alpha_2 (p_2 - p_0).
\]

The set of all affine combinations of three (non-collinear) points generates a plane. The set of all convex combinations of three points generates all the points of the triangle defined by the points. These shapes are called the affine span or affine closure, and convex closure of the points, respectively.

**Euclidean Geometry:** In affine geometry we have provided no way to talk about angles or distances. Euclidean geometry is an extension of affine geometry which includes one additional operation, called the inner product, which maps two real vectors (not points) into a nonnegative real. One important example of an inner product is the dot product, defined as follows. Suppose that the \( d \)-dimensional vectors \( \vec{u} \) and \( \vec{v} \) are represented by the (nonhomogeneous) coordinate vectors \((u_1, u_2, \ldots, u_d)\) and \((v_1, v_2, \ldots, v_d)\). Define

\[
\vec{u} \cdot \vec{v} = \sum_{i=1}^{d} u_i v_i.
\]

The dot product is useful in computing the following entities.

**Length:** of a vector \( \vec{v} \) is defined to be \( \|\vec{v}\| = \sqrt{\vec{v} \cdot \vec{v}} \).
Normalization: Given any nonzero vector $\vec{v}$, define the normalization to be a vector of unit length that points in the same direction as $\vec{v}$. We will denote this by $\hat{\vec{v}}$:

$$\hat{\vec{v}} = \frac{\vec{v}}{\|\vec{v}\|}.$$ 

Distance between points: Denoted either $\text{dist}(p, q)$ or $\|pq\|$ is the length of the vector between them, $\|p - q\|$.

Angle: between two nonzero vectors $\vec{u}$ and $\vec{v}$ (ranging from 0 to $\pi$) is

$$\text{ang}(\vec{u}, \vec{v}) = \cos^{-1}\left(\frac{\vec{u} \cdot \vec{v}}{\|\vec{u}\|\|\vec{v}\|}\right) = \cos^{-1}(\hat{\vec{u}} \cdot \hat{\vec{v}}).$$

This is easy to derive from the law of cosines.

Orientation of Points: In order to make discrete decisions, we would like a geometric operation that operates on points in a manner that is analogous to the relational operations ($<$, $=$, $>$) with numbers. There does not seem to be any natural intrinsic way to compare two points in $d$-dimensional space, but there is a natural relation between ordered $(d + 1)$-tuples of points in $d$-space, which extends the notion of binary relations in 1-space, called orientation.

Given an ordered triple of points $(p, q, r)$ in the plane, we say that they have positive orientation if they define a counterclockwise oriented triangle, negative orientation if they define a clockwise oriented triangle, and zero orientation if they are collinear (which includes as well the case where two or more of the points are identical). Note that orientation depends on the order in which the points are given.

![Figure 2.3: Orientations of the ordered triple $(p, q, r)$.](image)

Orientation is formally defined as the sign of the determinant of the points given in homogeneous coordinates, that is, by prepending a 1 to each coordinate. For example, in the plane, we define

$$\text{Orient}(p, q, r) = \det \begin{pmatrix} 1 & p_x & p_y \\ 1 & q_x & q_y \\ 1 & r_x & r_y \end{pmatrix}.$$ 

Observe that in the 1-dimensional case, Orient$(p, q)$ is just $q - p$. Hence it is positive if $p < q$, zero if $p = q$, and negative if $p > q$. Thus orientation generalizes $<$, $=$, $>$ in 1-dimensional space. Also note that the sign of the orientation of an ordered triple is unchanged if the points are translated, rotated, or scaled (by a positive scale factor). A reflection transformation, e.g., $f(x, y) = (-x, y)$, reverses the sign of the orientation. In general, applying any affine transformation to the point alters the sign of the orientation according to the sign of the matrix used in the transformation.

This generalizes readily to higher dimensions. For example, given an ordered 4-tuple points in 3-space, we can define their orientation as being either positive (forming a right-handed screw), negative
(a left-handed screw), or zero (coplanar). It can be computed as the sign of the determinant of an appropriate $4 \times 4$ generalization of the above determinant. This can be generalized to any ordered $(d + 1)$-tuple of points in $d$-space.

**Areas and Angles:** The orientation determinant, together with the Euclidean norm can be used to compute angles in the plane. This determinant $\text{Orient}(p, q, r)$ is equal to twice the signed area of the triangle $\triangle pqr$ (positive if CCW and negative otherwise). Thus the area of the triangle can be determined by dividing this quantity by 2. In general in dimension $d$ the area of the simplex spanned by $d + 1$ points can be determined by taking this determinant and dividing by $d! = d \cdot (d - 1) \cdots 2 \cdot 1$. Given the capability to compute the area of any triangle (or simplex in higher dimensions), it is possible to compute the volume of any polygon (or polyhedron), given an appropriate subdivision into these basic elements. (Such a subdivision does not need to be disjoint. The simplest methods that I know of use a subdivision into overlapping positively and negatively oriented shapes, such that the signed contribution of the volumes of regions outside the object cancel each other out.)

Recall that the dot product returns the cosine of an angle. However, this is not helpful for distinguishing positive from negative angles. The sine of the angle $\theta = \angle pqr$ (the signed angled from vector $p - q$ to vector $r - q$) can be computed as

$$\sin \theta = \frac{\text{Orient}(q, p, r)}{\|p - q\| \cdot \|r - q\|}.$$  

(Notice the order of the parameters.) By knowing both the sine and cosine of an angle we can unambiguously determine the angle.
3 Convex Hulls

**Reading:** Chapter 1 in the 4M’s (de Berg, van Kreveld, Overmars, Schwarzkopf). The divide-and-conquer and Jarvis march algorithms are given in Joseph O’Rourke’s, “Computational Geometry in C.” O’Rourke’s book is also a good source for information about orientations and some other geometric primitives.

**Convexity:** Next, we consider a fundamental structure in computational geometry, called the **convex hull**. We will give a more formal definition later, but the convex hull can be defined intuitively by surrounding a collection of points with a rubber band and then letting the rubber band “snap” tightly around the points.

![Figure 3.1: A point set and its convex hull.](image)

There are a number of reasons that the convex hull of a point set is an important geometric structure. One is that it is one of the simplest shape approximations for a set of points. (Other examples include minimum area enclosing rectangles, circles, and ellipses.) It can also be used for approximating more complex shapes. For example, the convex hull of a polygon in the plane or polyhedron in 3-space is the convex hull of its vertices.

Also many algorithms compute the convex hull as an initial stage in their execution or to filter out irrelevant points. For example, the **diameter** of a point set is the maximum distance between any two points of the set. It can be shown that the pair of points determining the diameter are both vertices of the convex hull. Also observe that minimum enclosing convex shapes (such as the minimum area rectangle, circle, and ellipse) depend only on the points of the convex hull.

**Convexity:** A set $S$ is **convex** if given any points $p, q \in S$ any convex combination of $p$ and $q$ is in $S$, or equivalently, the line segment $\overline{pq} \subseteq S$.

**Convex hull:** The **convex hull** of any set $S$ is the intersection of all convex sets that contains $S$, or more intuitively, the smallest convex set that contains $S$. We will denote this $\text{conv}(S)$.

Recall from the lecture on geometric basics that given two points $p$ and $q$, a **convex combination** is any point that can be expressed as $(1-\alpha)p + \alpha q$, for any scalar $\alpha$, where $0 \leq \alpha \leq 1$. More generally, given any finite subset of points $\{p_1, p_2, \ldots, p_m\}$ a convex combination is any point that can be expressed as a weighted sum $\sum_{i=1}^{m} \alpha_i p_i$, where $0 \leq \alpha_i \leq 1$ and $\sum_{i=1}^{m} \alpha_i = 1$. An equivalent definition of convex hull is the set of points that can be expressed as convex combinations of the points in $S$. (A proof can be found in any book on convexity theory.)

**Some Terminology:** Although we will not discuss topology with any degree of formalism, we will need to use some terminology from topology. These terms deserve formal definitions, but we are going to cheat and rely on intuitive definitions, which will suffice for the simple, well behaved geometric objects that we will be dealing with. Beware that these definitions are not fully general, and you are referred to a good text on topology for formal definitions.
For our purposes, for $r > 0$, define the $r$-neighborhood of a point $p$ to be the set of points whose distance to $p$ is strictly less than $r$, that is, it is the set of points lying within an open ball of radius $r$ centered about $p$. Given a set $S$, a point $p$ is an interior point of $S$ if for some radius $r$ the neighborhood about $p$ of radius $r$ is contained within $S$. A point is an exterior point if it lies in the interior of the complement of $S$. A point that is neither interior nor exterior is a boundary point. A set is open if it contains none of its boundary points and closed if its complement is open. If $p$ is in $S$ but is not an interior point, we will call it a boundary point.

We say that a geometric set is bounded if it can be enclosed in a ball of finite radius. A set is compact if it is both closed and bounded.

In general, convex sets may have either straight or curved boundaries and may be bounded or unbounded. Convex sets may be topologically open or closed. Some examples are shown in the figure below. The convex hull of a finite set of points in the plane is a bounded, closed, convex polygon.

![Diagram of convex hulls]

**Figure 3.2:** Terminology.

**Convex Hull Problem:** The (planar) convex hull problem is, given a set of $n$ points $P$ in the plane, output a representation of $P$’s convex hull. The convex hull is a closed convex polygon, the simplest representation is a counterclockwise enumeration of the vertices of the convex hull. Ideally, the hull should consist only of extreme points, in the sense that if three points lie on an edge of the boundary of the convex hull, then the middle point should not be output as part of the hull. (If we assume that the points are in general position, and in particular no three are collinear, then this is not an issue.)

There is a simple $O(n^3)$ convex hull algorithm, which operates by considering each ordered pair of points $(p, q)$, and the determining whether all the remaining points of the set lie within the half-plane lying to the right of the directed line from $p$ to $q$. (Observe that this can be tested using the orientation test.) The question is, can we do better?

**Graham’s scan:** We will present an $O(n \log n)$ algorithm for convex hulls. It is a simple variation of a famous algorithm for convex hulls, called Graham’s scan. This algorithm dates back to the early 70’s. The algorithm is loosely based on a common approach for building geometric structures called incremental construction. In such a algorithm object (points here) are added one at a time, and the structure (convex hull here) is updated with each new insertion.

An important issue with incremental algorithms is the order of insertion. If we were to add points in some arbitrary order, we would need some method of testing whether the newly added point is inside the existing hull. It will simplify matters to add points in some appropriately sorted order, in our case, in increasing order of $x$-coordinate. This guarantees that each newly added point is outside the current hull. (Note that Graham’s original algorithm sorted points in a different way. It found the lowest point in the data set and then sorted points cyclically around this point. Sorting by $x$-coordinate seems to be a bit easier to implement, however.)
Since we are working from left to right, it would be convenient if the convex hull vertices were also
ordered from left to right. The convex hull is a cyclically ordered sets. Cyclically ordered sets
are somewhat messier to work with than simple linearly ordered sets, so we will break the hull into
two hulls, an upper hull and lower hull. (See Fig. 3.3(a).) The break points common to both hulls will
be the leftmost and rightmost vertices of the convex hull. After building both, the two hulls can be
concatenated into a single cyclic counterclockwise list.

As is common throughout the semester, we will make the assumption that points are in general posi-
tion, which in this case means that no two points have the same x-coordinates and no three points are
collinear. (You might consider what changes would need to be made to the algorithm to handle these
special cases.)

Here is a brief presentation of the algorithm for computing the upper hull. We will store the hull
vertices in a stack $H$, where the top of the stack corresponds to the most recently added point. Let
$H$.first and $H$.second denote the top and second element from the top of $H$, respectively. Observe
that as we read the stack elements from top to bottom (that is, from right to left) consecutive triples of
points of the upper hull will make a (strict) left-hand turn, that is, they should have a positive (that is,
clockwise) orientation. (See Fig. 3.3(b).)

Graham’s scan works as follows. Let $p_i$ denote the next point to be added in the left-to-right ordering
of the points. If the triple $\langle p_i, H$.first, $H$.second $\rangle$ is positively oriented, then we can simply push $p_i$
onto the stack (see Fig. 3.3(c)). Otherwise, we can infer that the middle point of the triple $H$.first
cannot be on the upper hull, and so we pop it off the stack. We repeat this until reaching a positively
oriented triple (see Fig. 3.3(d)), or there are fewer than two elements on the stack.

Our algorithm makes use of the following orientation primitive, which was discussed in the course
introduction.

$$\text{Orient}(p, q, r) \begin{cases} 
> 0 & \text{if } \langle p, q, r \rangle \text{ is oriented counterclockwise} \\
= 0 & \text{if } \langle p, q, r \rangle \text{ are collinear} \\
< 0 & \text{if } \langle p, q, r \rangle \text{ is oriented clockwise}
\end{cases}$$

Why is the algorithm correct? We can show inductively that the contents of $H$ at any stage of the
algorithm constitute the upper hull of the points that have been processed so far. For the induction
basis ($H = \{p_1, p_2\}$) this is trivially true. For the induction step, observe that $p_i$ is the rightmost
point among the points processed so far, and therefore it must lie on the upper hull. Let $p_j$ be the
neighboring vertex to $p_i$ on the upper hull of the first $i$ points. (See Fig. 3.3(c).) It is easy to see that
(1) Sort the points according to increasing order of their \(x\)-coordinates, denoted \(p_1, p_2, \ldots, p_n\).

(2) Push \(p_1\) and then \(p_2\) onto \(H\).

(3) for \(i = 3\) to \(n\) do:
   
   (a) while \((\text{size}(H) \geq 2 \text{ and } \text{Orient}(p_i, H.\text{first}, H.\text{second}) \leq 0)\) pop \(H\).
   
   (b) Push \(p_i\) onto \(H\).

\(p_j\) must be in \(H\) prior to the addition of \(p_i\). Each point \(p_k\) in \(H\) that lies between \(p_j\) and \(p_i\) lies beneath the edge \(p_jp_i\), and so \(p_k\) should not be part of the upper hull after \(p_i\) is added. For each such point it is easy to see that \(\text{Orient}(p_i, p_k, p_j) \leq 0\). Thus, as each of these points \(p_k\) is tested within the while loop, it will be deleted. Finally, when \(p_j\) reaches the top of the stack either find that \(p_j = p_1\), and hence there are less than two elements on the stack, or we find that we finally have the triple that satisfies the orientation test. In either case, the loop terminates and \(p_i\) is pushed on the stack, as desired.

The lower hull can be computed by an essentially symmetric algorithm, but working from right to left instead. Once the two hulls are computed, we simply concatenate the two hulls into a single circular list.

**Analysis:** Let us prove the main result about the running time of Graham’s scan.

**Theorem:** Graham’s scan runs in \(O(n \log n)\) time.

**Proof:** Sorting the points according to \(x\)-coordinates can be done by any efficient sorting algorithm in \(O(n \log n)\) time. Let \(d_i\) denote the number of points that are popped (deleted) on processing \(p_i\). Because each orientation test takes \(O(1)\) time, the amount of time spent processing \(p_i\) is \(O(d_i + 1)\). (The extra +1 is for the last point tested, which is not deleted.) Thus, the total running time is proportional to

\[
\sum_{i=1}^{n} (d_i + 1) = n + \sum_{i=1}^{n} d_i.
\]

To bound \(\sum_i d_i\), observe that each of the \(n\) points is pushed onto the stack once. Once a point is deleted it can never be deleted again. Since each of \(n\) points can be deleted at most once, \(\sum_i d_i \leq n\). Thus after sorting, the total running time is \(O(n)\). Since this is true for the lower hull as well, the total time is \(O(2n) = O(n)\).

**Convex Hull by Divide-and-Conquer:** As with sorting, there are many different approaches to solving the convex hull problem for a planar point set \(P\). Next we will consider another \(O(n \log n)\) algorithm, which is based on the divide-and-conquer design technique. It can be viewed as a generalization of the famous MergeSort sorting algorithm (see any standard algorithms text). Here is an outline of the algorithm. It begins by sorting the points by their \(x\)-coordinate, in \(O(n \log n)\) time. The remainder of the algorithm is shown in the code section below.

The asymptotic running time of the algorithm can be expressed by a recurrence. Given an input of size \(n\), consider the time needed to perform all the parts of the procedure, ignoring the recursive calls. This includes the time to partition the point set, compute the two tangents, and return the final result. Clearly the first and third of these steps can be performed in \(O(n)\) time, assuming a linked list
1. If \(|P| \leq 3\), then compute the convex hull by brute force in \(O(1)\) time and return.

2. Otherwise, partition the point set \(P\) into two sets \(A\) and \(B\), where \(A\) consists of half the points with the lowest \(x\)-coordinates and \(B\) consists of half of the points with the highest \(x\)-coordinates.

3. Recursively compute \(H_A = \text{conv}(A)\) and \(H_B = \text{conv}(B)\).

4. Merge the two hulls into a common convex hull, \(H\), by computing the upper and lower tangents for \(H_A\) and \(H_B\) and discarding all the points lying between these two tangents.

![Figure 3.4: Computing the lower tangent.](image)

representation of the hull vertices. Below we will show that the tangents can be computed in \(O(n)\) time. Thus, ignoring constant factors, we can describe the running time by the following recurrence.

\[
T(n) = \begin{cases} 
1 & \text{if } n \leq 3 \\
1 + 2T(n/2) & \text{otherwise.}
\end{cases}
\]

This is the same recurrence that arises in Mergesort. It is easy to show that it solves to \(T(n) \in O(n \log n)\) (see any standard algorithms text). All that remains is showing how to compute the two tangents.

One thing that simplifies the process of computing the tangents is that the two point sets \(A\) and \(B\) are separated from each other by a vertical line (assuming no duplicate \(x\)-coordinates). Let’s concentrate on the lower tangent, since the upper tangent is symmetric. The algorithm operates by a simple “walking” procedure. We initialize \(a\) to be the rightmost point of \(H_A\) and \(b\) is the leftmost point of \(H_B\). (These can be found in linear time.) Lower tangency is a condition that can be tested locally by an orientation test involving the two vertices and neighboring vertices on the hull. We iterate the following two loops, which march \(a\) and \(b\) down, until they reach the points lower tangency. Given a point \(a\) on the hull, let \(a.\text{suc}\) and \(a.\text{pred}\) denote its successor and predecessor in CCW order about the hull.

The condition “\(ab\) is not the lower tangent of \(H_A\)” can be implemented with the orientation test \(\text{Orient}(b, a, a.\text{pred}) \geq 0\), and the other test for \(H_B\) is analogous. Proving the correctness of this procedure is a little tricky, but not too hard. (The issue is proving that the two inner while loops never go beyond the lower tangent points.) See O’Rourke’s book out for a careful proof. The important thing is that each vertex on each hull can be visited at most once by the search, and hence its running time is \(O(m)\), where \(m = |H_A| + |H_B| \leq |A| + |B|\). This is exactly what we needed to get the overall \(O(n \log n)\) running time.

QuickHull: If the divide-and-conquer algorithm can be viewed as a sort of generalization of MergeSort, one might ask whether there is corresponding generalization of other sorting algorithm for computing
LowerTangent\( (H_A, H_B) \):

1. Let \( a \) be the rightmost point of \( H_A \).
2. Let \( b \) be the leftmost point of \( H_B \).
3. While \( (ab) \) is not a lower tangent to \( H_A \) and \( H_B \) do
   a. While \( (ab) \) is not a lower tangent to \( H_A \) do \( a \) ← \( a \).pred (move \( a \) clockwise).
   b. While \( (ab) \) is not a lower tangent to \( H_B \) do \( b \) ← \( b \).succ (move \( b \) counterclockwise).
4. Return \( ab \).

convex hulls. In particular, the next algorithm that we will consider can be thought of as a generalization of the QuickSort sorting procedure. The resulting algorithm is called QuickHull.

Like QuickSort, this algorithm runs in \( O(n \log n) \) time for favorable inputs but can take as long as \( O(n^2) \) time for unfavorable inputs. However, unlike QuickSort, there is no obvious way to convert it into a randomized algorithm with \( O(n \log n) \) expected running time. Nonetheless, QuickHull tends to perform very well in practice.

The intuition is that in many applications most of the points lie in the interior of the hull. For example, if the points are uniformly distributed in a unit square, then it can be shown that the expected number of points on the convex hull is \( O(\log n) \).

The idea behind QuickHull is to discard points that are not on the hull as quickly as possible. QuickHull begins by computing the points with the maximum and minimum, \( x \)- and \( y \)-coordinates. Clearly these points must be on the hull. Horizontal and vertical lines passing through these points are support lines for the hull, and so define a bounding rectangle, within which the hull is contained. Furthermore, the convex quadrilateral defined by these four points lies within the convex hull, so the points lying within this quadrilateral can be eliminated from further consideration. All of this can be done in \( O(n) \) time.

![Figure 3.5: QuickHull’s initial quadrilateral.](image)

To continue the algorithm, we classify the remaining points into the four corner triangles that remain. In general, as this algorithm executes, we will have an inner convex polygon, and associated with each edge we have a set of points that lie “outside” of that edge. (More formally, these points are witnesses to the fact that this edge is not on the convex hull, because they lie outside the half-plane defined by this edge.) When this set of points is empty, the edge is a final edge of the hull. Consider some edge \( ab \). Assume that the points that lie “outside” of this hull edge have been placed in a bucket that is associated with \( ab \). Our job is to find a point \( c \) among these points that lies on the hull, discard the points in the triangle \( abc \), and split the remaining points into two subsets, those that lie outside \( ac \) and those than lie outside of \( cb \). We can classify each point by making two orientation tests.
How should \( c \) be selected? There are a number of possible selection criteria that one might think of. The method that is most often proposed is to let \( c \) be the point that maximizes the perpendicular distance from the line \( ab \). (For example, another possible choice might be the point that maximizes the angle \( cba \) or \( cab \). It turns out that these can be are very poor choices because they tend to produce imbalanced partitions of the remaining points.) We replace the edge \( ab \) with the two edges \( ac \) and \( cb \), and classify the points as lying in one of three groups: those that lie in the triangle \( abc \), which are discarded, those that lie outside of \( ac \), and those that lie outside of \( cb \). We put these points in buckets for these edges, and recurse. (We claim that it is not hard to classify each point \( p \), by computing the orientations of the triples \( acp \) and \( cbp \).)

The running time of QuickHull, as with QuickSort, depends on how evenly the points are split at each stage. Let \( T(n) \) denote the running time on the algorithm assuming that \( n \) points remain outside of some edge. In \( O(n) \) time we can select a candidate splitting point \( c \) and classify the points in the bucket in \( O(n) \) time. Let \( n_1 \) and \( n_2 \) denote the number of remaining points, where \( n_1 + n_2 \leq n \). Then the running time is given by the recurrence:

\[
T(n) = \begin{cases} 
1 & \text{if } n = 1 \\
T(n_1) + T(n_2) & \text{where } n_1 + n_2 \leq n.
\end{cases}
\]

In order to solve this recurrence, it would be necessary to determine the “reasonable” values for \( n_1 \) and \( n_2 \). If we assume that the points are “evenly” distributed, in the sense that \( \max(n_1, n_2) \leq \alpha n \) for some constant \( \alpha < 1 \), then by applying the same analysis as that used in QuickSort (see any standard algorithms text) the running time will solve to \( O(n \log n) \), where the constant factor depends on \( \alpha \). On the other hand, if the splits are not balanced, then the running time can easily increase to \( O(n^2) \).

Does QuickHull outperform Graham’s scan? This depends to a great extent on the distribution of the point set. There are variations of QuickHull that are designed for specific point distributions (e.g. points uniformly distributed in a square) and their authors claim that they manage to eliminate almost all of the points in a matter of only a few iterations.

**Gift-Wrapping and Jarvis’s March:** The next algorithm that we will consider is a variant on an \( O(n^2) \) sorting algorithm called SelectionSort. For sorting, this algorithm repeatedly finds the next element to add to the sorted order from the remaining items. The corresponding convex hull algorithm is called *Jarvis’s march*, which builds the hull in \( O(nh) \) time by a process called “gift-wrapping”. The algorithm operates by considering any one point that is on the hull, say, the lowest point. We then find the “next” edge on the hull in counterclockwise order. Assuming that \( p_k \) and \( p_{k-1} \) were the last two points added to the hull, compute the point \( q \) that maximizes the angle \( \angle p_{k-1}p_kq \). Thus, we can find the point \( q \) in \( O(n) \) time. After repeating this \( h \) times, we will return back to the starting point and we are done. Thus, the overall running time is \( O(nh) \). Note that if \( h \) is \( o(\log n) \) (asymptotically smaller than \( \log n \)) then this is a better method than Graham’s algorithm.
One technical detail is that when we to find an edge from which to start. One easy way to do this is to let \( p_1 \) be the point with the lowest \( y \)-coordinate, and let \( p_0 \) be the point \((−\infty, 0)\), which is infinitely far to the right. The point \( p_0 \) is only used for computing the initial angles, after which it is discarded.

![Diagram](image.png)

Figure 3.7: Jarvis’s march.
4 Line Segment Intersection

Reading: Chapter 2 in the 4M’s.

Geometric intersections: One of the most basic problems in computational geometry is that of computing intersections. Intersection computation in 2- and 3-space is basic to many different application areas.

- In solid modeling people often build up complex shapes by applying various boolean operations (intersection, union, and difference) to simple primitive shapes. The process in called constructive solid geometry (CSG). In order to perform these operations, the most basic step is determining the points where the boundaries of the two objects intersect.
- In robotics and motion planning it is important to know when two objects intersect for collision detection and collision avoidance.
- In geographic information systems it is often useful to overlay two subdivisions (e.g. a road network and county boundaries to determine where road maintenance responsibilities lie). Since these networks are formed from collections of line segments, this generates a problem of determining intersections of line segments.
- In computer graphics, ray shooting is an important method for rendering scenes. The computationally most intensive part of ray shooting is determining the intersection of the ray with other objects.

Most complex intersection problems are broken down to successively simpler and simpler intersection problems. Today, we will discuss the most basic algorithm, upon which most complex algorithms are based.

Line segment intersection: The problem that we will consider is, given \( n \) line segments in the plane, report all points where a pair of line segments intersect. We assume that each line segment is represented by giving the coordinates of its two endpoints.

Observe that \( n \) line segments can intersect in as few as 0 and as many as \( \binom{n}{2} = O(n^2) \) different intersection points. We could settle for an \( O(n^2) \) algorithm, claiming that it is worst-case asymptotically optimal, but it would not be very useful in practice, since in many instances of intersection problems intersections may be rare. Therefore it seems reasonable to look for an output sensitive algorithm, that is, one whose running time should be efficient both with respect to input and output size.

Let \( I \) denote the number of intersections. We will assume that the line segments are in general position, and so we will not be concerned with the issue of whether three or more lines intersect in a single point. However, generalizing the algorithm to handle such degeneracies efficiently is an interesting exercise. See our book for more discussion of this.

Complexity: We claim that best worst-case running time that one might hope for is \( O(n \log n + I) \) time algorithm. Clearly we need \( O(I) \) time to output the intersection points. What is not so obvious is that \( O(n \log n) \) time is needed. This results from the fact that the following problem is known to require \( \Omega(n \log n) \) time in the algebraic decision tree model of computation.

Element uniqueness: Given a list of \( n \) real numbers, are all of these numbers distinct? (That is, are there no duplicates.)
We will reduce the element uniqueness problem to intersection reporting in $O(n)$ time. This means that if we could solve intersection reporting in $o(n \log n)$ time then we could solve element uniqueness in $o(n \log n)$ time, which would contradict the known $\Omega(n \log n)$ lower bound.

Here is the reduction. Given a list of $n$ numbers, $(x_1, x_2, \ldots, x_n)$, in $O(n)$ time we can construct a set of $n$ horizontal line segments, all having the same $y$-coordinates. (E.g., map $x_i$ to the horizontal segment $p_iq_i$ where $p_i = (0, x_i)$ and $q_i = (1, x_i)$). Observe that if the numbers are distinct, then there are no intersections between these segments. On the other hand, if two or more numbers are equal, then the corresponding segments will be equal and hence there is at least one intersection. (It might bother you that we have created a degenerate configuration of segments. However, by applying a small perturbation to each segment, say by rotating about its midpoint by an infinitesimally small amount, this can be remedied.)

Note that this lower-bound result assumes the *algebraic decision tree model* of computation, in which all decisions are made by comparisons made based on exact algebraic operations ($+, -, \cdot, /$) applied to numeric inputs. Although this encompasses most of what one would consider to be “normal” geometric computations, there are alternative models of computation. For example, by taking mods, floors, or ceilings, you would be able to implement hashing. With hashing it is possible to solve the element uniqueness in expected $O(n)$ time. Unfortunately, even in this more powerful computational model, no one know how to determine whether any two line segments intersect in faster than $O(n \log n + I)$ time in the worst case.

We will present a (not quite optimal) $O(n \log n + I \log n)$ time algorithm for the line segment intersection problem. A natural question is whether this is optimal. Later in the semester we will discuss an optimal randomized $O(n \log n + I)$ time algorithm for this problem.

**Line segment intersection:** Like many geometric primitives, computing the point at which two line segments intersect can be reduced to solving a linear system of equations. Let $\overline{ab}$ and $\overline{cd}$ be two line segments, given by their endpoints. First observe that it is possible to determine whether these line segments intersect, simply by applying an appropriate combination of orientation tests. (We will leave this as an exercise.)

One way to determine the point at which the segments intersect is to use an old trick from computer graphics. We represent the segments using a *parametric representation*. Recall that any point on the line segment $\overline{ab}$ can be written as a convex combination involving a real parameter $s$:

$$ p(s) = (1-s)a + sb \quad \text{for} \quad 0 \leq s \leq 1. $$

Similarly for $\overline{cd}$ we may introduce a parameter $t$:

$$ q(t) = (1-t)c + td \quad \text{for} \quad 0 \leq t \leq 1. $$

An intersection occurs if and only if we can find $s$ and $t$ in the desired ranges such that $p(s) = q(t)$. Thus we obtain the two equations:

$$ (1-s)a_x + sb_x = (1-t)c_x + td_x, $$

$$ (1-s)a_y + sb_y = (1-t)c_y + td_y. $$

The coordinates of the points are all known, so it is just a simple exercise in linear algebra to solve for $s$ and $t$. The computation of $s$ and $t$ will involve a division. If the divisor is 0, this means that the

4-2
line segments are either parallel or collinear. These special cases should be dealt with carefully. If the divisor is nonzero, then we get values for \(s\) and \(t\) as rational numbers (the ratio of two integers). We can approximate them as floating point numbers, or if we want to perform exact computations it is possible to simulate rational number algebra exactly using high-precision integers (and multiplying through by least common multiples). Once the values of \(s\) and \(t\) have been computed all that is needed is to check that both are in the interval \([0, 1]\).

**Plane Sweep Algorithm:** Let \(S = \{s_1, s_2, \ldots, s_n\}\) denote the line segments whose intersections we wish to compute. The method is called *plane sweep*. Here are the main elements of any plane sweep algorithm, and how we will apply them to this problem:

**Sweep line:** We will simulate the sweeping of a vertical line \(\ell\), called the *sweep line* from left to right. (Our text uses a horizontal line, but there is obviously no significant difference.) We will maintain the line segments that intersect the sweep line in sorted order (say from top to bottom). Note that we cannot efficiently store the actual coordinates at which the segments intersect the sweep, since this would require too many updates. Instead we just store a pointer which provides us with access to the line segment’s equation. Given the current \(x\)-coordinate of the sweep line, we can compute the \(y\)-coordinate of the intersection point as needed.

**Events:** Although we might think of the sweep line as moving continuously, we only need to update data structures at points of some significant change in the sweep-line contents, called *event points*.

Different applications of plane sweep will have different notions of what event points are. For this application, event points will correspond to the following:

- **Endpoint events:** where the sweep line encounters an endpoint of a line segment, and
- **Intersection events:** where the sweep line encounters an intersection point of two line segments.

Note that endpoint events can be presorted before the sweep runs. In contrast, intersection events will be discovered as the sweep executes. For example, in the figure below, some of the intersection points lying to the right of the sweep line have not yet been “discovered” as events. However, we will show that every intersection point will be discovered as an event before the sweep line arrives here.

**Event updates:** When an event is encountered, we must update the data structures associated with the event. It is a good idea to be careful in specifying exactly what invariants you intend to maintain. For example, when we encounter an intersection point, we must interchange the order of the intersecting line segments along the sweep line.

There are a great number of nasty special cases that complicate the algorithm and obscure the main points. We will make a number of simplifying assumptions. They can be overcome through a more careful handling of these cases.

1. No line segment is vertical.
2. If two segments intersect, then they intersect in a single point (that is, they are not collinear).
3. No three line segments intersect in a common point.
Detecting intersections: We mentioned that endpoint events are all known in advance. But how do we detect intersection events. It is important that each event be detected before the actual event occurs. Our strategy will be as follows. Whenever two line segments become adjacent along the sweep line, we will check whether they have an intersection occurring to the right of the sweep line. If so, we will add this new event (assuming that it has not already been added).

A natural question is whether this is sufficient. In particular, if two line segments do intersect, is there necessarily some prior placement of the sweep line such that they are adjacent. Happily, this is the case, but it requires a proof.

Lemma: Given two segments $s_i$ and $s_j$, which intersect in a single point $p$ (and assuming no other line segment passes through this point) there is a placement of the sweep line prior to this event, such that $s_i$ and $s_j$ are adjacent along the sweep line (and hence will be tested for intersection).

Proof: From our general position assumption it follows that no three lines intersect in a common point. Therefore if we consider a placement of the sweep line that is infinitesimally to the left of the intersection point, lines $s_i$ and $s_j$ will be adjacent along this sweepline. Consider the event point $q$ with the largest $x$-coordinate that is strictly less than $p_x$. The order of lines along the sweep-line after processing $q$ will be identical the order of the lines along the sweep line just prior $p$, and hence $s_i$ and $s_j$ will be adjacent at this point.

Data structures: In order to perform the sweep we will need two data structures.

Event queue: This holds the set of future events, sorted according to increasing $x$-coordinate. Each event contains the auxiliary information of what type of event this is (segment endpoint or intersection) and which segment(s) are involved. The operations that this data structure should support are inserting an event (if it is not already present in the queue) and extracting the minimum event.
It seems like a heap data structure would be ideal for this, since it supports insertions and extract-min in $O(\log M)$ time, where $M$ is the number of entries in the queue. (See any standard algorithms text for details). However, a heap cannot support the operation of checking for duplicate events.

There are two ways to handle this. One is to use a more sophisticated data structure, such as an ordered dictionary (for example, a balanced binary tree or skip-list). This adds a small constant factor, but can check that there are no duplicates easily. The second is use the heap and accept the fact that there may be duplicate events present. Some mechanism will need to be implemented to detect and ignore these duplicate events. (Our book recommends the prior solution.)

If events have the same $x$-coordinate, then we can handle this by sorting points lexicographically by $(x, y)$. (This results in events be processed from bottom to top along the sweep line, and has the same geometric effect as imagining that the sweep line is rotated infinitesimally counterclockwise.)

**Sweep line status:** To store the sweep line status, we maintain an ordered dictionary (e.g., a balanced binary tree or skip-list) whose entries are pointers to the line segments, stored in increasing order of $y$-coordinate along the current sweep line.

Normally when storing items in a tree, the key values are constants. Since the sweep line varies, we need “variable” keys. To do this, let us assume that each line segment computes a line equation $y = mx + b$ as part of its representation. The “key” value in each node of the tree is a pointer to a line segment. To compute the $y$-coordinate of some segment at the location of the current sweep line, we simply take the current $x$-coordinate of the sweep line, plug it into the line equation for this line, and use the resulting $y$-coordinate in our comparisons.

The operations that we need to support are those of deleting a line segment, inserting a line segment, swapping the position of two line segments, and determining the immediate predecessor and successor of any item. Assuming any balanced binary tree or a skiplist, these operations can be performed in $O(\log n)$ time each.

![Figure 4.3: Plane-sweep algorithm events.](image)

The complete plane-sweep algorithm is presented below. Our algorithm considers the creation of new events, but ignores the issue of duplicate events. See our text for a more careful implementation. The various cases are illustrated in the following figure.

**Analysis:** The work done by the algorithm is dominated by the time spent updating the various data structures (since otherwise we spend only constant time per sweep event). We need to count two things:
Plane-Sweep Algorithm for Line Segment Intersection

(1) Insert all of the endpoints of the line segments of $S$ into the event queue. The initial sweep status is empty.

(2) While the event queue is nonempty, extract the next event in the queue. There are three cases, depending on the type of event:

- **Segment left endpoint**: Insert this line segment into the sweep line status, based on the $y$-coordinate of this endpoint and the $y$-coordinates of the other segments currently along the sweep line. Test for intersections with the segment immediately above and below.

- **Segment right endpoint**: Delete this line segment from the sweep line status. For the entries immediately preceding and succeeding this entry, test them for intersections.

- **Intersection point**: Swap the two line segments in order along the sweep line. For the new upper segment, test it against its predecessor for an intersection. For the new lower segment, test it against its successor for an intersection.

the number of operations applied to each data structure and the amount of time needed to process each operation.

For the sweep line status, there are at most $n$ elements intersecting the sweep line at any time, and therefore the time needed to perform any single operation is $O(\log n)$, from standard results on balanced binary trees.

Since we do not allow duplicate events to exist in the event queue, the total number of elements in the queue at any time is at most $2n + I$. Since we use a balanced binary tree to store the event queue, each operation takes time at most logarithmic in the size of the queue, which is $O(\log(2n + I))$. Since $I \leq n^2$, this is at most $O(\log n^2) = O(2 \log n) = O(\log n)$ time.

Each event involves a constant number of accesses or operations to the sweep status or the event queue, and since each such operation takes $O(\log n)$ time from the previous paragraph, it follows that the total time spent processing all the events from the sweep line is

$$O((2n + I) \log n) = O((n + I) \log n) = O(n \log n + I \log n).$$

Thus, this is the total running time of the plane sweep algorithm.
5 Planar Graphs, Polygons and Art Galleries

Reading: Chapter 3 in the 4M’s.

Topological Information: In many applications of segment intersection problems, we are not interested in just a listing of the segment intersections, but want to know how the segments are connected together. Typically, the plane has been subdivided into regions, and we want to store these regions in a way that allows us to reason about their properties efficiently.

This leads to the concept of a planar straight line graph (PSLG) or planar subdivision (or what might be called a cell complex in topology). A PSLG is a graph embedded in the plane with straight-line edges so that no two edges intersect, except possibly at their endpoints. (The condition that the edges be straight line segments may be relaxed to allow curved segments, but we will assume line segments here.) Such a graph naturally subdivides the plane into regions. The 0-dimensional vertices, 1-dimensional edges, and 2-dimensional faces. We consider these three types of objects to be disjoint, implying each edge is topologically open (it does not include it endpoints) and that each face is open (it does not include its boundary). There is always one unbounded face, that stretches to infinity. Note that the underlying planar graph need not be a connected graph. In particular, faces may contain holes (and these holes may contain other holes. A subdivision is called a convex subdivision if all the faces are convex.

Planar subdivisions form the basic objects of many different structures that we will discuss later this semester (triangulations and Voronoi diagrams in particular) so this is a good time to consider them in greater detail. The first question is how should we represent such structures so that they are easy to manipulate and reason about. For example, at a minimum we would like to be able to list the edges that bound each face of the subdivision in cyclic order, and we would like to be able to list the edges that surround each vertex.

Planar graphs: There are a number of important facts about planar graphs that we should discuss. Generally speaking, an (undirected) graph is just a finite set of vertices, and collection of unordered pairs of distinct vertices called edges. A graph is planar if it can be drawn in the plane (the edges need not be straight lines) so that no two distinct edges cross each other. An embedding of a planar graph is any such drawing. In fact, in specifying an embedding it is sufficient just to specify the counterclockwise cyclic list of the edges that are incident to each vertex. Since we are interested in geometric graphs, our embeddings will contain complete geometric information (coordinates of vertices in particular).

There is an important relationship between the number of vertices, edges, and faces in a planar graph (or more generally an embedding of any graph on a topological 2-manifold, but we will stick to the plane). Let \( V \) denote the number of vertices, \( E \) the number of edges, \( F \) the number of faces in a
connected planar graph. Euler’s formula states that
\[ V - E + F = 2. \]

The quantity \( V - E + F \) is called the *Euler characteristic*, and is an invariant of the plane. In general, given a orientable topological 2-manifold with \( g \) handles (called the *genus*) we have
\[ V - E + F = 2 - 2g. \]

Returning to planar graphs, if we allow the graph to be disconnected, and let \( C \) denote the number of connected components, then we have the somewhat more general formula
\[ V - E + F - C = 1. \]

In our example above we have \( V = 13, E = 12, F = 4 \) and \( C = 4 \), which clearly satisfies this formula. An important fact about planar graphs follows from this.

**Theorem:** A planar graph with \( V \) vertices has at most \( 3(V - 2) \) edges and at most \( 2(V - 2) \) faces.

**Proof:** We assume (as is typical for graphs) that there are no multiple edges between the same pair of vertices and no self-loop edges.

We begin by *triangulating* the graph. For each face that is bounded by more than three edges (or whose boundary is not connected) we repeatedly insert new edges until every face in the graph is bounded by exactly three edges. (Note that this is not a “straight line” planar graph, but it is a planar graph, nonetheless.) An example is shown in the figure below in which the original graph edges are shown as solid lines.

![Figure 5.2: Triangulating a planar graph.](image)

Let \( E' \geq E \) and \( F' \geq F \) denote the number edges and faces in the modified graph. The resulting graph has the property that it has one connected component, every face is bounded by exactly three edges, and each edge has a different face on either side of it. (The last claim may involve a little thought.)

If we count the number of faces and multiply by 3, then every edge will be counted exactly twice, once by the face on either side of the edge. Thus, \( 3F' = 2E' \), that is \( E' = 3F'/2 \). Euler’s formula states that \( V + E' - F' = 2 \), and hence
\[ V - \frac{3F'}{2} + F' = 2 \quad \Rightarrow \quad F \leq F' = 2(V - 2), \]

and using the face that \( F' = 2E'/3 \) we have
\[ V - E' + \frac{2E'}{3} = 2 \quad \Rightarrow \quad E \leq E' = 3(V - 2). \]

This completes the proof.
The fact that the numbers of vertices, edges, and faces are related by constant factors seems to hold only in 2-dimensional space. For example, a polyhedral subdivision of 3-dimensional space that has \( n \) vertices can have as many as \( \Theta(n^2) \) edges. (As a challenging exercise, you might try to create one.) In general, there are formulas, called the Dehn-Sommerville equations that relate the maximum numbers of vertices, edges, and faces of various dimensions.

There are a number of reasonable representations that for storing PSLG's. The most widely used on is the winged-edge data structure. Unfortunately, it is probably also the messiest. There is another called the quad-edge data structure which is quite elegant, and has the nice property of being self-dual. (We will discuss duality later in the semester.) We will not discuss any of these, but see our text for a presentation of the doubly-connected edge list (or DCEL) structure.

**Simple Polygons:** Now, let us change directions, and consider some interesting problems involving polygons in the plane. We begin study of the problem of triangulating polygons. We introduce this problem by way of a cute example in the field of combinatorial geometry.

We begin with some definitions. A **polygonal curve** is a finite sequence of line segments, called **edges** joined end-to-end. The endpoints of the edges are **vertices**. For example, let \( v_0, v_2, \ldots, v_n \) denote the set of \( n + 1 \) vertices, and let \( e_1, e_2, \ldots, e_n \) denote a sequence of \( n \) edges, where \( e_i = v_{i-1}v_i \). A polygonal curve is **closed** if the last endpoint equals the first \( v_0 = v_n \). A polygonal curve is **simple** if it is not self-intersecting. More precisely this means that each edge \( e_i \) does not intersect any other edge, except for the endpoints it shares with its adjacent edges.

![Figure 5.3: Polygonal curves](image)

The famous **Jordan curve theorem** states that every simple closed plane curve divides the plane into two regions (the **interior** and the **exterior**). (Although the theorem seems intuitively obvious, it is quite difficult to prove.) We define a **polygon** to be the region of the plane bounded by a simple, closed polygonal curve. The term **simple polygon** is also often used to emphasize the simplicity of the polygonal curve. We will assume that the vertices are listed in counterclockwise order around the boundary of the polygon.

**Art Gallery Problem:** We say that two points \( x \) and \( y \) in a simple polygon can see each other (or \( x \) and \( y \) are visible) if the open line segment \( xy \) lies entirely within the interior of \( P \). (Note that such a line segment can start and end on the boundary of the polygon, but it cannot pass through any vertices or edges.)

If we think of a polygon as the floor plan of an art gallery, consider the problem of where to place “guards”, and how many guards to place, so that every point of the gallery can be seen by some guard. Victor Klee posed the following question: Suppose we have an art gallery whose floor plan can be modeled as a polygon with \( n \) vertices. As a function of \( n \), what is the minimum number of guards that suffice to guard such a gallery? Observe that are you are told about the polygon is the number of sides, not its actual structure. We want to know the fewest number of guards that suffice to guard all polygons with \( n \) sides.
Before getting into a solution, let’s consider some basic facts. Could there be polygons for which no finite number of guards suffice? It turns out that the answer is no, but the proof is not immediately obvious. You might consider placing a guard at each of the vertices. Such a set of guards will suffice in the plane. But to show how counterintuitive geometry can be, it is interesting to note that there are simple nonconvex polyhedra in 3-space, such that even if you place a guard at every vertex there would still be points in the polygon that are not visible to any guard. (As a challenge, try to come up with one with the fewest number of vertices.)

An interesting question in combinatorial geometry is how does the number of guards needed to guard any simple polygon with \(n\) sides grow as a function of \(n\)? If you play around with the problem for a while (trying polygons with \(n = 3, 4, 5, 6 \ldots\) sides, for example) you will eventually come to the conclusion that \(\lfloor n/3 \rfloor\) is the right value. The figure above shows a worst-case example, where \(\lfloor n/3 \rfloor\) guards are required. A cute result from combinatorial geometry is that this number always suffices. The proof is based on three concepts: polygon triangulation, dual graphs, and graph coloring. The remarkably clever and simple proof was discovered by Fisk.

**Theorem:** (The Art-Gallery Theorem) Given a simple polygon with \(n\) vertices, there exists a guarding set with at most \(\lfloor n/3 \rfloor\) guards.

Before giving the proof, we explore some aspects of polygon triangulations. We begin by introducing a triangulation of \(P\). A triangulation of a simple polygon is a planar subdivision of (the interior of) \(P\) whose vertices are the vertices of \(P\) and whose faces are all triangles. An important concept in polygon triangulation is the notion of a diagonal, that is, a line segment between two vertices of \(P\) that are visible to one another. A triangulation can be viewed as the union of the edges of \(P\) and a maximal set of noncrossing diagonals.

**Lemma:** Every simple polygon with \(n\) vertices has a triangulation consisting of \(n - 3\) diagonals and \(n - 2\) triangles.

(We leave the proof as an exercise.) The proof is based on the fact that given any \(n\)-vertex polygon, with \(n \geq 4\) it has a diagonal. (This may seem utterly trivial, but actually takes a little bit of work to prove. In fact it fails to hold for polyhedra in 3-space.) The addition of the diagonal breaks the polygon into two polygons, of say \(m_1\) and \(m_2\) vertices, such that \(m_1 + m_2 = n + 2\) (since both share the vertices of the diagonal). Thus by induction, there are \((m_1 - 2) + (m_2 - 2) = n + 2 - 4 = n - 2\) triangles total. A similar argument holds for the case of diagonals.

It is a well known fact from graph theory that any planar graph can be colored with 4 colors. (The famous 4-color theorem.) This means that we can assign a color to each of the vertices of the graph,
from a collection of 4 different colors, so that no two adjacent vertices have the same color. However we can do even better for the graph we have just described.

Figure 5.5: Polygon triangulation and a 3-coloring.

**Lemma:** Let $T$ be the triangulation graph of a triangulation of a simple polygon. Then $T$ is 3-colorable.

**Proof:** For every planar graph $G$ there is another planar graph $G^*$ called its dual. The dual $G^*$ is the graph whose vertices are the faces of $G$, and two vertices of $G^*$ are connected by an edge if the two corresponding faces of $G$ share a common edge.

Since a triangulation is a planar graph, it has a dual, shown in the figure below. (We do not include the external face in the dual.) Because each diagonal of the triangulation splits the polygon into two, it follows that each edge of the dual graph is a cut edge, meaning that its deletion would disconnect the graph. As a result it is easy to see that the dual graph is a free tree (that is, a connected, acyclic graph), and its maximum degree is 3. (This would not be true if the polygon had holes.)

Figure 5.6: Dual graph of triangulation.

The coloring will be performed inductively. If the polygon consists of a single triangle, then just assign any 3 colors to its vertices. An important fact about any free tree is that it has at least one leaf (in fact it has at least two). Remove this leaf from the tree. This corresponds to removing a triangle that is connected to the rest triangulation by a single edge. (Such a triangle is called an ear.) By induction 3-color the remaining triangulation. When you add back the deleted triangle, two of its vertices have already been colored, and the remaining vertex is adjacent to only these two vertices. Give it the remaining color. In this way the entire triangulation will be 3-colored.

We can now give the simple proof of the guarding theorem.

**Proof:** (of the Art-Gallery Theorem:) Consider any 3-coloring of the vertices of the polygon. At least one color occurs at most $\lfloor n/3 \rfloor$ time. (Otherwise we immediately get there are more than $n$ vertices, a contradiction.) Place a guard at each vertex with this color. We use at most $\lfloor n/3 \rfloor$ guards. Observe that every triangle has at least one vertex of each of the tree colors (since you
cannot use the same color twice on a triangle). Thus, every point in the interior of this triangle is guarded, implying that the interior of $P$ is guarded. A somewhat messy detail is whether you allow guards placed at a vertex to see along the wall. However, it is not a difficult matter to push each guard infinitesimally out from his vertex, and so guard the entire polygon.
6 Polygon Triangulation

Reading: Chapter 3 in the 4M’s.

The Polygon Triangulation Problem: Triangulation is the general problem of subdividing a spatial domain into simplices, which in the plane means triangles. In its simplest form, a simple polygon is given (that is, a planar region that is defined by a closed, simple polygonal curve), and the objective is to subdivide the polygon into triangles. Triangulating simple polygons is important for many reasons. This operation useful, for example, whenever it is needed to decompose a complex shape into a set of disjoint simpler shapes. Note that in some applications it is desirable to produce “fat” (nearly equilateral) triangles, but we will not worry about this issue.

This simple problem has been the focus of a remarkably large number of papers in computational geometry spanning a number of years. There is a simple naive polynomial-time algorithm for the planar case (as opposed to possibly nonconvex polyhedra in higher dimensions). We say that two points on the boundary of the polygon are visible if the interior of the line segment lies entirely within the interior of the polygon. Define a diagonal of the polygon to be the line segment joining any pair of visible vertices. Observe that the addition of a diagonal splits the polygon into two polygons of smaller size. This naive algorithm operates by repeatedly adding diagonals. (Its correctness is based on the observation that every non-triangular planar polygon has at least one diagonal.) Unfortunately, this algorithm is not very efficient (unless the polygon has special properties, for example, convexity) because of the complexity of the visibility test.

There are very simple $O(n \log n)$ algorithms for this problem that have been known for many years. A longstanding open problem was whether there exists an $O(n)$ time algorithm. (Observe that the input polygon is presented as a cyclic list of vertices, and hence the data is in some sense “pre-sorted”, which precludes an $\Omega(n \log n)$ lower bound.) The problem of a linear time polygon triangulation was solved by Bernard Chazelle in 1991, but the algorithm is so amazingly intricate, it could never compete with the practical but asymptotically slower $O(n \log n)$ algorithms. In fact, there is no known algorithm that runs in less than $O(n \log n)$ time, that is really practical enough to replace the standard $O(n \log n)$ algorithms.

We will present one of many known $O(n \log n)$ algorithms. The approach we present today is a two-step process (although with a little cleverness, both steps can be combined into one algorithm). The first is to consider the special case of triangulating a monotone polygon. After this we consider how to convert an arbitrary polygon into a collection of disjoint monotone polygons. Then we will apply the first algorithm to each of the monotone pieces. The former algorithm runs in $O(n)$ time. The latter algorithm runs in $O(n \log n)$ time, for a total of $O(n \log n)$ time.
Monotone Polygons: A polygonal chain $C$ is said to be strictly monotone with respect to a given line $L$, if any line that is orthogonal to $L$ intersects $C$ in at most one point. A chain $C$ is monotone with respect to $L$ if each line that is orthogonal to $L$ intersects $C$ in a single connected component. Thus it may intersect, not at all, at a single point, or along a single line segment. A polygon $P$ is said to be monotone with respect to a line $L$ if its boundary, (sometimes denoted $\text{bnd}(P)$ or $\partial P$), can be split into two chains, each of which is monotone with respect to $L$.

Henceforth, let us consider monotonicity with respect to the $x$-axis. We will call these polygons horizontally monotone. It is easy to test whether a polygon is horizontally monotone. How?

(a) Find the leftmost and rightmost vertices (min and max $x$-coordinate) in $O(n)$ time.
(b) These vertices split the polygon’s boundary into two chains, an upper chain and a lower chain. Walk from left to right along each chain, verifying that the $x$-coordinates are nondecreasing. This takes $O(n)$ time.

As a challenge, consider the problem of determining whether a polygon is monotone in any (unspecified) direction. This can be done in $O(n)$ time, but is quite a bit harder.

Triangulation of Monotone Polygons: We can triangulate a monotone polygon by a simple variation of the plane-sweep method. We begin with the assumption that the vertices of the polygon have been sorted in increasing order of their $x$-coordinates. (For simplicity we assume no duplicate $x$-coordinates. Otherwise, break ties between the upper and lower chains arbitrarily, and within a chain break ties so that the chain order is preserved.) Observe that this does not require sorting. We can simply extract the upper and lower chain, and merge them (as done in MergeSort) in $O(n)$ time.

The idea behind the triangulation algorithm is quite simple: Try to triangulate everything you can to the left of the current vertex by adding diagonals, and then remove the triangulated region from further consideration.

In the example, there is obviously nothing to do until we have at least 3 vertices. With vertex 3, it is possible to add the diagonal to vertex 2, and so we do this. In adding vertex 4, we can add the diagonal to vertex 2. However, vertices 5 and 6 are not visible to any other nonadjacent vertices so no new diagonals can be added. When we get to vertex 7, it can be connected to 4, 5, and 6. The process continues until reaching the final vertex.

The important thing that makes the algorithm efficient is the fact that when we arrive at a vertex the untriangulated region that lies to the left of this vertex always has a very simple structure. This structure allows us to determine in constant time whether it is possible to add another diagonal. And in general we can add each additional diagonal in constant time. Since any triangulation consists of $n - 3$ diagonals, the process runs in $O(n)$ total time. This structure is described in the lemma below.
Lemma: (Main Invariant) For \( i \geq 2 \), let \( v_i \) be the vertex just processed by the triangulation algorithm. The untriangulated region lying to the left of \( v_i \) consists of two \( x \)-monotone chains, a lower chain and an upper chain each containing at least one edge. If the chain from \( v_i \) to \( u \) has two or more edges, then these edges form a reflex chain (that is, a sequence of vertices with interior angles all at least 180 degrees). The other chain consists of a single edge whose left endpoint is \( u \) and whose right endpoint lies to the right of \( v_i \).

We will prove the invariant by induction. As the basis case, consider the case of \( v_2 \). Here \( u = v_1 \), and one chain consists of the single edge \( v_2v_1 \) and the other chain consists of the other edge adjacent to \( v_1 \).

To prove the main invariant, we will give a case analysis of how to handle the next event, involving \( v_i \), assuming that the invariant holds at \( v_{i-1} \) and see that the invariant is satisfied after each event has been processed. There are the following cases that the algorithm needs to deal with.

Case 1: \( v_i \) lies on the opposite chain from \( v_{i-1} \):
In this case we add diagonals joining \( v_i \) to all the vertices on the reflex chain, from \( v_{i-1} \) back to (but not including) \( u \). Note that all of these vertices are visible from \( v_i \). Certainly \( u \) is visible to \( v_i \). Because the chain is reflex, \( x \)-monotone, and lies to the left of \( v_i \) it follows that the chain itself cannot block the visibility from \( v_i \) to some other vertex on the chain. Finally, the fact that the polygon is \( x \)-monotone implies that the unprocessed portion of the polygon (lying to the right of \( v_i \)) cannot “sneak back” and block visibility to the chain.

After doing this, we set \( u = v_{i-1} \). The invariant holds, and the reflex chain is trivial, consisting of the single edge \( v_i v_{i-1} \).

Case 2: \( v \) is on the same chain as \( v_{i-1} \):
We walk back along the reflex chain adding diagonals joining \( v_i \) to prior vertices until we find the first that is not visible to \( v_i \). As can be seen in the figure, this may involve connecting \( v_i \) to one
or more vertices (2a) or it may involve connecting \( v_i \) to no additional vertices (2b), depending on whether the first angle is less or greater than 180 degrees. In either case the vertices that were cut off by diagonals are no longer in the chain, and \( v_i \) becomes the new endpoint to the chain. Again, by \( x \)-monotonicity it follows that the unprocessed portion of the polygon cannot block visibility of \( v_i \) to the chain.

Note that when we are done the remaining chain from \( v_i \) to \( u \) is a reflex chain. (Note the similarity between this step and the main iteration in Graham’s scan.)

How is this implemented? The vertices on the reflex chain can be stored in a stack. We keep a flag indicating whether the stack is on the upper chain or lower chain, and assume that with each new vertex we know which chain of the polygon it is on. Note that decisions about visibility can be based simply on orientation tests involving \( v_i \) and the top two entries on the stack. When we connect \( v_i \) by a diagonal, we just pop the stack.

**Analysis:** We claim that this algorithm runs in \( O(n) \) time. As we mentioned earlier, the sorted list of vertices can be constructed in \( O(n) \) time through merging. The reflex chain is stored on a stack. In \( O(1) \) time per diagonal, we can perform an orientation test to determine whether to add the diagonal and (assuming a DCEL) the diagonal can be added in constant time. Since the number of diagonals is \( n - 3 \), the total time is \( O(n) \).

**Monotone Subdivision:** In order to run the above triangulation algorithm, we first need to subdivide an arbitrary simple polygon \( P \) into monotone polygons. This is also done by a plane-sweep approach. We will add a set of nonintersecting diagonals that partition the polygon into monotone pieces.

Observe that the absence of \( x \)-monotonicity occurs only at vertices in which the interior angle is greater than 180 degrees and both edges lie either to the left of the vertex or both to the right. Following our book’s notation, we call the first type a **merge vertex** (since as the sweep passes over this vertex the edges seem to be merging) and the latter type a **split vertex**.

Let’s discuss the case of a split vertex first (both edges lie to the right of the vertex). When a split vertex \( v \) is encountered in the sweep, there will be an edge \( e_a \) of the polygon lying above and an edge \( e_b \) lying below. We might consider attaching the split vertex to left endpoint of one of these two edges, but it might be that neither endpoint is visible to the split vertex. We need to maintain a vertex that is visible to any split vertex that may arise between \( e_a \) and \( e_b \). To do this, imagine a fluorescent light
shining down from every point on $e_a$ (these are the white vertices in the figure below, left). Note that $e_a$ left endpoint is considered in this set as well. We say that these vertices are vertically visible below $e_a$. Among all the vertices that are illuminated by these vertical light rays, let $u$ be the rightmost. We claim that $u$ is visible to every point along the sweep line between $e_a$ and $e_b$. This suggests the following concept, which is defined for each edge $e_a$ that intersects the sweep line, such that the polygon’s interior lies locally below $e_a$.

$$\text{helper}(e_a) :$$ Let $e_b$ be the edge of the polygon lying just below $e_a$ on the sweep line. The helper is the rightmost vertically visible vertex below $e_a$ on the polygonal chain between $e_a$ and $e_b$.

We join each split vertex to $\text{helper}(e_a)$, where $e_a$ is the edge of $P$ immediately above the split vertex. Note that it is possible that the helper is the left endpoint of $e_a$. Also note that $\text{helper}(e_a)$ is defined with respect to the current location of the sweep line. As the sweep line moves, its value changes. Also, it is only defined for those edges intersected by the sweep line.

**Events:** The endpoints of the edges of the polygon. These are sorted by increasing order of $x$-coordinates. Since no new events are generated, the events may be stored in a simple sorted list (i.e., no priority queue is needed).

**Sweep status:** The sweep line status consists of the list of edges that intersect the sweep line, sorted from top to bottom. Our book notes that we actually only need to store edges such that the polygon lies just below this edge (since these are the only edges that we evaluate $\text{helper}()$ from). These edges are stored in a dictionary (e.g., a balanced binary tree), so that the operations of insert, delete, find, predecessor and successor can be evaluated in $O(\log n)$ time each.

**Event processing:** There are six event types based on a case analysis of the local structure of edges around each vertex. Let $v$ be the current vertex encountered by the sweep.

- **Split vertex:** Search the sweep line status to find the edge $e$ lying immediately above $v$. Add a diagonal connecting $v$ to $\text{helper}(e)$. Add the two edges incident to $v$ in the sweep line status, and make $v$ the helper of the lower of these two edges and make $v$ the new helper of $e$.

- **Merge vertex:** Find the two edges incident to this vertex in the sweep line status (they must be adjacent). Delete them both. Let $e$ be the edge lying immediately above them. Make $v$ the new helper of $e$.

- **Start vertex:** (Both edges lie to the right of $v$, but the interior angle is less than 180 degrees.) Insert this vertex’s edges into the sweep line status. Set the helper of the upper edge to $v$.

Figure 6.5: Split vertices, merge vertices, and helpers.
**End vertex:** (Both edges lie to the left of $v$, but the interior angle is less than 180 degrees.) Delete both edges from the sweep line status.

**Upper-chain vertex:** (One edge is to the left, and one to the right, and the polygon interior is below.) Replace the left edge with the right edge in the sweep line status. Make $v$ the helper of the new edge.

**Lower-chain vertex:** (One edge is to the left, and one to the right, and the polygon interior is above.) Replace the left edge with the right edge in the sweep line status. Let $e$ be the edge lying above here. Make $v$ the helper of $e$.

Finally, in the cases of a End, Upper-chain or Lower-chain vertex, check whether the helper is a Merge vertex. If so, add a diagonal to it.

There are many special cases (what a pain!), but each one is fairly easy to deal with, so the algorithm is quite efficient. As with previous plane sweep algorithms, it is not hard to show that the running time is $O(\log n)$ times the number of events. In this case there is one event per vertex, so the total time is $O(n \log n)$. This gives us an $O(n \log n)$ algorithm for polygon triangulation.
7 Halfplane Intersection

Reading: Chapter 4 in the 4M’s, with some elements from Sections 8.2 and 11.4.

Halfplane Intersection: Today we begin studying another very fundamental topic in geometric computing, and along the way we will show a rather surprising connection between this topic and the topic of convex hulls, which we discussed earlier. Any line in the plane splits the plane into two regions, called halfplane, one lying on either side of the line. We may refer to a halfplane as being either closed or open depending on whether it contains the line itself. For this lecture we will be interested in closed halfplanes.

How do we represent lines and halfplanes? For the cleanest and most general understanding of representing lines, it is good to study projective geometry and homogeneous coordinates. However, for the sake of time, we will skip this. Typically, it will suffice to represent lines in the plane using the following equation:

\[ y = ax - b, \]

where \( a \) denotes the slope and \( b \) denotes the negation of the \( y \)-intercept. (We will see later why this representation is convenient.) Unfortunately, it is not fully general, since it cannot represent vertical lines. A more general line representation will generally involve three parameters, as in:

\[ ax + by = c. \]

In this case, it is easy to see that the line has slope \(-b/a\) and hence is perpendicular to the vector \((a, b)\). The equation is unchanged by an scalar multiplication, and so if \( c \neq 0 \) (the line does not pass through the origin) we could express this more succinctly as \( a'x + b'y = 1 \), where \( a' = a/c \) and \( b' = b/c \).

To represent a closed halfplane, we convert either representation into an inequality:

\[ y \leq ax - b \quad \text{or} \quad ax + by \leq c. \]

In the former case, this represents the halfplane lying below the line. The latter representation is more general, since we can represent halfplanes on either side of the line, simply by multiplying all the coefficients by \(-1\).

Halfplane intersection problem: The halfplane intersection problem is, given a set of \( n \) closed halfplanes, \( H = \{h_1, h_2, \ldots, h_n\} \) compute their intersection. A halfplane (closed or open) is a convex set, and hence the intersection of any number of halfplanes is also a convex set. Unlike the convex hull problem, the intersection of \( n \) halfplanes may generally be empty or even unbounded. A reasonable output representation might be to list the lines bounding the intersection in counterclockwise order, perhaps along with some annotation as to whether the final figure is bounded or unbounded.

How many sides can bound the intersection of \( n \) halfplanes in the worst case? Observe that by convexity, each of the halfplanes can appear only once as a side, and hence the maximum number of sides is \( n \). How fast can we compute the intersection of halfspaces? As with the convex hull problem, it can be shown through a suitable reduction from sorting that the problem has a lower bound of \( \Omega(n \log n) \).
Who cares about this problem? Our book discusses an application in the area of casting. Halfplane intersection and halfspace intersection in higher dimensions are also used in generating convex shape approximations. In computer graphics for example, a bounding box is often used to approximate a complex multi-sided polyhedral shape. Testing predicates (visibility or intersection, for example) may be much easier and faster with the bounding box than with the original object. We can generalize this to produce more accurate bounding objects by considering halfspaces that are not only axis-aligned, but those that are aligned along some fixed number of pre-chosen directions.

Solving the halfspace intersection problem in higher dimensions is quite a bit more challenging than in the plane. For example, the boundary of the intersection of halfspaces in dimension \(d\) is a \((d - 1)\)-dimensional cell complex, and would require an appropriate data structure for storing such objects.

We will discuss two algorithms for the halfplane intersection problem. The first is given in the text, and involves an interesting combination of two techniques we have discussed for geometric problems, geometric divide-and-conquer and plane sweep. For the other, we will consider somewhat simpler problem of computing something called the **lower envelope** of a set of lines, and show that it is closely related to the convex hull problem.

**Divide-and-Conquer Algorithm:** We begin by sketching a divide-and-conquer algorithm for computing the intersection of halfplanes. The basic approach is very simple:

1. If \(n = 1\), then just return this halfplane as the answer.
2. Split the \(n\) halfplanes of \(H\) into subsets \(H_1\) and \(H_2\) of sizes \([n/2]\) and \([n/2]\), respectively.
3. Compute the intersection of \(H_1\) and \(H_2\), each by calling this procedure recursively. Let \(K_1\) and \(K_2\) be the results.
4. Intersect the convex polygons \(K_1\) and \(K_2\) (which might be unbounded) into a single convex polygon \(K\), and return \(K\).

The running time of the resulting algorithm is most easily described using a **recurrence**, that is, a recursively defined equation. If we ignore constant factors, and assume for simplicity that \(n\) is a power of 2, then the running time can be described as:

\[
T(n) = \begin{cases} 
1 & \text{if } n = 1, \\
2T(n/2) + M(n) & \text{if } n > 1,
\end{cases}
\]

where \(M(n)\) is the time required to merge the two results, that is, to compute the intersection of two convex polygons whose total complexity is \(n\). We will show below that \(M(n) = O(n)\), and so it follows by standard results in recurrences that the overall running time \(T(n)\) is \(O(n \log n)\). (See any standard algorithms textbook.)
**Intersecting Two Convex Polygons:** The only nontrivial part of the process is implementing an algorithm that intersects two convex polygons, $K_1$ and $K_2$, into a single convex polygon. Note that these are somewhat special convex polygons because they may be empty or unbounded.

We know that it is possible to compute the intersection of line segments in $O((n + I) \log n)$ time, where $I$ is the number of intersecting pairs. Two convex polygons cannot intersect in more than $I = O(n)$ pairs. (This follows from the observation that each edge of one polygon can intersect at most two edges of the other polygon by convexity.) This would give $O(n \log n)$ algorithm for computing the intersection. This is too slow, however, and would result in an overall time of $O(n \log^2 n)$ for $T(n)$.

There are two common approaches for intersecting convex polygons. Both essentially involve merging the two boundaries. One works by a plane-sweep approach. The other involves a simultaneous counterclockwise sweep around the two boundaries. The latter algorithm is described in O'Rourke's book. We'll discuss the plane-sweep algorithm.

We perform a left-to-right plane sweep to compute the intersection. We begin by breaking the boundaries of the convex polygons into their upper and lower chains. By convexity, the sweep line intersects each convex polygon $K_i$ in at most two points, and hence, there are at most four points in the sweep line status at any time. Thus, we do not need a ordered dictionary for storing the sweep line status—a simple 4-element list suffices. Also, our event queue need only be of constant size. At any point there are at most 8 possible candidates for the next event, namely, the right endpoints of the four edges stabbed by the sweep line and the (up to four) intersection points of these upper and lower edges of $K_1$ with the upper and lower edges of $K_2$. Since there are only a constant number of possible events, and each can be handled in $O(1)$ time, the total running time is $O(n)$.

![Convex polygon intersection.](image)

**Lower Envelopes and Duality:** Next we consider a slight variant of this problem, to demonstrate some connections with convex hulls. These connections are very important to an understanding of computational geometry, and we see more about them in the future. These connections have to do with a concept called point-line duality. In a nutshell there is a remarkable similarity between how points interact with each other an how lines interact with each other. Sometimes it is possible to take a problem involving points and map it to an equivalent problem involving lines, and vice versa. In the process, new insights to the problem may become apparent.

The problem to consider is called the lower envelope problem, and it is a special case of the halfplane intersection problem. We are given a set of $n$ lines $L = \{\ell_1, \ell_2, \ldots, \ell_n\}$ where $\ell_i$ is of the form...
Think of these lines as defining \( n \) halfplanes, \( y \leq a_i x - b_i \), each lying below one of the lines. The lower envelope of \( L \) is the boundary of the intersection of these half planes. (There is also an upper envelope, formed by considering the intersection of the halfplanes lying above the lines.)

The lower envelope problem is a restriction of the halfplane intersection problem, but it an interesting restriction. Notice that any halfplane intersection problem that does not involve any vertical lines can be rephrased as the intersection of two envelopes, a lower envelope defined by the lower halfplanes and an upper envelope defined by the upward halfplanes.

We will see that solving the lower envelope problem is very similar to solving the upper convex hull problem. In fact, they are so similar that exactly the same algorithm will solve both problems, without changing even a single character of code. (!) All that changes is the way in which you view the two inputs and the two outputs.

**Duality:** Earlier this semester we presented the notion of polarity. This transformation maps points to hyperplanes and vice versa, and preserves incidence relationships. There are a number of different ways of defining such a transformation. We consider another such transformation here, which has many of the same general properties as polarity. As with polarity, this transformation can be applied in arbitrary dimensions, but here we will consider it in the plane.

Let us begin by considering lines in the plane. Each line can be represented in a number of ways, but for now, let us assume the representation \( y = ax - b \), for some scalar values \( a \) and \( b \). (Why \(-b\) rather than \(+b\)? The distinction is unimportant, but it will simplify some of the notation defined below.) We cannot represent vertical lines in this way, and for now we will just ignore them. Later in the semester we will fix this up.

Therefore, in order to describe a line in the plane, you need only give its two coordinates \((a, b)\). In some sense, lines in the plane can be thought of as points in a new plane in which the coordinate axes are labeled \((a, b)\), rather than \((x, y)\). Thus the line \( y = 7x - 4 \) corresponds to the point \((7, 4)\) in this new plane. Each point in this new plane of “lines” corresponds to a nonvertical line in the original plane. We will call the original \((x, y)\)-plane the *primal plane* and the new \((a, b)\)-plane the *dual plane*.

What is the equation of a line in the dual plane? Since the coordinate system uses \( a \) and \( b \), we might write a line in a symmetrical form, for example \( b = 3a - 5 \), where the values 3 and 5 could be replaced by any scalar values.

Consider a particular point \( p = (p_x, p_y) \) in the primal plane, and consider the set of all nonvertical lines passing through this point. Any such line must satisfy the equation \( p_y = ap_x - b \). The images
of all these lines in the dual plane is a set of points:

\[ \mathcal{L} = \{ (a, b) \mid p_y = a p_x - b \} \]
\[ = \{ (a, b) \mid b = p_x a - p_y \}. \]

Notice that this set is just the set of points that lie on a line in the dual \((a, b)\)-plane. (This explains why it was convenient to negate \(b\) in the line equation.) Thus, not only do lines in the primal plane map to points in the dual plane, but points in the primal plane correspond to lines in the dual plane.

To make this all more formal, we can define a function that maps points in the primal plane to lines in the dual plane, and lines in the primal plane to points in the dual plane. We denote it using a asterisk (*) as a superscript. Thus, given point \(p = (p_x, p_y)\) and line \(\ell : (y = \ell_a x - \ell_b)\) in the primal plane we define \(\ell^*\) and \(p^*\) to be a point and line respectively in the dual plane defined by:

\[ \ell^* = (\ell_a, \ell_b) \]
\[ p^* : (b = p_x a - p_y). \]

![Figure 7.4: Dual transformation.](image)

We can define the same mapping from dual to primal as well.

**Properties of Point-Line Duality:** Duality has a number of interesting properties, each of which is easy to verify by substituting the definition and a little algebra.

**Self Inverse:** \((p^*)^* = p\).

**Order reversing:** Point \(p\) lies above/on/below line \(\ell\) in the primal plane if and only if line \(p^*\) passes below/on/above point \(\ell^*\) in the dual plane, respectively.

**Intersection preserving:** Lines \(\ell_1\) and \(\ell_2\) intersect at point \(p\) if and only if line \(p^*\) passes through points \(\ell^*_1\) and \(\ell^*_2\) in the dual plane.

**Collinearity/Coincidence:** Three points are collinear in the primal plane if and only if their dual lines intersect in a common point.

For example, to verify the order reversing property, observe that \(p\) lies above \(\ell\) if and only if

\[ p_y > \ell_a p_x - \ell_b. \]

Rewriting this gives

\[ \ell_b > p_x \ell_a - p_y, \]
which (in the dual plane) is equivalent to saying that $\ell^*$ lies above $p^*$, that is, $p^*$ passes below $\ell^*$. To finish things up, we need to make the connection between the upper convex hull of a set of points and the lower envelope of a set of lines.

The polarity transformation introduced earlier and the duality transformation presented here have both similar and different characteristics. For example, in the polarity transformation each point is mapped to a line that does not pass through the origin. In contrast, with the duality transformation given here it is possible to generate lines that pass through the origin, but only nonvertical lines are generated. In general, the choice of duality transformation is based on the application at hand.

**Convex Hulls and Lower Envelopes:** The following lemma demonstrates the, under the duality transformation, the convex hull problem is transformed into the lower envelope problem.

**Lemma:** Let $P$ be a set of points in the plane. The counterclockwise order of the points along the upper (lower) convex hull of $P$, is equal to the left-to-right order of the sequence of lines on the lower (upper) envelope of the dual $P^*$.

**Proof:** We will prove the result just for the upper hull and lower envelope, since the other case is symmetrical. For simplicity, let us assume that no three points are collinear. Observe that a necessary and sufficient condition for a pair of points $p_i p_j$ to form an edge on the upper convex hull is that the line $\ell_{ij}$ that passes through both of these points has every other point in $P$ strictly beneath it.

Consider the dual lines $p_i^*$ and $p_j^*$. A necessary and sufficient condition that these lines are adjacent on the lower envelope is that the dual point at which they meet, $\ell_{ij}^*$, lies beneath all of the other dual lines in $P^*$.

The order reversing condition of duality assures us that the primal condition occurs if and only if the dual condition occurs. Therefore, the sequence of edges on the upper convex hull is identical to the sequence of vertices along the lower envelope.

As we move counterclockwise along the upper hull observe that the slopes of the edges increase monotonically. Since the slope of a line in the primal plane is the $a$-coordinate of the dual point, it follows that as we move counterclockwise along the upper hull, we visit the lower envelope from left to right.

One rather cryptic feature of this proof is that, although the upper and lower hulls appear to be connected, the upper and lower envelopes of a set of lines appears to consist of two disconnected sets. To make sense of this, we should interpret the primal and dual planes from the perspective of projective geometry, and think of the rightmost line of the lower envelope as “wrapping around” to the leftmost line of the upper envelope, and vice versa.

Another interesting question is that of orientation. We know the orientation of three points is positive if the points have a counterclockwise orientation. What does it mean for three lines to have a positive orientation? (The definition of line orientation is exactly the same, in terms of a determinant of the coefficients of the lines.)
8 Linear Programming

Reading: Chapter 4 in the 4M’s.

Linear Programming: Last time we considered the problem of computing the intersection of \( n \) halfplanes, and presented an optimal \( O(n \log n) \) algorithm for this problem. In many applications it is not important to know the entire polygon (or generally the entire polyhedron in higher dimensions), but only to find one particular point of interest.

One particularly important application is that of linear programming. In linear programming (or LP for short) we are given a set of linear inequalities, or constraints, which we may think of as defining a (possibly empty, possibly unbounded) polyhedron in space, called the feasible region, and we are given a linear objective function, which is to be minimized or maximized subject to the given constraints. A typical description of a \( d \)-dimensional linear programming problem might be:

Maximize: \( c_1 x_1 + c_2 x_2 + \cdots + c_d x_d \)

Subject to:
\[
\begin{align*}
a_{1,1} x_1 + \cdots + a_{1,d} x_d & \leq b_1 \\
a_{2,1} x_1 + \cdots + a_{2,d} x_d & \leq b_2 \\
& \vdots \\
a_{n,1} x_1 + \cdots + a_{n,d} x_d & \leq b_n
\end{align*}
\]

where \( a_{i,j}, c_i, \) and \( b_i \) are given real numbers. This can be also be expressed in matrix notation:

Maximize: \( c^T x, \)

Subject to: \( Ax \leq b. \)

where \( c \) and \( x \) are \( d \)-vectors, \( b \) is an \( n \)-vector and \( A \) is an \( n \times d \) matrix.

From a geometric perspective, the feasible region is the intersection of halfspaces, and hence is a (possibly unbounded) convex polyhedron in \( d \)-space. We can think of the objective function as a vector \( \vec{c} \), and the problem is to find the point of the feasible region that is farthest in the direction \( \vec{c} \), called the optimal vertex.

![Figure 8.1: 2-dimensional linear programming.](image)

Note that the magnitude of \( \vec{c} \) is irrelevant, since the problem is unchanged for any positive scalar multiple of \( \vec{c} \). In many of our examples, we will imagine that the vector \( \vec{c} \) is pointing straight down (that is, \( \vec{c} = (0, -1) \)) and hence the problem is just that of finding the lowest point (minimum \( y \)-coordinate) of the feasible region. This involves no loss of generality, since it is always possible to transform the problem into one in which \( \vec{c} = (0, -1) \) by an appropriate rotation.

There are three possible outcomes of a given LP problem:
**Feasible:** The extreme point exists (and assuming general position is unique) as a vertex of the feasible region.

**Infeasible:** The feasible region is empty and hence there is no solution.

**Unbounded:** The feasible region is unbounded in the direction of the objective function, and so no finite optimal solution exists.

As mentioned above, without the assumption of general position, it is possible for a feasible LP to have an infinite number of (finite) optimal solutions of equal value, because an edge or face of the feasible region is orthogonal to the objective function vector. In such instances it is common to break ties in some way to produce a unique solution. For example, we might require that the solution be lexicographically maximal, that is, among all optimal solutions, select the one with the lexicographically maximum vector.

**Linear Programming in High Dimensional Spaces:** Linear programming is a very important technique used in solving large optimization problems. Typical instances may involve hundreds to thousands of constraints in very high dimensional space. It is without doubt one of the most important formulations of general optimization problems.

The principal methods used for solving high-dimensional linear programming problems are the simplex algorithm and various interior-point methods. The simplex algorithm works by finding a vertex on the feasible polyhedron, then walking edge by edge downwards until reaching a local minimum. By convexity, the local minimum is the global minimum. It has been long known that there are instances where the simplex algorithm runs in exponential time. The question of whether linear programming was even solvable in polynomial time was unknown until Khachiyan’s ellipsoid algorithm (late 70’s) and Karmarkar’s more practical interior-point algorithm (mid 80’s).

**2-dimensional LP:** We will restrict ourselves to low dimensional instances of linear programming. There are a number of interesting optimization problems that can be posed as a low-dimensional linear programming problem, or as closely related optimization problems. One which we will see later is the problem of finding a minimum radius circle that encloses a given set of \(n\) points.

Let us consider the problem just in the plane. Here we know that there is an \(O(n \log n)\) algorithm based on just computing the feasible polyhedron, and finding its lowest vertex. However, since we are only interested in one point on the polyhedron, it seems that we might hope to do better. Also, in \(d\) dimensional space, a polytope bounded by \(n\) halfspaces may have as many as \(\Omega(n^{\lfloor d/2 \rfloor})\) complexity.

We will show that 2-dimensional LP problems can be solved in \(O(n)\) time. This algorithm can be applied in any dimension \(d\), but the constant factor hidden by the asymptotic notation grows as \(d!\), and so it is not practical except for very small dimensions.

Assume that we are given a set of \(n\) linear inequalities (halfplanes) \(H = \{h_1, \ldots, h_n\}\) of the form:

\[
h_i : a_{i,x}x + a_{i,y}y \leq b_i,
\]

and a nonzero objective function given by vector \(\vec{c} = (c_x, c_y)\). The problem is to determine whether the LP is feasible and bounded, and if so, to compute the feasible point \(p = (p_x, p_y)\) that is feasible and (treating the point \(p\) as a vector) maximizes the dot product \(c_x p_x + c_y p_y\). Henceforth, we will assume that \(\vec{c} = (0, -1)\). (Note: As we decrease the \(y\)-coordinate of a point, we increase the value of the objective function. So, bigger is not better!)
The two algorithms that we will discuss for linear programming are very simple, and are based on a method called \textit{incremental construction}. One method will be deterministic and the other will be randomized. Incremental construction is among the most common design techniques in computational geometry, and this is another important reason for studying the linear programming problem. First, we consider the deterministic incremental algorithm.

\textbf{Deterministic Incremental Construction:} Let us suppose for now that the LP problem is bounded, and furthermore we can find two halfplanes whose intersection (a cone) is bounded with respect to the objective function. Our book explains how to overcome these assumptions in $O(n)$ additional time. Let us assume that the halfplanes are renumbered so that these are $h_1$ and $h_2$, and let $v_2$ denote this first optimum vertex, where the two lines associated with these halfplanes intersect. We will then add halfplanes one by one, $h_3, h_4, \ldots$, and with each addition we update the current optimum vertex. Let $v_i$ denote the optimal feasible vertex after the addition of $\{h_1, h_2, \ldots, h_i\}$. Our job is to update $v_i$ with each new addition. Notice that with each new constraint, the feasible region generally becomes smaller, and hence the value of the objective function at optimum vertex can only decrease.

There are two cases that can arise when $h_i$ is added (see the figure below). In the first case, $v_{i-1}$ lies within the halfplane $h_i$, and so it satisfies this constraint. If so, then it is easy to see that the optimum vertex does not change, that is $v_i = v_{i-1}$. In the second case $v_{i-1}$ violates constraint $h_i$. In this case we need to find a new optimum vertex. Let us consider this case in greater detail.

The important observation is that (assuming that the feasible region is not empty) the new optimum vertex must lie on the line that bounds $h_i$. Call this line $\ell_i$. Our book proves this formally, but here is an intuitive argument. Suppose that the new optimum vertex does not lie on $\ell_i$. Draw a line segment from $v_{i-1}$ to the new optimum. Observe (1) that as you walk along this segment the value of the objective function is decreasing monotonically (by linearity), and (2) that this segment must cross $\ell_i$ (because it goes from being infeasible with respect to $h_i$ to being feasible). Thus, it is maximized at the crossing point, which lies on $\ell_i$. Convexity and linearity are both very important for the proof.

So this leaves the question of how do we find the optimum vertex lying on line $\ell_i$. This turns out to be a 1-dimensional LP problem. Simply intersect each of the halfplanes with this line. Each intersection will take the form of a ray that lies on the line. We can think of each ray as representing an interval (unbounded to either the left or to the right). All we need to do is to intersect these intervals, and find the point that maximizes the objective function (that is, the lowest point). Computing the intersection of a collection of intervals, is very easy and can be solved in linear time. We just need to find the smallest upper bound and the largest lower bound. We select the point that maximizes the objective function. If this interval is empty, then it follows that the feasible region is empty, and we may
terminate and report that there is no solution.

Notice that we have solved a 2-dimensional LP problem by a reduction to a 1-dimensional LP (which is easy to solve). This general framework can be applied to solving LP problems in any dimension, by repeatedly reducing to an LP problem in the next lower dimension whenever the current optimum vertex is not feasible.

**Analysis:** What is the worst-case running time of this algorithm? There are roughly $n$ halfspace insertions. In step $i$, we may either find that the current optimum vertex is feasible, in which case the processing time is constant. On the other hand, if the current optimum vertex is infeasible, then we must solve a 1-dimensional linear program with $i - 1$ constraints. In the worst case, this second step occurs all the time, and the overall running time is given by the summation:

$$
\sum_{i=3}^{n} (i-1) \leq \sum_{i=1}^{n-1} i = \frac{n(n-1)}{2} = O(n^2).
$$

This follows from standard results on summations.

However notice that this worst-case is based on the rather pessimistic assumption that the current vertex is almost always infeasible. Next we will consider whether this assumption is reasonable, and how to better deal with this.

**Randomized Algorithm:** The $O(n^2)$ time is not very encouraging (considering that we could compute the entire feasible region in $O(n \log n)$ time). But we presented it because it leads to a very elegant randomized algorithm whose expected running time is $O(n)$. (Note that the expectation is over the random number sequences, and does not depend on any probabilistic assumptions on the distribution of the input.) The algorithm operates in exactly the same way, but we insert the halfplanes in random order, each time updating the answer based on the existing halfplanes. This is an example of a general class of algorithms called randomized incremental algorithms. There is only one difference between this algorithm and the deterministic one, namely, just prior to running the incremental algorithm, we call a procedure that randomly permutes the initial input list. A description is given in the code block below.

---

**Randomized 2D Linear Programming**

**Input:** Let $H$ be a set of $n$ halfplanes, and $\vec{c}$ be the objective function vector.

1. Randomly permute the halfplanes of $H$, letting $(h_1, h_2, \ldots, h_n)$ denote the resulting sequence. We assume that $h_1$ and $h_2$ define a bounded LP relative to $\vec{c}$. (See our text for the general case.) Let $v_2$ be the intersection point of the two associated lines.

2. For $i = 3$ to $n$ do:
   
   (a) If $(v_{i-1} \in h_i)$ then $v_i \leftarrow v_{i-1}$.
   
   (b) Otherwise, project $\{h_1, h_2, \ldots, h_{i-1}\}$ onto the line $l_i$ that supports $h_i$. Solve the resulting 1-dimensional LP.

      (i) If the 1-dimensional LP is infeasible, terminate and report that the LP is infeasible.

      (ii) Otherwise, let $v_i$ be the solution to the $1$-dimensional LP.

3. Return $v_n$ as the final solution.

---

**Backwards Analysis:** We analyze the running time in the expected case where we average over all $n!$ possible permutations. Each permutation has an equal probability of $1/n!$ of occurring, and an associated
running time. However, presenting the analysis as sum of $n!$ terms does not lead to something that we can easily simplify. We will apply a technique called \textit{backwards analysis}, which is quite useful.

To motivate how backwards analysis works, let us consider a much simpler example, namely the problem of computing the minimum of a set of $n$ distinct numbers. We permute the numbers and inspect them in this order. We maintain a variable that holds the minimum value seen so far. If we see a value that is smaller than the current minimum, then we update the minimum. The question we will consider is, on average how many times is the minimum value updated? For example, in the following sequence, the minimum is updated four times.

\[
5 \quad 9 \quad 4 \quad 2 \quad 6 \quad 8 \quad 0 \quad 3 \quad 1 \quad 7
\]

Let $p_i$ denote the probability that the minimum value changes on inspecting the $i$th number of the random permutation. Thus, with probability $p_i$ the minimum changes (and we add 1 to the counter for the number of changes) and with probability $1 - p_i$ it does not (and we add 0 to the counter for the number of changes). The total expected number of changes is

\[
C(n) = \sum_{i=1}^{n} (p_i \cdot 1 + (1 - p_i) \cdot 0) = \sum_{i=1}^{n} p_i.
\]

It suffices to compute $p_i$. We might be tempted to reason as follows. Let us consider a random subset of the first $i - 1$ values, and then consider all the possible choices for the $i$th value. However, this leads to a complicated analysis involving conditional probabilities. Let us instead consider an alternative approach, in which we work backwards. In particular, let us consider a random set of $i$ values, and consider the probability the last value added to this set resulted in a change in the minimum. To make this more formal, let $S_i$ be an arbitrary subset of $i$ numbers from our initial set of $n$. (In theory, the probability is conditional on the fact that the elements of $S_i$ represent the first $i$ elements to be chosen, but since the analysis will not depend on the particular choice of $S_i$, it follows that the probability that we compute will hold unconditionally.) Among all $i!$ permutations of the elements of $S_i$, in how many of these does the minimum change in the transition from the $(i - 1)$-st stage to the $i$th stage? The key observation is that the minimum only changes for those sequences in which the minimum item appears with equal probability in each of the $i$ positions of a random sequence, the probability that it appears last is exactly $1/i$. Thus, $p_i = 1/i$. From this we have

\[
C(n) = \sum_{i=1}^{n} p_i = \sum_{i=1}^{n} \frac{1}{i} = \ln n + O(1).
\]

This summation $\sum (1/i)$ is called the \textit{Harmonic series} and the fact that it is nearly equal to $\ln n$ is a well known fact. (See any text on probability theory.)

This is called an \textit{backwards analysis} because the analysis works by consider the possible random transitions from $S_i$ back to $S_{i-1}$, as opposed to working forward from $S_{i-1}$ to $S_i$. Of course, the probabilities are no different whether we consider the random sequence backwards rather than forwards, so this is a perfectly accurate analysis. It’s just simpler and easier to understand.
**Backwards Analysis for Randomized LP:** Let us apply this same approach to the analysis of the running time of the randomized incremental linear programming algorithm. This time, let $p_i$ denote the probability that the insertion of the $i$th hyperplane in the random order resulted in a change in the optimum vertex. With probability $(1 - p_i)$ there is no change (Case 1), and it takes us $O(1)$ time to determine this. ($O(d)$ time in general in dimension $d$.) With probability $p_i$ we need to invoke a 1-dimensional LP on a set of $i - 1$ halfplanes in dimension 1, with a running time of $O(i)$. (In general, if $T(n, d)$ is the expected running time for $n$ halfspaces in dimension $d$, then this cost would be $T(i - 1, d - 1)$.) Combining this we have a total expected running time of

$$T(n) = \sum_{i=1}^{n} ((1 - p_i) \cdot 1 + p_i \cdot i) \leq n + \sum_{i=1}^{n} p_i \cdot i$$

All that remains is to determine $p_i$. We will apply the same technique. Let $S_i$ denote an arbitrary subset consisting of $i$ of the original halfplanes. Among all $i!$ permutations of $S_i$, in how many does the optimum vertex change with the $i$th step? Let $v_i$ denote the optimum vertex for these $i$ halfplanes. It is important to note that $v_i$ only depends on the set $S_i$ and not on the order of their insertion. (If you do not see why this is important, think about it.)

Assuming general position, there are two halfplanes $h'$ and $h''$ of $S_i$ passing through $v_i$. If neither of these was the last to be inserted, then $v_i = v_{i-1}$, and there is no change. If either $h'$ or $h''$ was the last to be inserted, then $v_i$ did not exist yet, and hence the optimum must have changed as a result of this insertion. Thus, the optimum changes if and only if either $h'$ or $h''$ was the last halfplane inserted. Since all of the $i$ halfplanes are equally likely to be last, this happens with probability $2/i$. Therefore, $p_i = 2/i$.

To illustrate this, consider the example shown in the following figure. We have $i = 7$ random halfplanes that have been added, so far and $v_i = v$ is the current optimum vertex and it is defined by $h_4$ and $h_8$. Let’s consider which halfplane was added last. Suppose that $h_5$ was the last halfplane to be added. Consider the state of affairs with all the halfplanes, but without $h_5$. Prior to this $v$ would already have been the optimum vertex. Therefore, if $h_5$ was added last, it would have been added with $O(1)$ cost. (There was nothing special about $h_5$. The same would be true for any of the five halfplanes other than $h_4$ and $h_8$.)

![Backwards analysis](https://via.placeholder.com/150)

**Figure 8.3:** Backwards analysis.

On the other hand, if $h_8$ was added last, then prior to its addition, we would have had a different optimum vertex, namely $v'$. (And the optimum would also change if $h_4$ was added last.) In summary,
in 2 out of 7 cases \((h_4 \text{ and } h_8)\) on the \(i\)th insertion we need to solve a 1-dimensional LP in \(O(i)\) time and in 5 out of 7 instances \(O(1)\) time suffices.

Returning to our analysis, since \(p_i = 2/i\) we have

\[
T(n) \leq n + \sum_{i=1}^{n} p_i \cdot i = n + \sum_{i=1}^{n} \frac{2i}{i} = n + 2n = O(n).
\]

Therefore, the expected running time is linear in \(n\). I’ll leave the analysis for the \(d\)-dimensional case as an exercise. (Note that in dimension \(d\) the optimum vertex is defined by \(d\) halfspaces, not 2.)
9 Planar Point Location and Trapezoidal Maps

Reading: Chapter 6 of the 4M’s.

Point Location: The point location problem (in 2-space) is: given a polygonal subdivision of the plane (that is, a cell complex bounded by line segment edges) with \( n \) vertices, preprocess this subdivision so that given a query point \( q \), we can efficiently determine which face of the subdivision contains \( q \). We may assume that each face has some identifying label, which is to be returned. We also assume that the subdivision is represented in any “reasonable” form. (We will discuss such representations in a different lecture.) In general \( q \) may coincide with an edge or vertex. To simplify matters, we will assume that \( q \) does not lie on an edge or vertex, but these special cases are not hard to handle.

Our goal is to develop a data structure of \( O(n) \) space that can answer queries in \( O(\log n) \) time. For many years the best methods known had an extra log factor, either in the space or in the query time. Kirkpatrick achieved a breakthrough by presenting a time/space optimal algorithm. Kirkpatrick’s algorithm has fairly high constant factors. Somewhat simpler and more practical optimal algorithms were discovered since then. We will present perhaps the simplest and most practical of the known optimal algorithms. The method is based on a randomized incremental construction, the same technique used in our linear programming algorithm.

Trapezoidal Map: The algorithm is based on a structure called a trapezoidal map, which also goes under many other names in the computational geometry literature. As mentioned above, we normally think of the input to a point location algorithm as being a planar polygonal subdivision, but for this problem we make the weaker assumption that the input is simply a set of nonvertical line segments, denoted \( S = \{s_1, s_2, \ldots, s_n\} \). We assume these line segments do not intersect except possibly at their endpoints. In this context the answer to a point location query is the index of the line segment of \( S \) lying immediately below the query point (or some special value if no segment is below the query point). Clearly, the original formulation of the point location problem can be easily recast in this manner.

To define the trapezoidal map, imagine shooting a bullet vertically upwards and downwards from each vertex in the polygonal subdivision until it hits another segment of \( S \). The resulting “bullet paths”, together with the initial line segments define the trapezoidal map. To avoid infinite bullet paths at the top and bottom of the subdivision, we may assume that the initial subdivision is contained entirely within a large bounding rectangle. (An example is shown in the figure below.)

![Figure 9.1: Trapezoidal map.](image-url)

In addition to assuming no vertical segments, we will make the additional general position assumption that no two distinct endpoints have the same \( x \)-coordinate. As usual, this assumption is easy to overcome, but complicates the description of the algorithm. Although the faces of this subdivision are
trapezoidal in shape, they may bounded by an arbitrarily large number of vertices and edges resulting from the vertices from surrounding trapezoids.

Observe that all the faces of the resulting subdivision are trapezoids with vertical sides. The left or right side might degenerate to a line segment of length zero, implying that the resulting trapezoid degenerates to a triangle. We claim that the process of converting an arbitrary polygonal subdivision into a trapezoidal decomposition increases its size by at most a constant factor. Actually this follows from the facts that we only increase the number of vertices by a constant factor and the graph is planar. But since constant factor expansions in space are significant, it is a good idea to work this through carefully.

Claim: Given a polygonal subdivision with \( n \) segments, the resulting trapezoidal map has at most \( 6n + 4 \) vertices and \( 3n + 1 \) trapezoids.

Proof: To prove the bound on the number of vertices, observe that each vertex shoots two bullet paths, each of which will result in the creation of a new vertex. Thus each original vertex gives rise to three vertices in the final map. Since each segment has two vertices, this implies at most \( 6n \) vertices.

To bound the number of trapezoids, observe that for each trapezoid in the final map, its left side (and its right as well) is bounded by a vertex of the original polygonal subdivision. The left endpoint of each line segment can serve as the left bounding vertex for two trapezoids (one above the line segment and the other below) and the right endpoint of a line segment can serve as the left bounding vertex for one trapezoid. Thus each segment of the original subdivision gives rise to at most three trapezoids, for a total of \( 3n \) trapezoids. The last trapezoid is the one bounded by the left side of the bounding box.

An important fact to observe about each trapezoid is that it is \textit{defined} (that is, its existence is determined) by exactly four entities from the original subdivision: a segment on top, a segment on the bottom, a bounding vertex on the left, and a bounding vertex on the right. This simple observation will play an important role later in the analysis.

Trapezoidal decompositions, like triangulations, are interesting data structures in their own right. It is another example of the idea of converting a complex shape into a disjoint collection of simpler objects. The fact that the sides are vertical makes trapezoids simpler than arbitrary quadrilaterals. Finally observe that the trapezoidal decomposition is a refinement of the original polygonal subdivision, and so once we know which face of the trapezoidal map a query point lies in, we will know which face of the original subdivision it lies in (either implicitly, or because we label each face of the trapezoidal map in this way).

Construction: We could construct the trapezoidal map easily by plane sweep. (This should be an easy exercise by this point, but think about how you would do it.) We will build the trapezoidal map by a randomized incremental algorithm, because the point location algorithm is based on this construction. (In fact, historically, this algorithm arose as a method for computing the trapezoidal decomposition of a collection of intersecting line segments, and the point location algorithm arose as an artifact that was needed in the construction.)

The incremental algorithm starts with the initial bounding rectangle (that is, one trapezoid) and then we add the segments of the polygonal subdivision one by one in random order. As each segment is
added, we update the trapezoidal map. Let $S_i$ denote the subset consisting of the first $i$ (random) segments, and let $T_i$ denote the resulting trapezoidal map.

To perform the update this we need to know which trapezoid the left endpoint of the segment lies in. We will address this question later, since it will be answered by the point location algorithm itself. Then we trace the line segment from left to right, determining which trapezoids it intersects. Finally, we go back to these trapezoids and “fix them up”. There are two things that are involved in fixing. First, the left and right endpoints of the new segment need to have bullets fired from them. Second, one of the earlier bullet paths might hit this line segment. When that happens the bullet path must be trimmed back. (We know which vertices are from the original subdivision vertices, so we know which side of the bullet path to trim.) The process is illustrated in the figure below.

![Figure 9.2: Incremental update.](image)

Observe that the structure of the trapezoidal decomposition does not depend on the order in which the segments are added. This observation will be important for the probabilistic analysis. The following is also important to the analysis.

**Claim:** Ignoring the time spent to locate the left endpoint of an segment, the time that it takes to insert the $i$th segment and update the trapezoidal map is $O(k_i)$, where $k_i$ is the number of newly created trapezoids.

**Proof:** Consider the insertion of the $i$th segment, and let $K$ denote the number of bullet paths that this segment intersects. We need to shoot four bullets (two from each endpoint) and then trim each of the $K$ bullet paths, for a total of $K + 4$ operations that need to be performed. If the new segment did not cross any of the bullet paths, then we would get exactly four new trapezoids. For each of the $K$ bullet paths we cross, we add one more to the number of newly created trapezoids, for a total of $K + 4$. Thus, letting $k_i = K + 4$ be the number of trapezoids created, the number of update operations is exactly $k_i$. Each of these operations can be performed in $O(1)$ time given any reasonable representation of the trapezoidal map as a planar subdivision. (For example, the *doubly connected edge list*, presented in Chapter 2 of the 4M’s).

**Analysis:** Let us consider the expected time to build the trapezoidal map. Later we will show that the time needed to determine the trapezoid containing the left endpoint of each new segment is $O(\log n)$ time on average. We will show that the expected time to add each new segment is $O(1)$. Since there are $n$ insertions, this will lead to a total expected time complexity of $O(n(1 + \log n)) = O(n \log n)$. As always, the expectation is over the sequence of random numbers, and does not depend on the distribution of the line segments.

We know that the size of the final trapezoidal map is $O(n)$. It turns out that the total size of the point location data structure will actually be proportional to the number of new trapezoids that are
created with each insertion. In the worst case, when we add the $i$th segment, it might cut through a large fraction of the existing $O(i)$ trapezoids, and this would lead to a total size proportional to $\sum_{i=1}^{n} i = \Theta(n^2)$. However, the magic of the incremental construction is that this does not happen very often on average. We will show that on average, each insertion results in only a constant number of trapezoids being created.

This is rather surprising at first. Clearly if the segments are short, then each segment might not intersect very many trapezoids. But if most of the segments are long, it might seem that almost every insertion cuts through $O(n)$ trapezoids. What saves us is that, although a long segment might cut through many trapezoids, it shields later segments from cutting through many trapezoids.

**Lemma:** Consider the randomized incremental construction of a trapezoidal map, and let $k_i$ denote the number of new trapezoids created when the $i$th segment is added. Then $E(k_i) = O(1)$, where the expectation is taken over all permutations of the segments.

**Proof:** The analysis will be based on a backwards analysis. Recall that such an analysis is based on analyzing the expected value assuming that the last insertion was random.
Let $T_i$ denote the trapezoidal map after the insertion of the $i$th segment. Because we are averaging over all permutations, among the $i$ segments that are present in $T_i$, each one has an equal probability $1/i$ of being the last one to have been added. For each of the segments $s$ we want to count the number of trapezoids that would have been created, had $s$ been the last segment to be added. Let’s say that a trapezoid $\Delta$ depends on an segment $s$, if $s$ would have caused $\Delta$ to be created, had $s$ been added last. We want to count the number of trapezoids that depend on each segment, and then compute the average over all segments. If we let $\delta(\Delta, s) = 1$ if segment $\Delta$ depends on $s$, and 0 otherwise, then the expected value is

$$E(k_i) = \frac{1}{i} \sum_{s \in S_i} (\text{no. of trapezoids that depend on } s) = \frac{1}{i} \sum_{s \in S_i} \sum_{\Delta \in T_i} \delta(\Delta, s).$$

![Figure 9.3: Trapezoid-segment dependencies.](image)

Some segments might have resulted in the creation of lots of trapezoids and other very few. How do we get a handle on this quantity? The trick is, rather than count the number of trapezoids that depend on each segment, we count the number segments that each trapezoid depends on. (This is the old combinatorial trick of reversing the order of summation.) In other words we can express the above quantity as:

$$E(k_i) = \frac{1}{i} \sum_{\Delta \in T_i} \sum_{s \in S_i} \delta(\Delta, s).$$
This quantity is much easier to analyze. In particular, each trapezoid is bounded by at most four sides. (The reason it is “at most” is that degenerate trapezoids are possible which may have fewer sides.) The top and bottom sides are each determined by a segment of $S_i$, and clearly if either of these was the last to be added, then this trapezoid would have come into existence as a result. The left and right sides are each determined by an endpoint of a segment in $S_i$, and clearly if either of these was the last to be added, then this trapezoid would have come into existence. (There is a bit of a subtlety here. What if multiple segments share the endpoint? Note that the trapezoid is only dependent on the first such segment to be added, since this is the segment that caused the vertex to come into existence. Also note that the same segment that forms the top or bottom side might also provide the left or right endpoint. These considerations only decrease the number of segments on which a trapezoid depends.)

In summary, each trapezoid is dependent on at most four segments, implying that $\sum_{s \in S_i} \delta(\Delta, s) \leq 4$. Since $T_i$ consists of $O(i)$ trapezoids we have

$$E(k_i) \leq \frac{1}{i} \sum_{\Delta \in T_i} 4 = \frac{1}{i} 4|T_i| = \frac{1}{i} 4O(i) = O(1).$$

Since the expected number of new trapezoids created with each insertion is $O(1)$, it follows that the total number of trapezoids that are created in the entire process is $O(n)$. This fact is important in bounding the total time needed for the randomized incremental algorithm.

The only question that we have not considered in the construction is how to locate the trapezoid that contains left endpoint of each newly added segment. We will consider this question, and the more general question of how to do point location next time.
10  More on Planar Point Location

Reading: Chapter 6 of the 4M’s.

Point Location: Last time we presented a randomized incremental algorithm for building a trapezoidal map. Today we consider how to modify this algorithm to answer point location queries for the resulting trapezoidal decomposition. The preprocessing time will be $O(n \log n)$ in the expected case (as was the time to construct the trapezoidal map), and the space and query time will be $O(n)$ and $O(\log n)$, respectively, in the expected case. Note that this may be applied to any spatial subdivision, by treating it as a set of line segments, and then building the resulting trapezoidal decomposition and using this data structure.

Recall that we treat the input as a set of segments $S = \{s_1, \ldots, s_n\}$ (permuted randomly), that $S_i$ denotes the subset consisting of the first $i$ segments of $S$, and $T_i$ denotes the trapezoidal map of $S_i$.

One important element of the analysis to remember from last time is that each time we add a new line segment, it may result in the creation of the collection of new trapezoids, which were said to depend on this line segment. We presented a backwards analysis that the number of new trapezoids that are created with each stage is expected to be $O(1)$. This will play an important role in today’s analysis.

Point Location Data Structure: The point location data structure is based on a rooted directed acyclic graph. Each node will either two or zero outgoing edges. Nodes with zero outgoing edges are called leaves. There will be one leaf for each trapezoid in the map. The other nodes are called internal nodes, and they are used to guide the search to the leaves. This is not a binary tree, however, because subtrees may be shared.

There are two types of internal nodes, $x$-nodes and $y$-nodes. Each $x$-node contains the $x$-coordinate $x_0$ of an endpoint of one of the segments. Its two children correspond to the points lying to the left and to the right of the vertical line $x = x_0$. Each $y$-node contains a pointer to a line segment of the subdivision. The left and right children correspond to whether the query point is above or below the line containing this segment, respectively. (Don’t be fooled by the name—$y$-node comparisons depend on both the $x$ and $y$ values of the query point.) Note that the search will reach a $y$-node only if we have already verified that the $x$-coordinate of the query point lies within the vertical slab that contains this segment.

Our construction of the point location data structure mirrors the incremental construction of the trapezoidal map. In particular, if we freeze the construction just after the insertion of any segment, the current structure will be a point location structure for the current trapezoidal map. In the figure below we show a simple example of what the data structure looks like for two line segments. The circular nodes are the $x$-nodes and the hexagonal nodes are the $y$-nodes. There is one leaf for each trapezoid. The $y$-nodes are shown as hexagons. For example, if the query point is in trapezoid $D$, we would first detect that it is to the right of endpoint $p_1$ (right child), then left of $q_1$ (left child), then below $s_1$ (right child), then right of $p_2$ (right child), then above $s_2$ (left child).

Incremental Construction: The question is how do we build this data structure incrementally? First observe that when a new line segment is added, we only need to adjust the portion of the tree that involves the trapezoids that have been deleted as a result of this new addition. Each trapezoid that is deleted will be replaced with a search structure that determines the newly created trapezoid that contains it.
Suppose that we add a line segment $s$. This results in the replacement of an existing set of trapezoids with a set of new trapezoids. As a consequence, we will replace the leaves associated with each such deleted trapezoid with a small search structure, which locates the new trapezoid that contains the query point. There are three cases that arise, depending on how many endpoints of the segment lie within the current trapezoid.

**Single (left or right) endpoint:** A single trapezoid $A$ is replaced by three trapezoids, denoted $X$, $Y$, and $Z$. Letting $p$ denote the endpoint, we create an $x$-node for $p$, and one child is a leaf node for the trapezoid $X$ that lies outside vertical projection of the segment. For the other child, we create a $y$-node whose children are the trapezoids $Y$ and $Z$ lying above and below the segment, respectively. (See the figure below left.)

**No segment endpoints:** This happens when the segment cuts completely through a trapezoid. A single trapezoid is replaced by two trapezoids, one above and one below the segment, denoted $Y$ and $Z$. We replace the leaf node for the original trapezoid with a $y$-node whose children are leaf nodes associated with $Y$ and $Z$. (This case is not shown in the figure.)

**Two segment endpoints:** This happens when the segment lies entirely inside the trapezoid. In this case one trapezoid $A$ is replaced by four trapezoids, $U$, $X$, $Y$, and $Z$. Letting $p$ and $q$ denote the left and right endpoints of the segment, we create two $x$-nodes, one for the $p$ and the other for $q$. We create a $y$-node for the line segment, and join everything together as shown in the figure.

It is important to notice that (through sharing) each trapezoid appears exactly once as a leaf in the resulting structure. An example showing the complete transformation to the data structure after adding a single segment is shown in the figure below.

![Figure 10.1: Trapezoidal map point location data structure.](image)

![Figure 10.2: Line segment insertion and updates to the point location structure. The single-endpoint case (left) and the two-endpoint case (right). The no-endpoint case is not shown.](image)
**Analysis:** We claim that the size of the point location data structure is $O(n)$ and the query time is $O(\log n)$, both in the expected case. As usual, the expectation depends only on the order of insertion, not on the line segments or the location of the query point.

To prove the space bound of $O(n)$, observe that the number of new nodes added to the structure with each new segment is proportional to the number of newly created trapezoids. Last time we showed that with each new insertion, the expected number of trapezoids that were created was $O(1)$. Therefore, we add $O(1)$ new nodes with each insertion in the expected case, implying that the total size of the data structure is $O(n)$.

Analyzing the query time is a little subtler. In a normal probabilistic analysis of data structures we think of the data structure as being fixed, and then compute expectations over random queries. Here the approach will be to imagine that we have exactly one query point to handle. The query point can be chosen arbitrarily (imagine an adversary that tries to select the worst possible query point) but this choice is made without knowledge of the random choices the algorithm makes. We will show that given a fixed query point, the expected search path length for this one query point is $O(\log n)$, where the expectation is over all segment insertion orders. (Note that this does not imply that the expected maximum depth of the tree is $O(\log n)$. We will discuss this below.)

Let $q$ denote the query point. Rather than consider the search path for $q$ in the final search structure, we will consider how $q$ moves incrementally through the structure with the addition of each new line segment. Let $\Delta_i$ denote the trapezoid of the map that $q$ lies in after the insertion of the first $i$ segments. Observe that if $\Delta_{i-1} = \Delta_i$, then insertion of the $i$th segment did not affect the trapezoid that $q$ was in, and therefore $q$ will stay where it is relative to the current search structure. (For example, if $q$ was in trapezoid $B$ prior to adding $s_3$ in the figure above, then the addition of $s_3$ does not incur any additional cost to locating $q$.) However, if $\Delta_{i-1} \neq \Delta_i$, then the insertion of the $i$th segment caused $q$’s trapezoid to be deleted. As a result, $q$ must locate itself with respect to the newly created trapezoids that overlap $\Delta_{i-1}$. Since there are a constant number of such trapezoids (at most four), there will be $O(1)$ work needed to locate $q$ with respect to these. In particular, $q$ may fall as most three levels in

![Diagram](image-url)
the search tree. (The worst case occurs in the two-endpoint case, if the query point falls into one of the trapezoids $X$ or $Y$ lying above or below the segment.)

Since a point can fall at most 3 levels with each change of its containing trapezoid, the expected length of the search path is at most 3 times the sum of probabilities that $q$ changes its trapezoid as a result of each insertion. For $1 \leq i \leq n$, let $P_i$ denote this probability (taken over random insertion orders, irrespective of the choice of $q$). Letting $D(q)$ denote the average depth of $q$ in the final search tree, we have

$$D(q) \leq 3 \sum_{i=1}^{n} P_i.$$  

We will show below that $P_i \leq 4/i$. From this it will follow that the expected path length is at most

$$D(q) \leq 3 \sum_{i=1}^{n} \frac{4}{i} = 12 \sum_{i=1}^{n} \frac{1}{i}.$$  

Recall that $\sum_{i=1}^{n} \frac{1}{i}$ is the Harmonic series, and for large $n$, its value is very nearly $\ln n$. Thus we have

$$D(q) \leq 12 \ln n = O(\log n).$$

To show that $P_i \leq 4/i$, we apply a backwards analysis. In particular, consider the trapezoid that contains $q$ after the $i$th insertion. Recall from last time that this trapezoid is dependent on at most four segments, which define the top and bottom edges, and the left and right sides of the trapezoid. Since each segment is equally likely to be the last segment to have been added, the probability that the last insertion caused $q$ to belong to a new trapezoid is at most $4/i$. This completes the proof.

**Guarantees on Search Time:** One shortcoming with this analysis is that even though the search time is provably small in the expected case for a given query point, it might still be the case that once the data structure has been constructed there is a single very long path in the search structure, and the user repeatedly performs queries along this path. Hence, the analysis provides no guarantees on the running time of all queries.

Although we will not prove it, the book presents a stronger result, namely that the length of the maximum search path is also $O(\log n)$ with high probability. In particular, they prove the following.

**Lemma:** Given a set of $n$ non-crossing line segments in the plane, and a parameter $\lambda > 0$, the probability that the total depth of the randomized search structure exceeds $3\lambda \ln(n + 1)$, is at most $2/(n + 1)^{\lambda \ln(1.25 - \delta)}$.

For example, for $\lambda = 20$, the probability that the search path exceeds $60 \ln(n + 1)$ is at most $2/(n + 1)^{1.5}$. (The constant factors here are rather weak, but a more careful analysis leads to a better bound.) Nonetheless, this itself is enough to lead to variant of the algorithm for which $O(\log n)$ time is guaranteed. Rather than just running the algorithm once and taking what it gives, instead run it repeatedly and keep track of the structure’s depth as you go. As soon as the depth exceeds $c \log n$ for some suitably chosen $c$, then stop and start over again with a new random sequence. For a suitable $c$, the above lemma implies that such a failure will occur with at most some very small constant probability. Therefore, after a constant number of trials, we will succeed in constructing a data structure of the desired depth bound. A similar argument can be applied to the space bounds.
**Theorem:** Given a set of $n$ non-crossing line segments in the plane, in expected $O(n \log n)$ time, it is possible to construct a point location data structure of (worst case) size $O(n)$ that can answer point location queries in (worst case) time $O(\log n)$.

**Line Segment Intersection Revisited:** Earlier this semester we presented a plane-sweep algorithm for computing line segment intersection. The algorithm had a running time of $O((n + I) \log n)$, where $I$ is the number of intersection points. It is interesting to note that the randomized approach we discussed today can be adapted to deal with intersecting segments as well. In particular, whenever a segment is added, observe that in addition to it stabbing vertical segments, it may generally cross over one of the existing segments. When this occurs, the algorithm must determine the trapezoid that is hit on the other side of the segment, and then continue the process of walking the segment. Note that the total size of the final decomposition is $O(n + I)$, which would suggest that the running time might be the same as the plane-sweep algorithm. It is remarkable, therefore, that the running time is actually better. Intuitively, the reason is that the $O(\log n)$ factor in the randomized algorithm comes from the point location queries, which are applied only to the left endpoint of each of the $n$ segments. With a bit of additional work, it can be shown that the adaptation of the randomized algorithm to general (intersecting) segments runs in $O(I + n \log n)$ time, thus removing the log factor from the $I$ term.
11 Voronoi Diagrams and Fortune’s Algorithm

Reading: Chapter 7 in the 4M’s.

Voronoi Diagrams: Voronoi diagrams are among the most important structures in computational geometry. A Voronoi diagram encodes proximity information, that is, what is close to what. Let \( P = \{p_1, p_2, \ldots, p_n\} \) be a set of points in the plane (or in any dimensional space), which we call sites. Define \( \mathcal{V}(p_i) \), the Voronoi cell for \( p_i \), to be the set of points \( q \) in the plane that are closer to \( p_i \) than to any other site. That is, the Voronoi cell for \( p_i \) is defined to be:

\[
\mathcal{V}(p_i) = \{ q \mid \|p_iq\| < \|p_jq\|, \forall j \neq i \},
\]

where \( \|pq\| \) denotes the Euclidean distance between points \( p \) and \( q \). The Voronoi diagram can be defined over any metric and in any dimension, but we will concentrate on the planar, Euclidean case here.

Another way to define \( \mathcal{V}(p_i) \) is in terms of the intersection of halfplanes. Given two sites \( p_i \) and \( p_j \), the set of points that are strictly closer to \( p_i \) than to \( p_j \) is just the open halfplane whose bounding line is the perpendicular bisector between \( p_i \) and \( p_j \). Denote this halfplane \( h(p_i, p_j) \). It is easy to see that a point \( q \) lies in \( \mathcal{V}(p_i) \) if and only if \( q \) lies within the intersection of \( h(p_i, p_j) \) for all \( j \neq i \). In other words,

\[
\mathcal{V}(p_i) = \cap_{j \neq i} h(p_i, p_j).
\]

Since the intersection of halfplanes is a (possibly unbounded) convex polygon, it is easy to see that \( \mathcal{V}(p_i) \) is a (possibly unbounded) convex polygon. Finally, define the Voronoi diagram of \( P \), denoted \( \text{Vor}(P) \) to be what is left of the plane after we remove all the (open) Voronoi cells. It is not hard to prove (see the text) that the Voronoi diagram consists of a collection of line segments, which may be unbounded, either at one end or both. An example is shown in the figure below.

![Voronoi diagram](image)

Voronoi diagrams have a number of important applications. These include:

**Nearest neighbor queries:** One of the most important data structures problems in computational geometry is solving nearest neighbor queries. Given a point set \( P \), and given a query point \( q \), determine the closest point in \( P \) to \( q \). This can be answered by first computing a Voronoi diagram and then locating the cell of the diagram that contains \( q \). (We have already discussed point location algorithms.)

**Computational morphology and shape analysis:** A useful structure in shape analysis is called the medial axis. The medial axis of a shape (e.g., a simple polygon) is defined to be the union of the center points of all locally maximal disks that are contained within the shape. If we generalize
the notion of Voronoi diagram to allow sites that are both points and line segments, then the
medial axis of a simple polygon can be extracted easily from the Voronoi diagram of these
generalized sites.

Figure 11.2: A simple polygon (left) and its medial axis (right), shown with a sample maximal disk.

**Facility location:** We want to open a new Blockbuster or Starbucks. It should be placed as far as
possible from any existing video stores. Where should it be placed? It turns out that the vertices
of the Voronoi diagram are the points that are locally at maximum distances from any other point
in the set, and hence they form a natural set of candidate locations.

**Neighbors and Interpolation:** Given a set of measured height values over some geometric terrain.
Each point has \((x, y)\) coordinates and a height value. We would like to interpolate the height
value of some query point that is not one of our measured points. To do so, we would like
to interpolate its value from neighboring measured points. One way to do this, called *natural
neighbor interpolation*, is based on computing the Voronoi neighbors of the query point, assuming
that it has one of the original set of measured points.

**Properties of the Voronoi diagram:** Here are some observations about the structure of Voronoi diagrams
in the plane.

**Voronoi complex:** Clearly the diagram is a cell complex whose faces are (possibly unbounded) con-
 vex polygons. Each point on an edge of the Voronoi diagram is equidistant from its two nearest
neighbors \(p_i\) and \(p_j\). Thus, there is a circle centered at such a point such that \(p_i\) and \(p_j\) lie on this
circle, and no other site is interior to the circle.

Figure 11.3: Properties of the Voronoi diagram.

**Voronoi vertices:** It follows that the vertex at which three Voronoi cells \(V(p_i)\), \(V(p_j)\), and \(V(p_k)\)
intersect, called a *Voronoi vertex* is equidistant from all sites. Thus it is the center of the circle
passing through these sites, and this circle contains no other sites in its interior.

**Degree:** Generally three points in the plane define a unique circle. If we make the general position
assumption that no four sites are cocircular, then the vertices of the Voronoi diagram all have
degree three.
Convex hull: A cell of the Voronoi diagram is unbounded if and only if the corresponding site lies on the convex hull. (Observe that a site is on the convex hull if and only if it is the closest point from some point at infinity.) Thus, given a Voronoi diagram, it is easy to extract the convex hull in linear time.

Size: If $n$ denotes the number of sites, then the Voronoi diagram is a planar graph (if we imagine all the unbounded edges as going to a common vertex infinity) with exactly $n$ faces. It follows from Euler’s formula that the number of Voronoi vertices is at most $2n - 5$ and the number of edges is at most $3n - 6$. (See the text for details. In higher dimensions the diagram’s combinatorial complexity ranges from $O(n)$ up to $O(n^{\lceil d/2 \rceil})$.)

Computing Voronoi Diagrams: There are a number of algorithms for computing the Voronoi diagram of a set of $n$ sites in the plane. Of course, there is a naive $O(n^2 \log n)$ time algorithm, which operates by computing $\mathcal{V}(p_i)$ by intersecting the $n-1$ bisector halfplanes $h(p_i, p_j)$, for $j \neq i$. However, there are much more efficient ways, which run in $O(n \log n)$ time. Since the convex hull can be extracted from the Voronoi diagram in $O(n)$ time, it follows that this is asymptotically optimal in the worst-case.

Historically, $O(n^2)$ algorithms for computing Voronoi diagrams were known for many years (based on incremental constructions). When computational geometry came along, a more complex, but asymptotically superior $O(n \log n)$ algorithm was discovered. This algorithm was based on divide-and-conquer. But it was rather complex, and somewhat difficult to understand. Later, Steven Fortune invented a plane sweep algorithm for the problem, which provided a simpler $O(n \log n)$ solution to the problem. It is his algorithm that we will discuss. Somewhat later still, it was discovered that the incremental algorithm is actually quite efficient, if it is run as a randomized incremental algorithm. We will discuss this algorithm later when we talk about the dual structure, called a Delaunay triangulation.

Fortune’s Algorithm: Before discussing Fortune’s algorithm, it is interesting to consider why this algorithm was not invented much earlier. In fact, it is quite a bit trickier than any plane sweep algorithm we have seen so far. The key to any plane sweep algorithm is the ability to discover all upcoming events in an efficient manner. For example, in the line segment intersection algorithm we considered all pairs of line segments that were adjacent in the sweep-line status, and inserted their intersection point in the queue of upcoming events. The problem with the Voronoi diagram is that of predicting when and where the upcoming events will occur. Imagine that you are designing a plane sweep algorithm. Behind the sweep line you have constructed the Voronoi diagram based on the points that have been encountered so far in the sweep. The difficulty is that a site that lies ahead of the sweep line may generate a Voronoi vertex that lies behind the sweep line. How could the sweep algorithm know of the existence of this vertex until it sees the site. But by the time it sees the site, it is too late. It is these unanticipated events that make the design of a plane sweep algorithm challenging. (See the figure below.)

Fortune made the clever observation of rather than computing the Voronoi diagram through plane sweep in its final form, instead to compute a “distorted” but topologically equivalent version of the diagram. This distorted version of the diagram was based on a transformation that alters the way that distances are measured in the plane. The resulting diagram has the same topological structure as the Voronoi diagram, but its edges are parabolic arcs, rather than straight line segments. Once this distorted diagram is generated, it was an easy matter to “undistort” it in $O(n)$ time to produce the correct Voronoi diagram.
Our presentation will be different from Fortune’s (and is based on our textbook’s construction). Rather than distort the diagram, we can think of this algorithm as distorting the sweep line. Actually, we will think of two objects that control the sweeping process. First, there will be a horizontal sweep line, moving from top to bottom. We will also maintain an \(x\)-monotonic curve called a \textit{beach line}. (It is so named because it looks like waves rolling up on a beach.) The beach line is a monotone curve formed from pieces of parabolic arcs. As the sweep line moves downward, the beach line follows just behind. The job of the beach line is to prevent us from seeing unanticipated events until the sweep line encounters the corresponding site.

The Beach Line: In order to make these ideas more concrete, recall that the problem with ordinary plane sweep is that sites that lie below the sweep line may affect the diagram that lies above the sweep line. To avoid this problem, we will maintain only the portion of the diagram that cannot be affected by anything that lies below the sweep line. To do this, we will subdivide the halfplane lying above the sweep line into two regions: those points that are closer to some site \(p\) above the sweep line than they are to the sweep line itself, and those points that are closer to the sweep line than any site above the sweep line.

What are the geometric properties of the boundary between these two regions? The set of points \(q\) that are equidistant from the sweep line to their nearest site above the sweep line is called the \textit{beach line}. Observe that for any point \(q\) above the beach line, we know that its closest site cannot be affected by any site that lies below the sweep line. Hence, the portion of the Voronoi diagram that lies above the beach line is “safe” in the sense that we have all the information that we need in order to compute it (without knowing about which sites are still to appear below the sweep line).

What does the beach line look like? Recall from high school geometry that the set of points that are equidistant from a point (in this a site) and a line (in this case the sweep line) is a parabola (see the figure below, left). Clearly the parabola’s shape changes continuously as the sweep line moves continuously. With a little analytic geometry, it is easy to show that the parabola becomes “skinnier” when the site is closer to the line and becomes “fatter” as the sweep line moves farther away. In the degenerate case when the line contains the site the parabola degenerates into a vertical ray shooting up from the site. (You should work through the distance equations to see why this is so.)

Thus, the beach line consists of the \textit{lower envelope} of these parabolas, one for each site. Note that the parabola of some sites above the beach line will not touch the lower envelope and hence will not contribute to the beach line. Because the parabolas are \(x\)-monotone, so is the beach line. Also observe that the point where two arcs of the beach line intersect, which we call a \textit{breakpoint}, is equidistant
from two sites and the sweep line, and hence must lie on some Voronoi edge. In particular, if the
beach line arcs corresponding to sites \( p_i \) and \( p_j \) share a common breakpoint on the beach line, then
this breakpoint lies on the Voronoi edge between \( p_i \) and \( p_j \). From this we have the following important
characterization.

**Lemma:** The beach line is an \( x \)-monotone curve made up of parabolic arcs. The breakpoints (that is, 
vertices) of the beach line lie on Voronoi edges of the final diagram.

Fortune’s algorithm consists of simulating the growth of the beach line as the sweep line moves
downward, and in particular tracing the paths of the breakpoints as they travel along the edges of the
Voronoi diagram. Of course, as the sweep line moves the parabolas forming the beach line change
their shapes continuously. As with all plane-sweep algorithms, we will maintain a sweep-line status
and we are interested in simulating the discrete event points where there is a “significant event”, that
is, any event that changes the topological structure of the Voronoi diagram and the beach line.

**Sweep-Line Status:** The algorithm maintain the current location (\( y \)-coordinate) of the sweep line. It
stores, in left-to-right order the sequence of sites that define the beach line. (We will say more
about this later.) **Important:** The algorithm does not store the parabolic arcs of the beach line.
It exists solely for conceptual purposes.

**Events:** There are two types of events:

- **Site events:** When the sweep line passes over a new site a new arc will be inserted into the
  beach line.

- **Voronoi vertex events:** (What our text calls *circle events.*) When the length of a parabolic arc
  shrinks to zero, the arc disappears and a new Voronoi vertex will be created at this point.

The algorithm consists of processing these two types of events. As the Voronoi vertices are being dis-
covered by Voronoi vertex events, it will be an easy matter to update the diagram as we go (assuming
any reasonable representation of this planar cell complex), and so to link the entire diagram together.
Let us consider the two types of events that are encountered.

**Site events:** A site event is generated whenever the horizontal sweep line passes over a site \( p_i \). As we
mentioned before, at the instant that the sweep line touches the point, its associated parabolic arc will
degenerate to a vertical ray shooting up from the point to the current beach line. As the sweep line
proceeds downwards, this ray will widen into an arc along the beach line. To process a site event we
will determine the arc of the sweep line that lies directly above the new site. (Let us make the general position assumption that it does not fall immediately below a vertex of the beach line.) Let \( p_j \) denote the site generating this arc. We then split this arc of the beach line in two by inserting a new entry at this point in the sweep-line status. (Initially this corresponds to an infinitesimally small arc along the beach line, but as the sweep line sweeps on, this arc will grow wider. Thus, the entry for \( \langle \ldots, p_j, \ldots \rangle \) on the sweep-line status is replaced by the triple \( \langle \ldots, p_j, p_i, p_j, \ldots \rangle \). (See the figure below.)

![Site event](image_url)

**Figure 11.6: Site event.**

It is important to consider whether this is the only way that new arcs can be introduced into the sweep line. In fact it is. We will not prove it, but a careful proof is given in the text. As a consequence of this proof, it follows that the maximum number of arcs on the beach line can be at most \( 2n - 1 \), since each new point can result in creating one new arc, and splitting an existing arc, for a net increase of two arcs per point (except the first). Note that a point may generally contribute more than one arc to the beach line. (As an exercise you might consider what is the maximum number of arcs a single site can contribute.)

The nice thing about site events is that they are all known in advance. Thus, after sorting the points by \( y \)-coordinate, all these events are known.

**Voronoi vertex events:** In contrast to site events, Voronoi vertex events are generated dynamically as the algorithm runs. As with the line segment plane sweep algorithm, the important idea is that each such event is generated by objects that are neighbors on the beach line. However, unlike the segment intersection where pairs of consecutive segments generated events, here triples of points generate the events.

In particular, consider any three consecutive sites \( p_i, p_j, \) and \( p_k \) whose arcs appear consecutively on the beach line from left to right. (See the figure below.) Further, suppose that the circumcircle for these three sites lies at least partially below the current sweep line (meaning that the Voronoi vertex has not yet been generated), and that this circumcircle contains no points lying below the sweep line (meaning that no future point will block the creation of the vertex).

Consider the moment at which the sweep line falls to a point where it is tangent to the lowest point of this circle. At this instant the circumcenter of the circle is equidistant from all three sites and from the sweep line. Thus all three parabolic arcs pass through this center point, implying that the contribution of the arc from \( p_j \) has disappeared from the beach line. In terms of the Voronoi diagram, the bisectors \( \langle p_i, p_j \rangle \) and \( \langle p_j, p_k \rangle \) have met each other at the Voronoi vertex, and a single bisector \( \langle p_i, p_k \rangle \) remains. Thus, the triple of consecutive sites \( p_i, p_j, p_k \) on the sweep-line status is replaced with \( p_i, p_k \). (See the figure below.)

**Sweep-line algorithm:** We can now present the algorithm in greater detail. The main structures that we will maintain are the following:
(Partial) Voronoi diagram: The partial Voronoi diagram that has been constructed so far will be stored in any reasonable data structure for storing planar subdivisions, for example, a doubly-connected edge list. There is one technical difficulty caused by the fact that the diagram contains unbounded edges. To handle this we will assume that the entire diagram is to be stored within a large bounding box. (This box should be chosen large enough that all of the Voronoi vertices fit within the box.)

Beach line: The beach line consists of the sorted sequence of sites whose arcs form the beach line. It is represented using a dictionary (e.g., a balanced binary tree or skip list). As mentioned above, we do not explicitly store the parabolic arcs. They are just there for the purposes of deriving the algorithm. Instead for each parabolic arc on the current beach line, we store the site that gives rise to this arc.

The key search operation is that of locating the arc of the beach line that lies directly above a newly discovered site. (As an exercise, before reading the next paragraph you might think about how you would design a binary search to locate this arc, given that you only have the sites, not the actual arcs.)

Between each consecutive pair of sites \( p_i \) and \( p_j \), there is a breakpoint. Although the breakpoint moves as a function of the sweep line, observe that it is possible to compute the exact location of the breakpoint as a function of \( p_i \), \( p_j \), and the current \( y \)-coordinate of the sweep line. In particular, the breakpoint is the center of a circle that passes through \( p_i \), \( p_j \) and is tangent to the sweep line. (Thus, as with beach lines, we do not explicitly store breakpoints. Rather, we compute them only when we need them.) Once the breakpoint is computed, we can then determine whether a newly added site is to its left or right. Using the sorted ordering of the sites, we use this primitive comparison to drive a binary search for the arc lying above the new site.

The important operations that we will have to support on the beach line are:

Search: Given the current \( y \)-coordinate of the sweep line and a new site \( p_i \), determine the arc of the beach line lies immediately above \( p_i \). Let \( p_j \) denote the site that contributes this arc. Return a reference to this beach line entry.

Insert and split: Insert a new entry for \( p_i \) within a given arc \( p_j \) of the beach line (thus effectively replacing the single arc \( \langle \ldots, p_j, \ldots \rangle \) with the triple \( \langle \ldots, p_j, p_i, p_j, \ldots \rangle \). Return a reference to the newly added beach line entry (for future use).

Delete: Given a reference to an entry \( p_j \) on the beach line, delete this entry. This replaces a triple \( \langle \ldots, p_i, p_j, p_k, \ldots \rangle \) with the pair \( \langle \ldots, p_i, p_k, \ldots \rangle \).

It is not difficult to modify a standard dictionary data structure to perform these operations in \( O(\log n) \) time each.
Event queue: The event queue is a priority queue with the ability both to insert and delete new events. Also the event with the largest $y$-coordinate can be extracted. For each site we store its $y$-coordinate in the queue. All operations can be implemented in $O(\log n)$ time assuming that the priority queue is stored as an ordered dictionary.

For each consecutive triple $p_i, p_j, p_k$ on the beach line, we compute the circumcircle of these points. (We’ll leave the messy algebraic details as an exercise, but this can be done in $O(1)$ time.) If the lower endpoint of the circle (the minimum $y$-coordinate on the circle) lies below the sweep line, then we create a Voronoi vertex event whose $y$-coordinate is the $y$-coordinate of the bottom endpoint of the circumcircle. We store this in the priority queue. Each such event in the priority queue has a cross link back to the triple of sites that generated it, and each consecutive triple of sites has a cross link to the event that it generated in the priority queue.

The algorithm proceeds like any plane sweep algorithm. The algorithm starts by inserting the topmost vertex into the sweep-line status. We extract an event, process it, and go on to the next event. Each event may result in a modification of the Voronoi diagram and the beach line, and may result in the creation or deletion of existing events.

Here is how the two types of events are handled in somewhat greater detail.

Site event: Let $p_i$ be the new site. (See Fig. 11.6 above.)

(1) Advance the sweep line so that it passes through $p_i$. Apply the above search operation to determine the beach line arc that lies immediately above $p_i$. Let $p_j$ be the corresponding site.

(2) Applying the above insert-and-split operation, inserting a new entry for $p_i$, thus replacing \(\langle\ldots, p_j, \ldots\rangle\) with \(\langle\ldots, p_j, p_i, p_j, \ldots\rangle\).

(3) Create a new (dangling) edge in the Voronoi diagram, which lies on the bisector between $p_i$ and $p_j$.

(4) Some old triples that involved $p_j$ may need to be deleted and some new triples involving $p_i$ will be inserted, based on the change of neighbors on the beach line. (The straightforward details are omitted.)

Note that the newly created beach-line triple $p_j, p_i, p_j$ cannot generate an event because it only involves two distinct sites.

Voronoi vertex event: Let $p_i, p_j, p_k$ be the three sites that generated this event, from left to right. (See Fig. 11.7 above.)

(1) Using the above delete operation, delete the entry for $p_j$ from the beach line. (Thus eliminating its arc.)

(2) Create a new vertex in the Voronoi diagram (at the circumcenter of \(\{p_i, p_j, p_k\}\)) and join the two Voronoi edges for the bisectors \(\langle p_i, p_j \rangle\), \(\langle p_j, p_k \rangle\) to this vertex. (These edges were generated during an earlier site event in Step (3) above.)

(3) Create a new (dangling) edge for the bisector between $p_i$ and $p_k$.

(4) Delete any events that arose from triples involving the arc of $p_j$, and generate new events corresponding to consecutive triples involving $p_i$ and $p_k$. (There are two of them. The straightforward details are omitted.)
The analysis follows a typical analysis for plane sweep. Each event involves $O(1)$ processing time plus a constant number operations to the various data structures. The size of the data structures is $O(n)$, and each of these operations takes $O(\log n)$ time. Thus the total time is $O(n \log n)$, and the total space is $O(n)$. 
12 Delaunay Triangulations

Reading: Chapter 9 in the 4M’s.

Delaunay Triangulations: Recently we discussed the topic of Voronoi diagrams. Today we consider the related structure, called a Delaunay triangulation (DT). The Voronoi diagram of a set of sites in the plane is a planar subdivision, that is, cell complex. The dual of such subdivision is another subdivision that is defined as follows. For each face of the Voronoi diagram, we create a vertex (corresponding to the site). For each edge of the Voronoi diagram lying between two sites \( p_i \) and \( p_j \), we create an edge in the dual connecting these two vertices. Finally, each vertex of the Voronoi diagram corresponds to a face of the dual.

The resulting dual graph is a planar subdivision. Assuming general position, the vertices of the Voronoi diagram have degree three, it follows that the faces of the resulting dual graph (excluding the exterior face) are triangles. Thus, the resulting dual graph is a triangulation of the sites, called the Delaunay triangulation. This is illustrated in the figure below.

![Figure 12.1: The Delaunay triangulation of a set of points (solid lines) and the Voronoi diagram (broken lines).](image)

Delaunay triangulations have a number of interesting properties, that are consequences of the structure of the Voronoi diagram.

**Convex hull:** The boundary of the exterior face of the Delaunay triangulation is the boundary of the convex hull of the point set.

**Circumcircle property:** The circumcircle of any triangle in the Delaunay triangulation is empty (contains no sites of \( P \)).

Proof: This is because the center of this circle is the corresponding dual Voronoi vertex, and by definition of the Voronoi diagram, the three sites defining this vertex are its nearest neighbors.

**Empty circle property:** Two sites \( p_i \) and \( p_j \) are connected by an edge in the Delaunay triangulation, if and only if there is an empty circle passing through \( p_i \) and \( p_j \).

Proof: If two sites \( p_i \) and \( p_j \) are neighbors in the Delaunay triangulation, then their cells are neighbors in the Voronoi diagram, and so for any point on the Voronoi edge between these sites, a circle centered at this point passing through \( p_i \) and \( p_j \) cannot contain any other point (since they must be closest). Conversely, if there is an empty circle passing through \( p_i \) and \( p_j \), then the center \( c \) of this circle is a point on the edge of the Voronoi diagram between \( p_i \) and \( p_j \), because \( c \) is equidistant from each of these sites and there is no closer site. Thus the Voronoi cells of two sites are adjacent in the Voronoi diagram, implying that there edge is in the Delaunay triangulation.
**Closest pair property:** The closest pair of sites in $P$ are neighbors in the Delaunay triangulation.

Proof: Suppose that $p_i$ and $p_j$ are the closest sites. The circle having $p_i$ and $p_j$ as its diameter cannot contain any other site, since otherwise such a site would be closer to one of these two points, violating the hypothesis that these points are the closest pair. Therefore, the center of this circle is on the Voronoi edge between these points, and so it is an empty circle.

If the sites are not in general position, in the sense that four or more are cocircular, then the Delaunay triangulation may not be a triangulation at all, but just a planar graph (since the Voronoi vertex that is incident to four or more Voronoi cells will induce a face whose degree is equal to the number of such cells). In this case the more appropriate term would be Delaunay graph. However, it is common to either assume the sites are in general position (or to enforce it through some sort of symbolic perturbation) or else to simply triangulate the faces of degree four or more in any arbitrary way. Henceforth we will assume that sites are in general position, so we do not have to deal with these messy situations.

Given a point set $P$ with $n$ sites where there are $h$ sites on the convex hull, it is not hard to prove by Euler’s formula that the Delaunay triangulation has $2n - 2 - h$ triangles, and $3n - 3 - h$ edges. The ability to determine the number of triangles from $n$ and $h$ only works in the plane. In 3-space, the number of tetrahedra in the Delaunay triangulation can range from $O(n)$ up to $O(n^2)$. In dimension $d$, the number of simplices (the $d$-dimensional generalization of a triangle) can range as high as $O(n^{\lceil d/2 \rceil})$.

**Minimum Spanning Tree:** The Delaunay triangulation possesses some interesting properties that are not directly related to the Voronoi diagram structure. One of these is its relation to the minimum spanning tree. Given a set of $n$ points in the plane, we can think of the points as defining a Euclidean graph whose edges are all $(n \choose 2)$ (undirected) pairs of distinct points, and edge $(p_i, p_j)$ has weight equal to the Euclidean distance from $p_i$ to $p_j$. A minimum spanning tree is a set of $n - 1$ edges that connect the points (into a free tree) such that the total weight of edges is minimized. We could compute the MST using Kruskal’s algorithm. Recall that Kruskal’s algorithm works by first sorting the edges and inserting them one by one. We could first compute the Euclidean graph, and then pass the result on to Kruskal’s algorithm, for a total running time of $O(n^2 \log n)$.

However there is a much faster method based on Delaunay triangulations. First compute the Delaunay triangulation of the point set. We will see later that it can be done in $O(n \log n)$ time. Then compute the MST of the Delaunay triangulation by Kruskal’s algorithm and return the result. This leads to a total running time of $O(n \log n)$. The reason that this works is given in the following theorem.

**Theorem:** The minimum spanning tree of a set of points $P$ (in any dimension) is a subgraph of the Delaunay triangulation.

**Proof:** Let $T$ be the MST for $P$, let $w(T)$ denote the total weight of $T$. Let $a$ and $b$ be any two sites such that $ab$ is an edge of $T$. Suppose to the contrary that $ab$ is not an edge in the Delaunay triangulation. This implies that there is no empty circle passing through $a$ and $b$, and in particular, the circle whose diameter is the segment $ab$ contains a site, call it $c$. (See the figure below.)

The removal of $ab$ from the MST splits the tree into two subtrees. Assume without loss of generality that $c$ lies in the same subtree as $a$. Now, remove the edge $ab$ from the MST and add the edge $bc$ in its place. The result will be a spanning tree $T'$ whose weight is

$$w(T') = w(T) + |bc| - |ab| < w(T).$$
The last inequality follows because $ab$ is the diameter of the circle, implying that $|bc| < |ab|$. This contradicts the hypothesis that $T$ is the MST, completing the proof.

By the way, this suggests another interesting question. Among all triangulations, we might ask, does the Delaunay triangulation minimize the total edge length? The answer is no (and there is a simple four-point counterexample). However, this claim was made in a famous paper on Delaunay triangulations, and you may still hear it quoted from time to time. The triangulation that minimizes total edge weight is called the \textit{minimum weight triangulation}. Recently it was proved that this problem is NP-hard. (This problem has been open for many years, dating back to the original development of the theory of NP-completeness back in the 1970’s.)

\textbf{Spanner Properties:} A natural observation about Delaunay triangulations is that its edges would seem to form a reasonable transportation road network between the points. On inspecting a few examples, it is natural to conjecture that the length of the shortest path between two points in a planar Delaunay triangulation is not significantly longer than the straight-line distance between these points.

This is closely related to the theory of geometric spanners, that is, geometric graphs whose shortest paths are not too long. Consider any point set $P$ and a straight-line graph $G$ whose vertices are the points of $P$. For any two points $p, q \in P$, let $\delta_G(p, q)$ denote the length of the shortest path from $p$ to $q$ in $G$, where the weight of each edge is its Euclidean length. Given any parameter $t \geq 1$, we say that $G$ is a $t$-\textit{spanner} if for any two points $p, q \in P$, the shortest path length between $p$ and $q$ in $G$ is at most a factor $t$ longer than the Euclidean distance between these points, that is

$$\delta_G(p, q) \leq t||pq||$$

Observe that when $t = 1$, the graph $G$ must be the complete graph, consisting of $\binom{n}{2} = O(n^2)$ edges. Of interest is whether there exist spanners having $O(n)$ edges.

It can be proved that the edges of the Delaunay triangulation form a spanner. We will not prove the following result, which is due to Keil and Gutwin.

\textbf{Theorem:} Given a set of points $P$ in the plane, the Delaunay triangulation of $P$ is a $t$-spanner for $t = 4\pi\sqrt{3}/9 \approx 2.4$.

In fact, it is conjectured that the Delaunay triangulation is a $(\pi/2)$-spanner, but this has never been proved (and it seems to be a hard problem).

\textbf{Maximizing Angles and Edge Flipping:} Another interesting property of Delaunay triangulations is that among all triangulations, the Delaunay triangulation maximizes the minimum angle. This property
is important, because it implies that Delaunay triangulations tend to avoid skinny triangles. This is useful for many applications where triangles are used for the purposes of interpolation.

In fact a much stronger statement holds as well. Among all triangulations with the same smallest angle, the Delaunay triangulation maximizes the second smallest angle, and so on. In particular, any triangulation can be associated with a sorted angle sequence, that is, the increasing sequence of angles \((\alpha_1, \alpha_2, \ldots, \alpha_m)\) appearing in the triangles of the triangulation. (Note that the length of the sequence will be the same for all triangulations of the same point set, since the number depends only on \(n\) and \(h\).)

**Theorem:** Among all triangulations of a given planar point set, the Delaunay triangulation has the lexicographically largest angle sequence, and in particular, it maximizes the minimum angle.

Before getting into the proof, we should recall a few basic facts about angles from basic geometry. First, recall that if we consider the circumcircle of three points, then each angle of the resulting triangle is exactly half the angle of the minor arc subtended by the opposite two points along the circumcircle. It follows as well that if a point is inside this circle then it will subtend a larger angle and a point that is outside will subtend a smaller angle. This in the figure part (a) below, we have \(\theta_1 > \theta_2 > \theta_3\).

![Figure 12.3: Angles and edge flips.](image)

We will not give a formal proof of the theorem. (One appears in the text.) The main idea is to show that for any triangulation that fails to satisfy the empty circle property, it is possible to perform a local operation, called an edge flip, which increases the lexicographical sequence of angles. An edge flip is an important fundamental operation on triangulations in the plane. Given two adjacent triangles \(\Delta abc\) and \(\Delta cda\), such that their union forms a convex quadrilateral \(abcd\), the edge flip operation replaces the diagonal \(ac\) with \(bd\). Note that it is only possible when the quadrilateral is convex. Suppose that the initial triangle pair violates the empty circle condition, in that point \(d\) lies inside the circumcircle of \(\Delta abc\). (Note that this implies that \(b\) lies inside the circumcircle of \(\Delta cda\).) If we flip the edge it will follow that the two circumcircles of the two resulting triangles, \(\Delta abd\) and \(\Delta bcd\) are now empty (relative to these four points), and the observation above about circles and angles proves that the minimum angle increases at the same time. In particular, in the figure above, we have

\[\phi_{ab} > \theta_{ab}, \phi_{bc} > \theta_{bc}, \phi_{cd} > \theta_{cd}, \phi_{da} > \theta_{da}.\]

There are two other angles that need to be compared as well (can you spot them?). It is not hard to show that, after swapping, these other two angles cannot be smaller than the minimum of \(\theta_{ab}, \theta_{bc}, \theta_{cd}\), and \(\theta_{da}\). (Can you see why?)

Since there are only a finite number of triangulations, this process must eventually terminate with the lexicographically maximum triangulation, and this triangulation must satisfy the empty circle condition, and hence is the Delaunay triangulation.
Note that the process of edge-flipping can be generalized to simplicial complexes in higher dimensions. However, the process does not generally replace a fixed number of triangles with the same number, as it does in the plane (replacing two old triangles with two new triangles). For example, in 3-space, the most basic flip can replace two adjacent tetrahedra with three tetrahedra, and vice versa. Although it is known that in the plane any triangulation can be converted into any other through a judicious sequence of edge flips, this is not known in higher dimensions.
13 Delaunay Triangulations: Incremental Construction

Reading: Chapter 9 in the 4M’s.

Constructing the Delaunay Triangulation: We will present a simple randomized $O(n \log n)$ expected time algorithm for constructing Delaunay triangulations for $n$ sites in the plane. The algorithm is remarkably similar in spirit to the randomized algorithm for trapezoidal map algorithm in that not only builds the triangulation but also provides a point-location data structure as well. We will not discuss the point-location data structure in detail, but the details are easy to fill in.

As with any randomized incremental algorithm, the idea is to insert sites in random order, one at a time, and update the triangulation with each new addition. The issues involved with the analysis will be showing that after each insertion the expected number of structural changes in the diagram is $O(1)$.

As with other incremental algorithm, we need some way of keeping track of where newly inserted sites are to be placed in the diagram. We will describe a somewhat simpler method than the one we used in the trapezoidal map. Rather than building a data structure, this one simply puts each of the uninserted points into a bucket according to the triangle that contains it in the current triangulation. In this case, we will need to argue that the expected number of times that a site is re-bucketed is $O(\log n)$.

Incircle Test: The basic issue in the design of the algorithm is how to update the triangulation when a new site is added. In order to do this, we first investigate the basic properties of a Delaunay triangulation.

Recall that a triangle $\triangle abc$ is in the Delaunay triangulation, if and only if the circumcircle of this triangle contains no other site in its interior. (Recall that we make the general position assumption that no four sites are cocircular.) How do we test whether a site $d$ lies within the interior of the circumcircle of $\triangle abc$? It turns out that this can be reduced to a determinant computation. First off, let us assume that the sequence $\langle abcd \rangle$ defines a counterclockwise convex polygon. [WE VIOLATE THIS ORDERING ASSUMPTION LATER IN THE ALGORITHM—FIX] (If it does not because $d$ lies inside the triangle $\triangle abc$ then clearly $d$ lies in the circumcircle for this triangle. Otherwise, we can always relabel $abc$ so this is true.) Under this assumption, $d$ lies in the circumcircle determined by the $\triangle abc$ if and only if the following determinant is positive. This is called the incircle test. We will assume that this primitive is available to us.

$$\text{inCircle}(a, b, c, d) = \det \begin{pmatrix} a_x & a_y & a_x^2 + a_y^2 & 1 \\ b_x & b_y & b_x^2 + b_y^2 & 1 \\ c_x & c_y & c_x^2 + c_y^2 & 1 \\ d_x & d_y & d_x^2 + d_y^2 & 1 \end{pmatrix} > 0.$$  

We will not prove the correctness of this test, but we will show a somewhat simpler assertion, namely that if the above determinant is equal to zero, then the four points are cocircular. The four points are cocircular if there exists a center point $q = (q_x, q_y)$ and a radius $r$ such that

$$(a_x - q_x)^2 + (a_y - q_y)^2 = r^2,$$

and similarly for the other three points. Expanding this and collecting common terms we have

$$(a_x^2 + a_y^2) - 2q_x a_x - 2q_y a_y + (q_x^2 + q_y^2 - r^2) = 0,$$
and similarly for the other three points, \(b, c, \) and \(d\). If we let \(X_1, X_2, X_3\) and \(X_4\) denote the columns of the above matrix we have

\[
X_3 - 2q_x X_1 - 2q_y X_2 + (q_x^2 + q_y^2 - r^2)X_4 = 0.
\]

Since there is a linear combination of these vectors that sums to 0, it follows that these vector are linearly dependent. From standard linear algebra, we know that the columns of a matrix are linearly dependent if and only if the determinant of the matrix is zero. We will leave the completion of the proof (involving inside and outside) as an exercise.

**Incremental update:** When we add the next site, \(p_i\), the problem is to convert the current Delaunay triangulation into a new Delaunay triangulation containing this site. This will be done by creating a non-Delaunay triangulation containing the new site, and then incrementally “fixing” this triangulation to restore the Delaunay properties. The fundamental changes will be: (1) adding a site to the middle of a triangle, and creating three new edges, and (2) performing an edge flip. Both of these operations can be performed in \(O(1)\) time, assuming that the triangulation is maintained in any reasonable way, say, as a double-connected edge list.

The algorithm that we will describe has been known for many years, but was first analyzed by Guibas, Knuth, and Sharir. The algorithm starts within an initial triangulation such that all the points lie in the convex hull. This can be done by enclosing the points in a suitably large triangle. Our book suggests an alternative, namely computing a triangle that contains all the points, but then “fudging” the incircle test so that the vertices of this enclosing triangle “behave” as if they were infinitely far away.

The sites are added in random order. When a new site \(p\) is added, we find the triangle \(\triangle abc\) of the current triangulation that contains this site (we will see how later), insert the site in this triangle, and join this site to the three surrounding vertices. This creates three new triangles, \(\triangle pab, \triangle pbc,\) and \(\triangle pca\), each of which may or may not satisfy the empty-circle condition. How do we test this? For each of the triangles that have been added, we check the vertex of the triangle that lies on the opposite side of the edge that does not include \(p\). (If there is no such vertex, because this edge is on the convex hull, then we are done.) If this vertex fails the incircle test (that is, if it is inside the circumcircle), then we swap the edge (creating two new triangles that are adjacent to \(p\)). This replaces one triangle that was incident to \(p\) with two new triangles. We repeat the same test with these triangles. An example is shown in the figure below.

The algorithm for the incremental algorithm is shown below, and an example is presented in Fig. 13.2. The current triangulation is kept in a global data structure. The edges in the following algorithm should be thought of as pointers to an reasonable representation of the simplicial complex.

As you can see, the algorithm is very simple. There are only two elements that have not been shown are the implementation. The first is the update operations on the data structure for the simplicial

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2Some care must be taken in the construction of this enclosing triangle. It is not sufficient that it simply contains all the points. It should be so large that the vertices of the triangle do not lie in the circumcircles of any of the triangles of the final triangulation.
Randomized Incremental Delaunay Triangulation Algorithm

\[\text{Insert}(p) \{\]
\[\quad \text{Find the triangle } \triangle abc \text{ containing } p;\]
\[\quad \text{Insert edges } pa, pb, \text{ and } pc \text{ into triangulation;}\]
\[\quad \text{SwapTest}(ab); \quad \text{// Fix the surrounding edges}\]
\[\quad \text{SwapTest}(bc);\]
\[\quad \text{SwapTest}(ca);\]
\[\}\]

\[\text{SwapTest}(ab) \{\]
\[\quad \text{if } (ab \text{ is an edge on the exterior face}) \text{ return;}\]
\[\quad \text{Let } d \text{ be the vertex to the right of edge } ab;\]
\[\quad \text{if } (\text{inCircle}(p, a, b, d)) \{ \quad \text{// } d \text{ violates the incircle test}\]
\[\quad \quad \text{Flip edge } ab \text{ for } pd;\]
\[\quad \quad \text{SwapTest}(ad); \quad \text{// Fix the new suspect edges}\]
\[\quad \quad \text{SwapTest}(db);\]
\[\quad \}\]
complex. These can be done in $O(1)$ time each on any reasonable representation. The other issue is locating the triangle that contains $p$. We will discuss this below.

**Correctness:** There is one major issue in establishing the correctness of the algorithm. When we performed empty-circle tests, we only tested the empty circle tests for the newly created triangles containing the site $p$, and then only for sites that lay on the opposite side of an edge of each such triangle.

This is related to an important issue in Delaunay triangulations. We say that a triangulation is *locally Delaunay* if for each triangle the vertices lying on the opposite side of each edge of the (up to) three neighboring triangles satisfy the empty-circle condition. But to establish complete correctness of the triangulation, we need to show that the triangulation is *globally Delaunay*, meaning that empty-circle condition is satisfied for all triangles, and all points of $P$. That is, it suffices to show that if a triangulation is locally Delaunay, then it is globally Delaunay. This important theorem (called *Delaunay’s Theorem*), and we will sketch a proof of this theorem below for this special context.

First, to see that it suffices to consider only triangles that are incident to $p$, observe that $p$ is the only newly added site, and hence it is the only site that can cause a violation of the empty-circle condition.

To finish the argument, it suffices to see why “locally Delaunay” implies “globally Delaunay.” Consider a triangle $\triangle pab$ that contains $p$ and consider the vertex $d$ belonging to the triangle that lies on the opposite side of edge $ab$. We argue that if $d$ lies outside the circumcircle of $pab$, then no other point of the point set can lie within this circumcircle. A complete proof of this takes some effort, but here is a simple justification. What could go wrong? It might be that $d$ lies outside the circumcircle, but there is some other site, say, a vertex $e$ of a triangle adjacent to $d$, that lies inside the circumcircle. (This is illustrated in the figure below). We claim that this cannot happen. It can be shown that if $e$ lies within the circumcircle of $\triangle pab$, then $a$ must lie within the circumcircle of $\triangle bde$. (The argument is an exercise in the geometry of circles.) However, this would violate the assumption that the initial triangulation (before the insertion of $p$) was a Delaunay triangulation.

![Figure 13.3: Proof of sufficiency of testing neighboring sites.](image)

**Point Location:** The point location can be accomplished by one of two means. Our text discusses the idea of building a history graph point-location data structure, just as we did in the trapezoid map case. A simpler approach is based on the idea of maintaining the uninserted sites in a set of *buckets*. Think of each triangle of the current triangulation as a *bucket* that holds the sites that lie within this triangle and have yet to be inserted. Whenever an edge is flipped, or when a triangle is split into three triangles through point insertion, some old triangles are destroyed and are replaced by a constant number of new triangles. When this happens, we lump together all the sites in the buckets corresponding to the deleted triangles, create new buckets for the newly created triangles, and reassign each site into its new bucket. Since there are a constant number of triangles created, this process requires $O(1)$ time per site that is rebucketed.

13-4

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To analyze the expected running time of algorithm we need to bound two quantities: (1) how many structural changes are made in the triangulation on average with the addition of each new site, and (2) how much effort is spent in rebucketing sites. As usual, our analysis will be in the worst-case (for any point set) but averaged over all possible insertion orders.

**Structural Changes:** We argue first that the expected number of edge changes with each insertion is $O(1)$ by a simple application of backwards analysis. First observe that (assuming general position) the structure of the Delaunay triangulation is independent of the insertion order of the sites so far. Thus, any of the existing sites is equally likely to have been the last site to be added to the structure. Suppose that some site $p$ was the last to have been added. How much work was needed to insert $p$? Observe that the initial insertion of $p$ involved the creation of three new edges, all incident to $p$. Also, whenever an edge swap is performed, a new edge is added to $p$. These are the only changes that the insertion algorithm can make. Therefore the total number of changes made in the triangulation for the insertion of $p$ is proportional to the degree of $p$ after the insertion is complete. Thus the work needed to insert $p$ is proportional to $p$’s degree after the insertion.

To perform the backwards analysis, we consider the situation after the insertion of the $i$th site. Since the diagram’s structure does not depend on the order of insertion, every one of the $i$ sites appearing in the diagram was equally likely to be the last one added. Thus, by a backwards analysis, the expected time to insert the $i$th site is equal to the average degree of a vertex in the triangulation of $i$ sites. (The only exception are the three initial vertices at infinity, which must be the first sites to be inserted.) By Euler’s formula we know that the average degree of a vertex in any planar graph is at most 6. (Recall that a planar graph with $n$ vertices can have at most $3n$ edges, and the sum of vertex degrees is equal to twice the number of edges, which is at most $6n$.) Thus, irrespective of the stage number, the expected number of edge changes is proportional to the expected vertex degree, which is $O(1)$.

Summing over all $n$ insertions, the total number of structural changes is $O(n)$. Recall that each structural change (new edges and edge flips) can be performed in $O(1)$ time.

**Rebucketing:** Next we argue that the total expected time spent in rebucketing points is $O(n \log n)$. From this it will follow that the overall expected running time is dominated by the rebucketing process, and so is $O(n \log n)$.

To do this, we will show that the expected number of times that any site is rebucketed (as to which triangle it lies in) is $O(\log n)$. Again this is done by a standard application of backwards analysis. Let us fix a site $q \in P$. Consider the situation just after the insertion of the $i$th site. If $q$ has already been inserted, then it is not involved in the rebucketing process, so let us assume that $q$ has not yet been inserted. As above we make use of the fact that any of the existing sites is equally likely to be the last site inserted.

We assert that the probability that $q$ was rebucketed as a result of the last insertion is at most $3/i$. To see this, let $\Delta$ be the triangle containing $q$ after the $i$th insertion. As observed above, after we insert the $i$th site all of the newly created triangles are now incident to this new site. $\Delta$ would have come into existence as a result of the last insertion if and only one of its three vertices was the last to be added. (Think about this a moment, since it is critical!) Since $\Delta$ is incident to exactly three sites, and every site is equally likely to be the last inserted, it follows that the probability that $\Delta$ came into existence is $3/i$. (We are cheating a bit here by ignoring the three initial sites at infinity.) Thus, the probability that $q$ required rebucketing after the last insertion is at most $3/i$. 

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After stage $i$ there are $n - i$ points that might be subject to rebucketing, and each has probability $3/i$ of being rebucketed. Thus, the expected number of points that require rebucketing as part of the last insertion is $(n - i)3/i$. By the linearity of expectation, to obtain the total number of rebucketings, we sum these up over all stages, yielding

$$
\sum_{i=1}^{n} \frac{3}{i}(n - i) \leq \sum_{i=1}^{n} \frac{3}{i}n = 3n \sum_{i=1}^{n} \frac{1}{i} = 3n \ln n + O(1),
$$

(where as usual we have applied the bound on the Harmonic series.) Thus, the total expected time spent in rebucketing is $O(n \log n)$, as desired.

There is one place in the proof that we were sloppy. (Can you spot it?) We showed that the number of points that required rebucketing is $O(n \log n)$, but notice that when a point is inserted, many rebucketing operations may be needed (one for the initial insertion and one for each additional edge flip). We will not give a careful analysis of the total number of individual rebucketing operations per point, but it is not hard to show that the expected total number of individual rebucketing operations will not be larger by more than a constant factor. The reason is that (as argued above) each new insertion only results in a constant number of edge flips, and hence the number of individual rebucketings per insertion is also a constant. But a careful proof should consider this. Such a proof is given in our textbook.


14 Line Arrangements and the Zone Theorem

Reading: Chapter 8 in the 4M’s.

Arrangements: So far we have studied a few of the most important structures in computational geometry: convex hulls, Voronoi diagrams and Delaunay triangulations. Perhaps, the next most important structure is that of a line arrangement. As with hulls and Voronoi diagrams, it is possible to define arrangements (of \((d-1)\)-dimensional hyperplanes) in any dimension, but we will concentrate on the plane. As with Voronoi diagrams, a line arrangement is a polygonal subdivision of the plane. Unlike most of the structures we have seen up to now, a line arrangement is not defined in terms of a set of points, but rather in terms of a set \(L\) of lines. However, line arrangements are used mostly for solving problems on point sets. The connection is that the arrangements are typically constructed in the dual plane. We will begin by defining arrangements, discussing their combinatorial properties and how to construct them, and finally discuss applications of arrangements to other problems in computational geometry.

Before discussing algorithms for computing arrangements and applications, we first provide definitions and some important theorems that will be used in the construction. A finite set \(L\) of lines in the plane subdivides the plane into a cell complex. The result is called an arrangement, denoted \(A(L)\). (Arrangements can also be defined for curves and surfaces, but the theory is simplest and for lines and planes.) In the plane, the arrangement defines a planar graph whose vertices are the points where two or more lines intersect, edges are the intersection free segments (or rays) of the lines, and faces are (possibly unbounded) convex regions containing no line. An example is shown in the figure below on the left.

![Figure 14.1: Arrangement of lines.](image)

An arrangement is said to be simple if no three lines intersect at a common point. We will make the usual general position assumptions that no three lines intersect in a single point. This assumption can be overcome either through a more careful implementation of special cases or through an appropriate symbolic perturbation of the line coefficients.

An arrangement is not formally a planar graph, because it has unbounded edges. We can fix this (topologically) by imagining that a vertex is added at infinity, and all the unbounded edges are attached to this vertex. A somewhat more geometric way to fix this is to imagine that there is a bounding box which is large enough to contain all the vertices, and we tie all the unbounded edges off at this box. It is obvious that we can compute such a box in \(O(n^2)\) time, but an \(O(n)\) time algorithm exist. (This is a nice exercise.) However, rather than computing the coordinates of this huge box, it is simpler to imagine the existence of an abstract “box at infinity.” Intersection computations involving this box are performed as if its bounding sides are at \(x, y \in \{-\infty, +\infty\}\). If you think about it a moment, you
can convince yourself that this is equivalent to processing the lines in slope order. (If you don’t see this right away, think about it.)

The **combinatorial complexity** of an arrangement is the total number of vertices, edges, and faces in the arrangement. The following shows that all of these quantities are $O(n^2)$.

**Theorem:** Give a set of $n$ lines $L$ in the plane in general position,

(i) the number of vertices in $\mathcal{A}(L)$ is $\binom{n}{2}$,

(ii) the number of edges in $\mathcal{A}(L)$ is $n^2$, and

(iii) the number of faces in $\mathcal{A}(L)$ is $\binom{n}{2} + n + 1$.

**Proof:** The fact that the number of vertices is $\binom{n}{2}$ is clear from the fact that each pair of lines intersects in a single point. To prove that the number of edges is $n^2$, we use induction. The basis case is trivial (1 line and 1 edge). When we add a new line to an arrangement of $n-1$ lines (having $(n-1)^2$ edges by the induction hypothesis) we split $n-1$ existing edges, thus creating $n-1$ new edges, and we add $n$ new edges from the $n-1$ intersections with the new line. This gives a total of $(n-1)^2 + (n-1) + n = n^2$. The number of faces comes from Euler’s formula, $v - e + f = 2$ (with the little trick that we need to create one extra vertex to attach all the unbounded edges to).

**Incremental Construction:** Arrangements are used for solving many problems in computational geometry. But in order to use arrangements, we first must be able to construct arrangements. We will present a simple incremental algorithm, which builds an arrangement by adding lines one at a time. Unlike most of the other incremental algorithms we have seen so far, this one is not randomized. Its asymptotic running time will be the same, $O(n^2)$, no matter what order we insert the lines. This is asymptotically optimal, since this is the size of the arrangement. The algorithm will also require $O(n^2)$ space, since this is the amount of storage needed to store the final result. (Later we will consider the question of whether it is possible to traverse an arrangement without actually building it.)

As usual, we will assume that the arrangement is represented in any reasonable data structure for planar graphs (e.g., as a doubly-connected edge list). Let $L = \{\ell_1, \ell_2, \ldots, \ell_n\}$ denote the lines. We will simply add lines one by one to the arrangement. (The order does not matter.) We will show that the $i$-th line can be inserted in $O(i)$ time. If we sum over $i$, this will give $O(n^2)$ total time.

Suppose that the first $i-1$ lines have already been added, and consider the effort involved in adding $\ell_i$. Recall our assumption that the arrangement is assumed to lie within a large bounding box. Since each line intersects this box twice, the first $i-1$ lines have subdivided the boundary of the box into $2(i-1)$ edges. We determine where $\ell_i$ intersects the box, and which of these edges it crosses intersects. This will tell us which face of the arrangement $\ell_i$ first enters.

Next we trace the line through the arrangement, from one face to the next. Whenever we enter a face, the main question is where does the line exit the face? We answer the question by a very simple strategy. We walk along the edges of the face, say in a counterclockwise direction. Recall that any reasonable subdivision representation allows such local traversal in time linear in the number of elements visited. As we visit each edge we ask whether $\ell_i$ intersects this edge. When we find the edge through which $\ell_i$ exits the face, we jump to the face on the other side of this edge and continue the trace with the neighboring face. This is illustrated in the figure below.

Once we know the points of entry and exit into each face, the last matter is to update the subdivision representation by inserting edges for each entry-exit pair. This is easy to handle in constant time, on any of the standard representations.
The zone of a line $\ell$ traversing the lower part of the zone.

Figure 14.2: Traversing an arrangement and zones.

Clearly the time that it takes to perform the insertion is proportional to the total number of edges that have been traversed in this tracing process. A naive argument says that we encounter $i - 1$ lines, and hence pass through $i$ faces (assuming general position). Since each face is bounded by at most $i$ lines, each facial traversal will take $O(i)$ time, and this gives a total $O(i^2)$. Hey, what went wrong? Above we said that we would do this in $O(i)$ time. The claim is that the traversal does indeed traverse only $O(i)$ edges, but to understand why, we need to delve more deeply into a concept of a zone of an arrangement.

**Zone Theorem:** The most important combinatorial property of arrangements (which is critical to their efficient construction) is a rather surprising result called the zone theorem. Given an arrangement $A$ of a set $L$ of $n$ lines, and given a line $\ell$ that is not in $L$, the zone of $\ell$ in $A(\ell)$, denoted $Z_A(\ell)$, is the set of faces whose closure intersects $\ell$. The figure above illustrates a zone for the line $\ell$. For the purposes of the above construction, we are only interested in the edges of the zone that lie below $\ell_i$, but if we bound the total complexity of the zone, then this will be an upper bound on the number of edges traversed in the above algorithm. The combinatorial complexity of a zone (as argued above) is at most $O(n^2)$. The Zone theorem states that the complexity is actually much smaller, only $O(n)$.

**Theorem: (Zone Theorem)** Given an arrangement $A(L)$ of $n$ lines in the plane, and given any line $\ell$ in the plane, the total number of edges in all the cells of the zone $Z_A(\ell)$ is at most $6n$.

**Proof:** As with most combinatorial proofs, the key is to organize everything so that the counting can be done in an easy way. Note that this is not trivial, because it is easy to see that any one line of $L$ might contribute many segments to the zone of $\ell$. The key in the proof is finding a way to add up the edges so that each line appears to induce only a constant number of edges into the zone. The proof is based on a simple inductive argument. We will first imagine, through a suitable rotation, that $\ell$ is horizontal, and further that none of the lines of $L$ is horizontal (through an infinitesimal rotation). We split the edges of the zone into two groups, those that bound some face from the left side and those that bound some face from the right side. More formally, since each face is convex, if we split it at its topmost and bottommost vertices, we get two convex chains of edges. The left-bounding edges are on the left chain and the right-bounding edges are on the right chain. We will show that there are at most $3n$ lines that bound faces from the left, and a symmetrical argument applies to the right-bounding edges. (Note that an edge of the zone that crosses $\ell$ itself contributes only once to the complexity of the zone. In the book’s proof they seem to count this edge twice, and hence their bound they get a bound of $4n$ instead. We will also ignore the edges of the bounding box.) For the base case, when $n = 1$, then there is exactly one left bounding edge in $\ell$’s zone, and $1 \leq 3n$. Assume that the hypothesis is true for any set of $n - 1$ lines. Consider the rightmost line.
of the arrangement to intersect $\ell$. Call this $\ell_1$. (Selecting this particular line is very important for the proof.) Suppose that we consider the arrangement of the other $n-1$ lines. By the induction hypothesis there will be at most $3(n-1)$ left-bounding edges in the zone for $\ell$.

![Figure 14.3: Proof of the Zone Theorem.](image)

Now let us add back $\ell_1$ and see how many more left-bounding edges result. Consider the rightmost face of the arrangement of $n-1$ lines. (Shaded in the figure above.) Note that all of its edges are left-bounding edges. Line $\ell_1$ intersects $\ell$ within this face. By convexity, $\ell_1$ intersects the boundary of this face in two edges, denoted $e_a$ and $e_b$, where $e_a$ is above $\ell$ and $e_b$ is below. The insertion of $\ell_1$ creates a new left bounding edge along $\ell_1$ itself, and splits the left bounding edges $e_a$ and $e_b$ into two new left bounding edges for a net increase of three edges. Observe that $\ell_1$ cannot contribute any other left-bounding edges to the zone, because (depending on slope) either the line supporting $e_a$ or the line supporting $e_b$ blocks $\ell_1$’s visibility from $\ell$. (Note that it might provide right-bounding edges, but we are not counting them here.) Thus, the total number of left-bounding edges on the zone is at most $3(n-1) + 3 \leq 3n$, and hence the total number of edges is at most $6n$, as desired.
15 Applications of Arrangements

Reading: Chapter 8 in the 4M’s. Some of this material is not covered in the book.

Sweeping Arrangements: Last time we showed how to construct an arrangement of lines in the plane in $O(n^2)$ time. Since an arrangement is of size $O(n^2)$, this is optimal. Usually it is not sufficient just to build the arrangement, but necessary to traverse the arrangement as well. In some instances, any graph traversal will suffice. For other problems it is nice to perform the traversal in some order, for example as a plane sweep from left to right.

If an arrangement is to be built just so it can be swept, then maybe you didn’t need to construct the entire arrangement after all. You can just perform the plane sweep on the lines, exactly as we did for the line segment intersection algorithm. Assuming that we are sweeping from left to right, the initial position of the sweep line is at $x = -\infty$ (computed symbolically). The sweep line status maintains the lines in top to bottom order according to their intersection with the sweep line. (The initial order is according to increasing order of slope.) Then the sweep line proceeds exactly as it did in the algorithm for determining line segment intersections, but the only events are intersection events (since there are no endpoints). Sweeping an arrangement in this way takes $O(n^2 \log n)$ time, but only $O(n)$ space (since we need only maintain the current sweep line), and this may be an attractive alternative to constructing the entire subdivision (assuming the computational problem can be formulated in terms of sweeping).

But is the $O(\log n)$ factor necessary? In many applications involving plane sweep it is not necessary to sweep in strictly left-to-right order. A variant of the classical plane sweep, which is often applicable to problems on planar arrangements is called topological plane sweep. You can think of this as sweeping the arrangement with a “wiggly” sweep line, which need not be strictly vertical, but is subject to the constraint that it can intersect each line of the arrangement exactly once. (Such a sweep line will, for example, visit the vertices along each line in left-to-right order.) It can be shown that this sort of sweep can be performed in $O(n^2)$ time, thus saving a factor of $O(\log n)$. Note that it only works for line arrangements—it cannot be applied to the line segment intersection problem, for example.

![Figure 15.1: One possible topological ordering of the vertices of an arrangement.](image)

When is topological plane sweep applicable? Unfortunately, this is not easy to characterize without a more detailed investigation of this algorithm, which we do not have time for. Nonetheless, I will often declare (without proof) that a certain problem either can or cannot be solved by topological plane sweep.
Applications of Arrangements and Duality: The computational and mathematical tools that we have developed with geometric duality and arrangements allow a large number of problems to be solved. Here are some examples. Unless otherwise stated, all these problems can be solved in $O(n^2)$ time and $O(n^2)$ space by constructing a line arrangement, or in $O(n^2)$ time and $O(n)$ space through topological plane sweep.

General position test: Given a set of $n$ points in the plane, determine whether any three are collinear.

Minimum area triangle: Given a set of $n$ points in the plane, determine the minimum area triangle whose vertices are selected from these points.

Minimum $k$-corridor: Given a set of $n$ points, and an integer $k$, determine the narrowest pair of parallel lines that enclose at least $k$ points of the set. The distance between the lines can be defined either as the vertical distance between the lines or the perpendicular distance between the lines.

Visibility graph: Given line segments in the plane, we say that two points are visible if the interior of the line segment joining them intersects none of the segments. Given a set of $n$ non-intersecting line segments, compute the visibility graph, whose vertices are the endpoints of the segments, and whose edges are pairs of visible endpoints.

Maximum stabbing line: Given a set of $n$ line segments in the plane, compute the line that stabs (intersects) the maximum number of these line segments.

Hidden surface removal: Given a set of $n$ non-intersecting polygons in 3-space, imagine projecting these polygon onto a plane (either orthogonally or using perspective). Determine which portions of the polygons are visible from the viewpoint under this projection.

Note that in the worst case, the complexity of the final visible scene may be as high as $O(n^2)$, so this is asymptotically optimal. However, since such complex scenes rarely occur in practice, this algorithm is really only of theoretical interest.

Ham Sandwich Cut: Given $n$ red points and $m$ blue points, find a single line that simultaneously bisects these point sets. It is a famous fact from mathematics (called the Ham-Sandwich Theorem) that such a line always exists. If the point sets are separated by a line, then this can be done in time: $O(n \log n)$, space: $O(n+m)$.

Sorting all angular sequences: Here is a natural application of duality and arrangements that turns out to be important for the problem of computing visibility graphs. Consider a set of $n$ points in the plane. For each point $p$ in this set we want to perform an angular sweep, say in counterclockwise order, visiting the other $n-1$ points of the set. For each point, it is possible to compute the angles between this point and the remaining $n-1$ points and then sort these angles. This would take $O(n \log n)$ time per point, and $O(n^2 \log n)$ time overall.

With arrangements we can speed this up to $O(n^2)$ total time, getting rid of the extra $O(\log n)$ factor. Here is how. Recall the point-line dual transformation. A point $p = (p_x, p_y)$ and line $\ell : (y = ax - b)$ in the primal plane are mapped through to a dual point $\ell^*$ and dual line $p^*$ as follows:

$$\ell^* = (a, b)$$
$$p^* : (b = p_x a - p_y).$$

Suppose that $p$ is the point that we want to sort around, and let $p_1, p_2, \ldots, p_n$ be the points in final angular order about $p$. 

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Consider the arrangement defined by the dual lines $p_i^*$. How is this order revealed in the arrangement? Consider the dual line $p^*$, and its intersection points with each of the dual lines $p_i^*$. These form a sequence of vertices in the arrangement along $p^*$. Consider this sequence ordered from left to right. It would be nice if this order were the desired circular order, but this is not quite correct. It follows from the definition of our dual transformation that the $a$-coordinate of each of these vertices in the dual arrangement is the slope of some line of the form $pp_i$ in the primal plane. Thus, the sequence in which the vertices appear on the line is a slope ordering of the points about $p_i$, not an angular ordering.

However, given this slope ordering, we can simply test which primal points lie to the left of $p$ (that is, have a smaller $x$-coordinate in the primal plane), and separate them from the points that lie to the right of $p$ (having a larger $x$-coordinate). We partition the vertices into two sorted sequences, and then concatenate these two sequences, with the points on the right side first, and the points on the left side later. The resulting is an angular sequence starting with the angle $-90$ degrees and proceeding up to $+270$ degrees.

Thus, once the arrangement has been constructed, we can reconstruct each of the angular orderings in $O(n)$ time, for a total of $O(n^2)$ time.

**Narrowest 3-corridor:** As mentioned above, in this problem we are given a set $P$ of $n$ points in the plane, and an integer $k$, $1 \leq k \leq n$, and we wish to determine the narrowest pair of parallel lines that enclose at least $k$ points of the set. In this case we will define the vertical distance between the lines as the distance to minimize. (It is easy, but a bit tricky to adapt the algorithm for perpendicular distance.)

To simplify the presentation, we assume that $k = 3$. The generalization to general $k$ is an exercise. We will assume that no three points of $P$ are collinear. This will allow us to assume that the narrowest corridor contains exactly three points and has width strictly greater than zero. (Observe that if three points were collinear, then they would correspond to three lines that intersect at a common vertex in the arrangement, which could be tested as we build the arrangement.) We will also assume that no two points have the same $x$-coordinate. The dual transformation that we consider cannot represent vertical lines, but this assumption will imply that the solution is not a vertical line. We could fix this by using homogeneous coordinates, but we will not worry about it now.

If we dualize the points of $P$, then in dual space we have a set $L$ of $n$ lines, $\{\ell_1, \ell_2, \ldots, \ell_n\}$. The slope of each dual-line is the $x$-coordinate of the corresponding point of $P$, and its $y$-intercept is the negation of the point’s $y$-coordinate.

A narrowest 3-corridor in the primal plane consists of two parallel lines $\ell_a$ and $\ell_b$ in primal space. Their duals $\ell^*_a$ and $\ell^*_b$ are dual points, which have the same $x$-coordinates (since the lines are parallel),
and the vertical distance between these points, is the difference in the $y$-intercepts of the two primal lines. Thus the vertical width of the corridor, is the vertical length of the line segment.

In the primal plane, there are exactly three points lying in the corridor, that is, there are three points that are both above $\ell_b$ and below $\ell_a$. Thus, by the order reversing property, in the dual plane, there are three dual lines that pass both below point $\ell^*_b$ and above $\ell^*_a$. Combining all these observations it follows that the dual formulation of the narrowest 3-corridor problem is the following:

**Shortest vertical 3-stabber:** Given an arrangement of $n$ lines, determine the shortest vertical segment that stabs three lines of the arrangement.

![Primal Plane and Dual Plane](image.png)

Figure 15.3: Narrowest 3-corridor: primal and dual form.

It is easy to show (by a simple perturbation argument) that the shortest vertical 3-stabber may be assumed to have one of its endpoints on a vertex of the arrangement, implying that the other endpoint lies on the line of the arrangement lying immediately above or below this vertex. (In the primal plane the significance is that we can assume that the minimum 3-corridor one of the lines passes through 2 of the points, and the other passes through a third point, and there are no points within the interior of the corridor. This is shown in the figure below.

We can compute the minimum 3-stabber in an arrangement, by a simple plane sweep of the arrangement (using a vertical sweep line). Whenever we encounter a vertex of the arrangement, we consider the distance to the edge of the arrangement lying immediately above this vertex and the edge lying immediately below. We can solve this problem by topological plane sweep in $O(n^2)$ time and $O(n)$ space.

We can also solve this by constructing the arrangement and the computing the (vertical) trapezoidal map. Each trapezoidal edge will correspond to a corridor. The shortest such edge is the final answer. This leads to an $O(n^2)$ time and space solution.

![Narrowest Corridor and Trapezoidal Maps](image.png)

Figure 15.4: Narrowest corridor and trapezoidal maps.
Maximum Discrepancy: Next we consider a problem derived from computer graphics and sampling. Suppose that we are given a collection of \( n \) points \( S \) lying in a unit square \( U = [0, 1]^2 \). We want to use these points for random sampling purposes. In particular, the property that we would like these points to have is that for any halfplane \( h \), we would like the size of the fraction of points of \( P \) that lie within \( h \) should be roughly equal to the area of intersection of \( h \) with \( U \). That is, if we define \( \mu(h) \) to be the area of \( h \cap U \), and \( \mu_S(h) = |S \cap h|/|S| \), then we would like \( \mu(h) \approx \mu_S(h) \) for all \( h \). This property is important when point sets are used for things like sampling and Monte-Carlo integration.

To this end, we define the discrepancy of \( S \) with respect to a halfplane \( h \) to be

\[
\Delta_S(h) = |\mu(h) - \mu_S(h)|.
\]

For example, in the figure below (a), the area of \( h \cap U \) is \( \mu(h) = 0.625 \), and there are 7 out of 13 points in \( h \), thus \( \mu_S(h) = 7/13 = 0.538 \). Thus the discrepancy of \( h \) is \( |0.625 - 0.538| = 0.087 \).

Define the halfplane discrepancy of \( S \) to be the maximum (or more properly the supremum, or least upper bound) of this quantity over all halfplanes:

\[
\Delta(S) = \sup_h \Delta_S(h).
\]

Figure 15.5: Discrepancy of a point set.

Since there are an uncountably infinite number of halfplanes, it is important to derive some sort of finiteness criterion on the set of halfplanes that might produce the greatest discrepancy.

**Lemma:** Let \( h \) denote the halfplane that generates the maximum discrepancy with respect to \( S \), and let \( \ell \) denote the line that bounds \( h \). Then either (i) \( \ell \) passes through at least two points of \( S \), or (ii) \( \ell \) passes through one point of \( S \), and this point is the midpoint of the line segment \( \ell \cap U \).

**Remark:** If a line passes through one or more points of \( S \), then should this point be included in \( \mu_S(h) \)? For the purposes of computing the maximum discrepancy, the answer is to either include or omit the point, whichever will generate the larger discrepancy. The justification is that it is possible to perturb \( h \) infinitesimally so that it includes none or all of these points without altering \( \mu(h) \).

**Proof:** If \( \ell \) does not pass through any point of \( S \), then (depending on which is larger \( \mu(h) \) or \( \mu_S(h) \)) we can move the line up or down without changing \( \mu_S(h) \) and increasing or decreasing \( \mu(h) \) to increase their difference. If \( \ell \) passes through a point \( p \in S \), but is not the midpoint of the line segment \( \ell \cap U \), then we claim that we can rotate this line about \( p \) and hence increase or decrease \( \mu(h) \) without altering \( \mu_S(h) \), to increase their difference.

To establish the claim, consider the figure above (b). Suppose that the line \( \ell \) passes through point \( p \) and let \( r_1 < r_2 \) denote the two lengths along \( \ell \) from \( p \) to the sides of the square. Observe that
if we rotate $\ell$ through a small angle $\theta$, then to a first order approximation, the loss due to area of the triangle on the left is $r_1^2 \theta / 2$, since this triangle can be approximated by an angular sector of a circle of radius $r_1$ and angle $\theta$. (CHECK: Based on the figure do we mean "decrease" rather than "gain" in the next sentence.) The gain due to the area of the triangle on the right is $r_2^2 \theta / 2$. Thus, since $r_1 < r_2$ this rotation will increase the area of region lying below $h$ infinitesimally. A rotation in the opposite decreases the area infinitesimally. Since the number of points bounded by $h$ does not change as a function of $\theta$, the discrepancy cannot be achieved as long as such a rotation is possible.

(Note that this proof reveals something slightly stronger. If $\ell$ contacts two points, the line segment between these points must contain the midpoint of the $\ell \cap U$. Do you see why?)

Since for each point $p \in S$ there are only a constant number of lines $\ell$ (at most two, I think) through this point such that $p$ is the midpoint of $\ell \cap U$, it follows that there are at most $O(n)$ lines of type (i) above, and hence the discrepancy of all of these lines can be tested in $O(n^2)$ time.

To compute the discrepancies of the other lines, we can dualize the problem. In the primal plane, a line $\ell$ that passes through two points $p_i, p_j \in S$, is mapped in the dual plane to a point $\ell^*$ at which the lines $p_i^*$ and $p_j^*$ intersect. This is just a vertex in the arrangement of the dual lines for $S$. So, if we have computed the arrangement, then all we need to do is to visit each vertex and compute the discrepancy for the corresponding primal line.

It is easy to see that the area $\ell \cap U$ of each corresponding line in the primal plane can be computed in $O(1)$ time. So, all that is needed is to compute the number of points of $S$ lying below each such line. In the dual plane, the corresponds to determining the number of dual lines that lie below or above each vertex in the arrangement. If we know the number of dual lines that lie strictly above each vertex in the arrangement, then it is trivial to compute the number of lines that lie below by subtraction.

We define a point to be at level $k$, denoted $L_k$, in an arrangement if there are at most $k - 1$ lines above this point and at most $n - k$ lines below this point. The $k$-th level of an arrangement is an $x$-monotone polygonal curve, as shown below. For example, the upper envelope of the lines is level 1 of the arrangement, and the lower envelope is level $n$ of the arrangement. Note that a vertex of the arrangement can be on multiple levels. (Also note that our definition of level is exactly one greater than our text’s definition.)

![Figure 15.6: Examples of levels in an arrangement.](image)

We claim that it is an easy matter to compute the level of each vertex of the arrangement (e.g. by plane sweep). The initial levels at $x = -\infty$ are determined by the slope order of the lines. As the plane
sweep proceeds, the index of a line in the sweep line status is its level. Thus, by using topological plane sweep, in $O(n^2)$ time we can compute the minimum and maximum level number of each vertex in the arrangement. From the order reversing property, for each vertex of the dual arrangement, the minimum level number minus one indicates the number of primal points that lie strictly below the corresponding primal line and the maximum level number is the number of primal points that lie on or below this line. Thus, given the level numbers and the fact that areas can be computed in $O(1)$ time, we can compute the discrepancy in $O(n^2)$ time and $O(n)$ space, through topological plane sweep.
16 Orthogonal Range Searching and kd-Trees

Reading: Chapter 5 in the 4M’s.

Range Queries: We will shift our focus from algorithm problems to data structures for the next few lectures. We will consider the following class of problems. Given a collection of objects, preprocess them (storing the results in a data structure of some variety) so that queries of a particular form can be answered efficiently. Generally we measure data structures in terms of two quantities, the time needed to answer a query and the amount of space needed by the data structure. Often there is a trade-off between these two quantities, but most of the structures that we will be interested in will have either linear or near linear space. Preprocessing time is an issue of secondary importance, but most of the algorithms we will consider will have either linear or \( O(n \log n) \) preprocessing time.

In a range queries we are given a set \( P \) of points and region \( R \) in space (e.g., a rectangle, polygon, halfspace, or disk) and are asked list (or count or compute some accumulation function of) the subset of \( P \) lying within the region. To get the best possible performance, the design of the data structure is tailored to the particular type of region, but there are some data structures that can be used for a wide variety of regions.

An important concept behind all geometric range searching is that the subsets that can be formed by simple geometric ranges is much smaller than the set of possible subsets (called the power set) of \( P \). We can define any range search problem abstractly as follows. Given a particular class of ranges, a range space is a pair \((P, R)\) consisting of the points \( P \) and the collection \( R \) of all subsets of \( P \) that be formed by ranges of this class. For example, the following figure shows the range space assuming rectangular ranges for a set of points in the plane. In particular, note that the sets \( \{1,4\} \) and \( \{1,2,4\} \) cannot be formed by rectangular ranges. Observe that each rectangular range is effectively defined by at most four points along each of its four sides, which implies that there are \( O(n^4) \) distinct ranges in this space, which is generally much smaller than the \( 2^n \) power set.

![Figure 16.1: Rectangular range space.](image)

Today we consider orthogonal rectangular range queries, that is, ranges defined by rectangles whose sides are aligned with the coordinate axes. One of the nice things about rectangular ranges is that they can be decomposed into a collection of 1-dimensional searches.

Canonical Subsets: A common approach used in solving almost all range queries is to represent \( P \) as a collection of canonical subsets \( \{S_1, S_2, \ldots, S_k\} \), each \( S_i \subseteq S \) (where \( k \) is generally a function of \( n \) and the type of ranges), such that any set can be formed as the disjoint union of canonical subsets. Note that these subsets may generally overlap each other.

There are many ways to select canonical subsets, and the choice affects the space and time complexities. For example, the canonical subsets might be chosen to consist of \( n \) singleton sets, each of the...
form \{p_i\}. This would be very space efficient, since we need only \(O(n)\) total space to store all the canonical subsets, but in order to answer a query involving \(k\) objects we would need \(k\) sets. (This might not be bad for reporting queries, but it would be too long for counting queries.) At the other extreme, we might let the canonical subsets be the power set of \(P\). Now, any query could be answered with a single canonical subset (assuming we could determine which one), but we would have \(2^n\) different canonical subsets to store. (A more realistic solution would be to use the set of all ranges, but this would still be quite large for most interesting range spaces.) The goal of a good range data structure is to strike a balance between the total number of canonical subsets (space) and the number of canonical subsets needed to answer a query (time).

**One-dimensional range queries:** Before we consider how to solve general range queries, let us consider how to answer 1-dimension range queries, or *interval queries*. Let us assume that we are given a set of points \(P = \{p_1, p_2, \ldots, p_n\}\) on the line, which we will preprocess into a data structure. Then, given an interval \([x_{lo}, x_{hi}]\), the goal is to report all the points lying within the interval. Ideally we would like to answer a query in time \(O(\log n + k)\) time, where \(k\) is the number of points reported (an output sensitive result). Range counting queries can be answered in \(O(\log n)\) time with minor modifications.

Clearly one way to do this is to simply sort the points, and apply binary search to find the first point of \(P\) that is greater than or equal to \(x_{lo}\), and less than or equal to \(x_{hi}\), and then list all the points between. This works fine in dimension 1, but does not generalize readily to any higher dimensions.

Let us consider a different approach, which will generalize to higher dimensions. Sort the points of \(P\) in increasing order and store them in the leaves of a balanced binary search tree. Each internal node of the tree is labeled with the largest key appearing in its left child. We can associate each node of this tree (implicitly or explicitly) with the subset of points stored in the leaves that are descendants of this node. This gives rise to the \(O(n)\) canonical subsets. For now, these canonical subsets will not be stored explicitly as part of the data structure, but this will change later when we talk about range trees. This is illustrated in the figure below.

![Figure 16.2: Canonical sets for interval queries.](https://example.com/figure.png)

We claim that the canonical subsets corresponding to any range can be identified in \(O(\log n)\) time from this structure. Given any interval \([x_{lo}, x_{hi}]\), we search the tree to find the leftmost leaf \(u\) whose key is greater than or equal to \(x_{lo}\) and the rightmost leaf \(v\) whose key is less than or equal to \(x_{hi}\). Clearly all the leaves between \(u\) and \(v\), together possibly with \(u\) and \(v\), constitute the points that lie within the range. If \(\text{key}(u) = x_{lo}\) then we include \(u\)'s canonical (single point) subset and if \(\text{key}(v) = x_{hi}\) then we do the same for \(v\). To form the remaining canonical subsets, we take the subsets of all the maximal subtrees lying between \(u\) and \(v\).
Here is how to compute these subtrees. The search paths to \( u \) and \( v \) may generally share some common subpath, starting at the root of the tree. Once the paths diverge, as we follow the left path to \( u \), whenever the path goes to the left child of some node, we add the canonical subset associated with its right child. Similarly, as we follow the right path to \( v \), whenever the path goes to the right child, we add the canonical subset associated with its left child.

To answer a range reporting query we simply traverse these canonical subtrees, reporting the points of their leaves. Each tree can be traversed in time proportional to the number of leaves in each subtree. To answer a range counting query we store the total number of points in each subtree (as part of the preprocessing) and then sum all of these over all the canonical subtrees.

Since the search paths are of length \( O(\log n) \), it follows that \( O(\log n) \) canonical subsets suffice to represent the answer to any query. Thus range counting queries can be answered in \( O(\log n) \) time.

In summary, 1-dimensional range queries can be answered in \( O(\log n) \) time, using \( O(n) \) storage. This concept of finding maximal subtrees that are contained within the range is fundamental to all range search data structures. The only question is how to organize the tree and how to locate the desired sets. Let see next how we can extend this to higher dimensional range queries.

**Kd-trees:** The natural question is how to extend 1-dimensional range searching to higher dimensions. First we will consider kd-trees. This data structure is easy to implement and quite practical and useful for many different types of searching problems (nearest neighbor searching for example). However it is not the asymptotically most efficient solution for the orthogonal range searching, as we will see later.

Our terminology is a bit nonstandard. The data structure was designed by Jon Bentley. In his notation, these were called “k-d trees,” short for “k-dimensional trees”. The value \( k \) was the dimension, and thus there are 2-d trees, 3-d trees, and so on. However, over time, the specific value of \( k \) was lost. Our text uses the somewhat nonstandard form “kd-tree” rather than “k-d tree.” By the way, there are many variants of the kd-tree concept. We will describe the most commonly used one, which is quite similar to Bentley’s original design. In our trees, points will be stored only at the leaves. There are variants in which points are stored at internal nodes as well.

The idea behind a kd-tree is to extend the notion of a 1-dimensional tree. But for each node we subdivide space either by splitting along \( x \)-coordinate of the points or along the \( y \)-coordinates. Each internal node \( t \) of the kd-tree is associated with the following quantities:

- \( t.cutDim \) the cutting dimension
- \( t.cutVal \) the cutting value
- \( t.size \) the number of points in \( t \)’s subtree

In dimension \( d \), the cutting dimension may be represented as in integer ranging from 0 to \( d-1 \). If the cutting dimension is \( i \), then all points whose \( i \)th coordinate is less than or equal to \( t.cutVal \) are stored in the left subtree and the remaining points are stored in the right subtree. (See the figure below.) If a point’s coordinate is equal to the cutting value, then we may allow the point to be stored on either side. This is done to allow us to balance the number of points in the left and right subtrees if there are many equal coordinate values. When a single point remains (or more generally a small constant number of
points), we store it in a leaf node, whose only field \textit{t.point} is this point (or generally a pointer to this point).

![Subdivision]

**Figure 16.3:** A kd-tree and the associated spatial subdivision.

The cutting process has a geometric interpretation. Each node of the tree is associated implicitly with a rectangular region of space, called a \textit{cell}. (In general these rectangles may be unbounded, but in many applications it is common to restrict ourselves to some bounded rectangular region of space before splitting begins, and so all these rectangles are bounded.) The cells are nested in the sense that a child’s cell is contained within its parent’s cell. Hence, these cells define a \textit{hierarchical decomposition} of space. This is illustrated in figure.

There are two key decisions in the design of the tree.

**How is the cutting dimension chosen?** The simplest method is to cycle through the dimensions one by one. (This method is shown in the above figure.) Since the cutting dimension depends only on the level of a node in the tree, one advantage of this rule is that the cutting dimension need not be stored explicitly in each node, instead we keep track of it while traversing the tree.

One disadvantage of this splitting rule is that, depending on the data distribution, this simple cyclic rule may produce very skinny (elongated) cells, and such cells may adversely affect query times. Another method is to select the cutting dimension to be the one along which the points have the greatest \textit{spread}, defined to be the difference between the largest and smallest coordinates. Bentley call the resulting tree an \textit{optimized kd-tree}.

**How is the cutting value chosen?** To guarantee that the tree has height \(O(\log n)\), the best method is to let the cutting value be the median coordinate along the cutting dimension. If there are an even number of points in the subtree, we may take either the upper or lower median, or we may simply take the midpoint between these two points. In our example, when there are an odd number of points, the median is associated with the left (or lower) subtree.

A kd-tree is a special case of a more general class of hierarchical spatial subdivisions, called \textit{binary space partition trees} (or \textit{BSP trees}) in which the splitting lines (or hyperplanes in general) may be oriented in any direction. In the case of BSP trees, the cells are convex polygons.

**Constructing the kd-tree:** It is possible to build a kd-tree in \(O(n \log n)\) time by a simple top-down recursive procedure. The most costly step of the process is determining the median coordinate for splitting purposes. One way to do this is to maintain two lists of pointers to the points, one sorted by \(x\)-coordinate and the other containing pointers to the points sorted according to their \(y\)-coordinates. (In dimension \(d\), \(d\) such arrays would be maintained.) Using these two lists, it is an easy matter to find the median at each step in constant time. In linear time it is possible to split each list about this median
element. For example, if \( x = s \) is the cutting value, then all points with \( p_x \leq s \) go into one list and those with \( p_x > s \) go into the other. (In dimension \( d \) this generally takes \( O(d) \) time per point.) This leads to a recurrence of the form \( T(n) = 2T(n/2) + n \), which solves to \( O(n \log n) \). Since there are \( n \) leaves and each internal node has two children, it follows that the number of internal nodes is \( n - 1 \). Hence the total space requirements are \( O(n) \).

**Theorem:** A balanced kd-tree of \( n \) points can be constructed in \( O(n \log n) \) time and has \( O(n) \) space.

**Range Searching in kd-trees:** Let us consider how to answer orthogonal range counting queries. Range reporting queries are an easy extension. Let \( Q \) denote the desired range, \( t \) denote the current node in the kd-tree, and let \( C \) denote the rectangular cell associated with \( t \). The search algorithm traverses the tree recursively. If it arrives at a leaf cell, we check to see whether the associated point, \( t.\text{point} \), lies within \( Q \) in \( O(1) \) time, and if so we count it. Otherwise, \( t \) is an internal node. If \( t \)'s cell \( C \) is disjoint from \( Q \) (which can be tested in \( O(1) \) time since both are rectangles), then we know that no point in the subtree rooted at \( t \) is in the query range, and so there is nothing to count. If \( C \) is entirely contained within \( Q \) (again testable in \( O(1) \) time), then every point in the subtree rooted at \( t \) can be counted. (These points constitute a canonical subset.) Otherwise, \( t \)'s cell partially overlaps \( Q \). In this case we recurse on \( t \)'s two children and update the count accordingly.

```c
int rangeCount(Range Q, KDNode t, Rectangle C)
    (1) if (t is a leaf)
        (a) if (Q contains t.\text{point}) return 1,
        (b) else return 0.
    (2) if (t is not a leaf)
        (a) if (C \( \cap \) Q = \( \emptyset \)) return 0.
        (b) else if (C \( \subseteq \) Q) return t.\text{size}.
        (c) else, split C along \( t \)'s cutting dimension and cutting value, letting \( C_1 \) and \( C_2 \) be the two rectangles. Return
            \( \text{rangeCount}(Q,t.\text{left},C_1) + \text{rangeCount}(Q,t.\text{right},C_2) \).
```

The figure below shows an example of a range search. White nodes have been visited in the search. Light shaded nodes were not visited because their cell is contained entirely within \( Q \). Dark shaded nodes are not visited because their cell is disjoint from \( Q \).

![Figure 16.4: Range search in a kd-tree. (Note: This particular tree was not generated by the algorithm described above.)](image_url)
Analysis of query time: How many nodes does this method visit altogether? We claim that the total number of nodes is \( O(\sqrt{n}) \) assuming a balanced kd-tree. Recall from the discussion above that a node is processed (both children visited) if and only if the cell overlaps the range without being contained within the range. We say that such a cell is \textit{stabbed} by the query. To bound the total number of nodes that are processed in the search, we first bound the number of nodes that are stabbed.

Lemma: Given a balanced kd-tree with \( n \) points using the alternating splitting rule, any vertical or horizontal line stabs \( O(\sqrt{n}) \) cells of the tree.

Proof: Let us consider the case of a vertical line \( x = x_0 \). The horizontal case is symmetrical.

Consider a processed node which has a cutting dimension along \( x \). The vertical line \( x = x_0 \) either stabs the left child or the right child but not both. If it fails to stab one of the children, then it cannot stab any of the cells belonging to the descendents of this child either. If the cutting dimension is along the \( y \)-axis (or generally any other axis in higher dimensions), then the line \( x = x_0 \) stabs both children’s cells.

Since we alternate splitting on left and right, this means that after descending two levels in the tree, we may stab at most two of the possible four grandchildren of each node. In general each time we descend two more levels we double the number of nodes being stabbed. Thus, we stab the root node, at most 2 nodes at level 2 of the tree, at most 4 nodes at level 4, 8 nodes at level 6, and generally at most \( 2^i \) nodes at level \( 2^i \).

Because we have an exponentially increasing number, the total sum is dominated by its last term. Thus is suffices to count the number of nodes stabbed at the lowest level of the tree. Since the tree is balanced, it has roughly \( \lg n \) levels. Thus the number of leaf nodes processed at the bottommost level is \( 2^{(\lg n)/2} = 2^{\lg \sqrt{n}} = \sqrt{n} \). This completes the proof.

We have shown that any vertical or horizontal line can stab only \( O(\sqrt{n}) \) cells of the tree. Thus, if we were to extend the four sides of \( Q \) into lines, the total number of cells stabbed by all these lines is at most \( O(4\sqrt{n}) = O(\sqrt{n}) \). Thus the total number of cells stabbed by the query range is \( O(\sqrt{n}) \). Since we only make recursive calls when a cell is stabbed, it follows that the total number of nodes visited by the search is \( O(\sqrt{n}) \).

Theorem: Given a balanced kd-tree with \( n \) points, orthogonal range counting queries can be answered in \( O(\sqrt{n}) \) time and reporting queries can be answered in \( O(\sqrt{n} + k) \) time. The data structure uses space \( O(n) \).
17 Orthogonal Range Trees

Reading: Chapter 5 in the 4M’s.

Orthogonal Range Trees: Last time we saw that kd-trees could be used to answer orthogonal range queries in the plane in $O(\sqrt{n} + k)$ time. Today we consider a data structure, which is more highly tuned to this particular problem, called an orthogonal range tree.

An orthogonal range tree is a data structure which, in all dimensions $d \geq 2$, uses $O(n \log^{(d-1)} n)$ space, and can answer orthogonal rectangular range queries in $O(\log^{(d-1)} n + k)$ time, where $k$ is the number of points reported. Preprocessing time is the same as the space bound. Thus, in the plane, we can answer range queries in time $O(\log n)$ and space $O(n \log n)$. We will present the data structure in two parts, the first is a version that can answer queries in $O(\log^2 n)$ time and then we will show how to improve this in order to strip off a factor of $\log n$ from the query time.

Multi-level Search Trees: The data structure is based on the concept of a multi-level search tree. In this method, a complex search is decomposed into a constant number of simpler range searches. We cascade a number of search structures for simple ranges together to answer the complex range query. In this case we will reduce a $d$-dimensional range search to a series of 1-dimensional range searches.

Suppose you have a query which can be stated as the intersection of a small number of simpler queries. For example, a rectangular range query in the plane can be stated as two range queries: Find all the points whose $x$-coordinates are in the range $[Q.x_{lo}, Q.x_{hi}]$ and all the points whose $y$-coordinates are in the range $[Q.y_{lo}, Q.y_{hi}]$. Let us consider how to do this for 2-dimensional range queries, and then consider how to generalize the process. First, we assume that we have preprocessed the data by building a range tree for the first range query, which in this case is just a 1-dimensional range tree for the $x$-coordinates. Recall that this is just a balanced binary tree on these points sorted by $x$-coordinates. Also recall that each node of this binary tree is implicitly associated with a canonical subset of points, that is, the points lying in the subtree rooted at this node.

Observe that the answer to any 1-dimensional range query can be represented as the disjoint union of a small collection of $m = O(\log n)$ canonical subsets, $\{S_1, S_2, \ldots, S_m\}$, where each subset corresponds to a node $t$ in the range tree. This constitutes the first level of the search tree. To continue the preprocessing, for the second level, for each node $t$ in this $x$-range tree, we build an auxiliary tree, $t_{aux}$, each of which is a $y$-coordinate range tree, which contains all the points in the canonical subset associated with $t$.

The final data structure consists of two levels: an $x$-range tree, such that each node in this tree points to auxiliary $y$-range tree. This notion of a tree of trees is basic to solving range queries by leveling. (For example, for $d$-dimensional range trees, we will have $d$-levels of trees.)

Query Processing: Given such a 2-level tree, let us consider how to answer a rectangular range query $Q$. First, we determine the nodes of the tree that satisfy the $x$ portion of the range query. Each such node $t$ is associated with a canonical set $S_t$, and the disjoint union of these $O(\log n)$ sets consistute all the points of the data set that lie within the $x$ portion of the range. Thus to finish the query, we need to find out which points from each canonical subset lie within the range of $y$-coordinates. To do this, for each such node $t$, we access the auxiliary tree for this node, $t_{aux}$ and perform a 1-dimensional range search on the $y$-range of the query. This process is illustrated in the following figure.
What is the query time for a range tree? Recall that it takes $O(\log n)$ time to locate the nodes representing the canonical subsets for the 1-dimensional range query. For each, we invoke a 1-dimensional range search. Thus there are $O(\log n)$ canonical sets, each invoking an $O(\log n)$ range search, for a total time of $O(\log^2 n)$. As before, listing the elements of these sets can be performed in additional $O(k)$ time by just traversing the trees. Counting queries can be answered by precomputing the subtree sizes, and then just adding them up.

**Space:** The space used by the data structure is $O(n \log n)$ in the plane (and $O(n \log^{(d-1)} n)$ in dimension $d$). The reason comes by summing the sizes of the two data structures. The tree for the $x$-coordinates requires only $O(n)$ storage. But we claim that the total storage in all the auxiliary trees is $O(n \log n)$. We want to count the total sizes of all these trees. The number of nodes in a tree is proportional to the number of leaves, and hence the number of points stored in this tree. Rather than count the number of points in each tree separately, instead let us count the number of trees in which each point appears. This will give the same total. Observe that a point appears in the auxiliary trees of each of its ancestors. Since the tree is balanced, each point has $O(\log n)$ ancestors, and hence each point appears in $O(\log n)$ auxiliary trees. It follows that the total size of all the auxiliary trees is $O(n \log n)$. By the way, observe that because the auxiliary trees are just 1-dimensional trees, we could just store them as a sorted array.

We claim that it is possible to construct a 2-dimensional range tree in $O(n \log n)$ time. Constructing the 1-dimensional range tree for the $x$-coordinates is easy to do in $O(n \log n)$ time. However, we need to be careful in constructing the auxiliary trees, because if we were to sort each list of $y$-coordinates separately, the running time would be $O(n \log^2 n)$. Instead, the trick is to construct the auxiliary trees in a bottom-up manner. The leaves, which contain a single point are trivially sorted. Then we simply merge the two sorted lists for each child to form the sorted list for the parent. Since sorted lists can be merged in linear time, the set of all auxiliary trees can be constructed in time that is linear in their total since, or $O(n \log n)$. Once the lists have been sorted, then building a tree from the sorted list can be done in linear time.

**Multilevel Search and Decomposable Queries:** Summarizing, here is the basic idea to this (and many other query problems based on leveled searches). Let $(S, R)$ denote the range space, consisting of points $S$ and range space $R$. Suppose that each range of $R$ can be expressed as an intersection of...
simpler ranges \( R = R_1 \cap R_2 \cap \ldots \cap R_c \), and assume that we know how to build data structures for each of these simpler ranges.

The multilevel search structure is built by first building a range search tree for query \( R_1 \). In this tree, the answer to any query can be represented as the disjoint union of some collection \( \{ S_1, S_2, \ldots, S_m \} \) of canonical subsets, each a subset of \( S \). Each canonical set corresponds to a node of the range tree for \( R_1 \).

For each node \( t \) of this range tree, we build an auxiliary range tree for the associated canonical subset \( S_t \) for ranges of class \( R_2 \). This forms the second level. We can continue in this manner, with each node of the range tree for the ranges \( R_t \) being associated with an auxiliary range tree for ranges \( R_{t+1} \).

To answer a range query, we solve the first range query, resulting in a collection of canonical subsets whose disjoint union is the answer to this query. We then invoke the second range query problem on each of these canonical subsets, and so on. Finally we take the union of all the answers to all these queries.

**Fractional Cascading:** Can we improve on the \( O(\log^2 n) \) query time? We would like to reduce the query time to \( O(\log n) \). As we descend the search the \( x \)-interval tree, for each node we visit, we need to search the corresponding \( y \)-interval tree. It is this combination that leads to the squaring of the logarithms. If we could search each \( y \)-interval in \( O(1) \) time, then we could eliminate this second log factor. The trick to doing this is used in a number of places in computational geometry, and is generally a nice idea to remember. We are repeatedly searching different lists, but always with the same key. The idea is to merge all the different lists into a single massive list, do one search in this list in \( O(\log n) \) time, and then use the information about the location of the key to answer all the remaining queries in \( O(1) \) time each. This is a simplification of a more general search technique called fractional cascading.

In our case, the massive list on which we will do one search is the entire set of points, sorted by \( y \)-coordinate. In particular, rather than store these points in a balanced binary tree, let us assume that they are just stored as sorted arrays. (The idea works for either trees or arrays, but the arrays are a little easier to visualize.) Call these the auxiliary lists. We will do one (expensive) search on the auxiliary list for the root, which takes \( O(\log n) \) time. However, after this, we claim that we can keep track of the position of the \( y \)-range in each auxiliary list in constant time as we descend the tree of \( x \)-coordinates.

Here is how we do this. Let \( v \) be an arbitrary internal node in the range tree of \( x \)-coordinates, and let \( v_L \) and \( v_R \) be its left and right children. Let \( A_v \) be the sorted auxiliary list for \( v \) and let \( A_L \) and \( A_R \) be the sorted auxiliary lists for its respective children. Observe that \( A_v \) is the disjoint union of \( A_L \) and \( A_R \) (assuming no duplicate \( y \)-coordinates). For each element in \( A_v \), we store two pointers, one to the item of equal or larger value in \( A_L \) and the other to the item of equal or larger value in \( A_R \). (If there is no larger item, the pointer is null.) Observe that once we know the position of an item in \( A_v \), then we can determine its position in either \( A_L \) or \( A_R \) in \( O(1) \) additional time.

Here is a quick illustration of the general idea. Let \( v \) denote a node of the \( x \)-tree, and let \( v_L \) and \( v_R \) denote its left and right children. Suppose that (in bottom to top order) the associated nodes within this range are: \( \langle p_1, p_2, p_3, p_4, p_5, p_6 \rangle \), and suppose that in \( v_L \) we store the points \( \langle p_2, p_3, p_5 \rangle \) and in \( v_R \) we store \( \langle p_1, p_4, p_6 \rangle \). (This is shown in the figure below.) For each point in the auxiliary list for \( v \), we store a pointer to the lists \( v_L \) and \( v_R \), to the position this element would be inserted in the other list.
(assuming sorted by $y$-values). That is, we store a pointer to the largest element whose $y$-value is less than or equal to this point.

![Figure 17.2: Cascaded search in range trees.](image)

At the root of the tree, we need to perform a binary search against all the $y$-values to determine which points lie within this interval, for all subsequent levels, once we know where the $y$-interval falls with respect to the order points here, we can drop down to the next level in $O(1)$ time. Thus (as with fractional cascading) the running time is $O(2 \log n)$, rather than $O(\log^2 n)$. It turns out that this trick can only be applied to the last level of the search structure, because all other levels need the full tree search to compute canonical sets.

**Theorem:** Given a set of $n$ points in $R^d$, orthogonal rectangular range queries can be answered in $O(\log^{(d-1)} n + k)$ time, from a data structure of size $O(n \log^{(d-1)} n)$ which can be constructed in $O(n \log^{(d-1)} n)$ time.
18 Geometric Approximations and WSPDs

Reading: This material is not covered in our text. My presentation is taken from lecture notes by Sariel Har-Peled (which do not appear to be online anymore). The original paper on WSPDs is “A Decomposition of Multidimensional Point Sets with Applications to $k$-Nearest-Neighbors and $n$-Body Potential Fields,” by P. B. Callahan and S. Rao Kosaraju, *J. ACM*, 42, 67–90, 1995.

Approximation Algorithms in Computational Geometry: For the next few lectures we will consider an emerging area of computational geometry, namely geometric approximation algorithms. What is geometric approximation? Traditionally, approximation refers to approximate solutions to optimization problems. An optimization problem involves computing a discrete structure of some special type that minimizes or maximizes a given objective function, that is, a function representing the cost or profit of the solution. A geometric example would be computing the minimum traveling salesperson tour of a set of points in the plane, that is, a path of minimum length path that visits every point of the set. There are no polynomial time solutions to this problem, and the general problem is NP-hard. In this context, approximation means finding a feasible solution to the problem, but one that is (ideally) only moderately larger (assuming a minimization problem) than the optimal solution with respect to the cost function.

In computational geometry, approximation can have a second meaning. This involves computing a geometric structure that approximates some desired structure. An example of such a problem would be, given a set of $n$ points in space, output a convex polytope that is an approximation to the convex hull of the point set. Of course, to make this precise, we would need to make clear what it means for one shape to approximate another. (For example, we might require that the distance from every point of the boundary of the approximate hull be within some small distance of the boundary of the true convex hull.) In general, the notion of what it means to approximate a geometric structure depends very much on the purposes for which the structure is being used.

When is it acceptable to produce an approximate result? Of course, this depends on the desired application of the algorithm’s output. Typically, the goal of approximation is to replace exact, slow computations with approximate, faster computations. Note that approximation is often acceptable in geometry problem solving. One reason is that geometric inputs are often subject to the limits of accuracy of the measurement devices, and hence, our only approximations to begin with. It does not make sense to use inputs of low precision to derive answers of high precision. Another reason is that the output of a geometric algorithm may not be an end result, but merely an intermediate result to guide a larger computation, which itself has many sources of errors. If it is known that the output from one algorithm will be fed into another program, and this second program is an approximation or a heuristic, exactness may not be needed in the first program. This often happens with geometric algorithms are used as preprocessing steps in inexact or heuristic pattern analysis programs.

WSPD Motivation: We begin our discussion of approximation algorithms in geometry with a simple and powerful example. To motivate this example, consider an application in physics involving the simulation of the motions of a large collection of bodies (e.g., planets or stars) subject to their own mutual gravitational forces. In physics, such a simulation is often called the $n$-body problem. Exact analytical solutions are known to exist in only extremely small special cases. Even determining a good numerical solution is relative costly. In order to determine the motion of a single object in the simulation, we need to know the gravitational force induced by the other $n - 1$ bodies of the system. In order to
compute this force, it would seem that at a minimum we would need $O(n)$ computations per point, for a total of $O(n^2)$ total computations. The question is whether there is a way to do this faster?

An approximate answer to this question was discovered by Greengard and Rokhlin, and forms the basis of a technique called the fast multipole method (or FMM for short). We will not discuss the FMM, since it would take us out of the way, but will instead discuss the geometric structure that encodes much of the information that made the FMM such a popular technique. Furthermore, this structure, called a well-separated pair decomposition, or WSPD for short, is itself useful for solving a number of geometric problems.

Well Separated Pairs: A set of $n$ points in space defines a set of $\binom{n}{2} = \Theta(n^2)$ distinct pairs. The WSPD is a concise encoding of all these pairs that takes only $O(n)$ space. To see how this is done, let us return briefly to the $n$-body example. Suppose that we wish to determine the gravitational effect of a large number of bodies lying in a one galaxy on the planets of another. Assuming that the two galaxies are far enough away from each other relative to their respective sizes, the individual influences of the bodies in each galaxy can be aggregated into a single physical force. If there are $n_1$ and $n_2$ points in the respective galaxies, the interactions due to all $n_1 \cdot n_2$ pairs can be encoded by a single interaction pair involving the centers of the two galaxies.

To make this more precise, assume that we are given an $n$-element point set $P$ in $\mathbb{R}^d$, and a separation factor $s > 0$. We say that two disjoint sets of $A$ and $B$ are $s$-well separated if the sets $A$ and $B$ can be enclosed within two spheres of radius $r$ such that the closest distance between these spheres is at least $sr$. (This definition differs slightly from the one given by Callahan and Kosaraju but is a little easier to visualize.)

Observe that if a pair of points is $s$-well separated, it is also $s'$-well separated for all $s' < s$. Of course, since any point lies within a (degenerate) sphere of radius 0, it follows that a pair of singleton sets, $\{\{a\}, \{b\}\}$, for $a \neq b$, is well-separated for any $s$.

![Figure 18.1: A well separated pair.](image)

Well Separated Pair Decomposition: Once we see that we can aggregate pairs generated by well separated clusters in this manner, the question that remains is can we replace all the $O(n^2)$ distinct pairs of points by well separated pairs? The answer is, of course, yes (since singleton pairs are always well separated). However, what is somewhat surprising is that for any fixed $s > 0$, there exists such a decomposition consisting of only $O(n)$ pairs, where the constant factor depends inversely on $s$. This is true even if the point set is not naturally clustered.

Given arbitrary sets $A$ and $B$, define $A \otimes B$ to be the set of all distinct (unordered) pairs from these sets, that is

$$A \otimes B = \{\{a, b\} \mid a \in A, \ b \in B, \ a \neq b\}.$$
Observe that \( A \otimes A \) consists of all the \( \binom{n}{2} \) distinct pairs of \( A \). Given a point set \( P \) and separation factor \( s > 0 \), we define an \( s \)-well separated pair decomposition (\( s \)-WSPD) to be a collection of pairs of subsets of \( P \), denoted \( \{\{A_1, B_1\}, \{A_2, B_2\}, \ldots, \{A_m, B_m\}\} \), such that

1. \( A_i, B_i \subset P \), for \( 1 \leq i \leq m \)
2. \( A_i \cap B_i = \emptyset \), for \( 1 \leq i \leq m \)
3. \( \bigcup_{i=1}^{m} A_i \otimes B_i = P \otimes P \)
4. \( A_i \) and \( B_i \) are \( s \)-well separated, for \( 1 \leq i \leq m \)

The first two conditions assert that the sets \( A_i \) and \( B_i \) form disjoint pairs of \( P \). The third condition asserts that these pairs cover all the pairs of \( P \). Although these conditions alone do not imply that every pair from \( P \) arises from exactly one \( \{A_i, B_i\} \) pair, the construction that we will present later has this property. The last condition states that each pair is well separated.

An example is shown below. Take a close look at this example. Although their appears to be some sort of hierarchical structure here, note that the pairs are not properly nested within one another.

Trivially, there exists a WSPD of size \( O(n^2) \) by setting the \( \{A_i, B_i\} \) pairs to each of the distinct pair singletons of \( P \). Next we shall see that a linear-sized WSPD exists for any fixed \( s > 0 \).

**Digression: Quadtrees:** Our goal is to show that, given an \( n \)-element point set \( P \) in \( \mathbb{R}^d \) and any \( s > 0 \), there exists a \( s \)-WSPD of size \( O(n) \) (where the constant depends on \( s \) and \( d \)). Before doing this, we must make a brief digression to discuss the quadtree data structure, on which our construction is based.

The construction is based on a quadtree decomposition of the point set. A *quadtree* is a hierarchical subdivision of space into regions, called *cells*, that are hypercubes. The decomposition begins by assuming that the points of \( P \) lie within a bounding hypercube. For simplicity we may assume that \( P \) has been scaled and translated so it lies within the unit hypercube \([0,1]^d\). The initial cell, associated with the *root* of the tree, is the unit hypercube. The following process is then repeated recursively. Consider any unprocessed cell and its associated node \( u \) in the current tree. If this cell contains either 0 or 1 point of \( P \), then this is declared a leaf node of the quadtree, and the subdivision process terminates for this cell. Otherwise, the cell is subdivided into \( 2^d \) hypercubes whose side lengths are exactly half that of the original hypercube. For each of these \( 2^d \) cells we create a node of the tree, which is then made a child of \( u \) in the quadtree. The process is illustrated in the figure below, where the points and associated spatial subdivision are shown on the left and the resulting tree structure is shown on the right.
Although in practice, quadtrees as described above tend to be reasonably efficient in fairly small dimensions, there are a number of important issues in their efficient implementation in the worst case. The first is that the tree may not be of linear size. The reason is that if two points are extremely close to one another, there may be an arbitrarily long path of trivial nodes, in which only one of the $2^d$ children is an internal node. This can be fixed by compressing the quadtree. A compressed quadtree is one in which each such trivial paths is compressed into a single link. (See the figure below.) The link is given an appropriate annotation to encode the path’s structure. (If you are familiar with the concept of path compression in the PATRICIA trie data structure, the idea here is analogous.) Each internal node of the resulting tree separates at least two points into separate subtrees. Thus, there can be no more than $n - 1$ internal nodes, and hence the total number of nodes is $O(n)$.

A second issue involves the efficient computation of the quadtree. It is well known that the tree can be computed in time $O(hn)$, where $h$ is the height of the tree. However, for a compressed quadtree the tree height can be as high as $n$, which would imply an $O(n^2)$ construction time. We will not discuss it here, but it can be shown that in any fixed dimension it is possible to construct the quadtree of an $n$-element point set in $O(n \log n)$ time. (The key is handling uneven splits efficiently. Such splits arise when one child contains almost all of the points, and all the others contain only a small constant number.)

The key facts that we will use about quadtrees below are:

(a) Given an $n$-element point set $P$ in a space of fixed dimension $d$, a quadtree for $P$ of size $O(n)$ can be constructed in $O(n \log n)$ time.

(b) Each internal node has a constant number ($2^d$) children.

(c) The cell associated with each node of the quadtree is a $d$-dimensional hypercube, and as we descend from the parent to a child (in the uncompressed quadtree), the size (side length) of the cells decreases by a factor of 2.
(d) The cells associated with any level of the tree have the same size and pairwise disjoint interiors.

An important consequence stemming from (c) and (d) is the following lemma, which provides an upper bound on the number of quadtree disjoint quadtree cells of size at least $x$ that can overlap a ball of radius $r$.

**Packing Lemma:** Consider a ball $b$ of radius $r$ in any fixed dimension $d$, and consider a collection $X$ of pairwise disjoint quadtree cells of side lengths at least $x$ that overlap $b$. Then

$$|X| \leq \left(1 + \left\lceil \frac{2r}{x} \right\rceil \right)^d \leq O\left(\max\left(2, \frac{r}{x}\right)^d\right)$$

**Proof:** We may assume that all the cells of $X$ are of side length exactly equal to $x$, since making cells larger only reduces the number of overlapping cells. By the nature of a quadtree decomposition, the cells of side length $x$ form a hypercube grid $G$ of side length $x$. Consider a hypercube $H$ of side length $2r$ that encloses $b$. Clearly every cell of $X$ overlaps this hypercube. Along each dimension, the number of cells of $G$ that can overlap an interval of side length $2r$ is at most $1 + \left\lceil \frac{2r}{x} \right\rceil$. Thus, the number of grid cubes of $G$ that overlap $H$ is at most $(1 + \left\lceil \frac{2r}{x} \right\rceil)^d$. If $2r < x$, this quantity is at most $2^d$, and otherwise it is $O((r/x)^d)$.

![Figure 18.5: Proof of the Packing Lemma.](image)

For the construction of the WSPD, we need to make a small augmentation to the quadtree structure. We wish to associate each node of the tree, both leaves and internal nodes, with a point that lies within its cell (if such a point exists). Given a node $u$, we will call this point $u$’s *representative* and denote this as $\text{rep}(u)$. We do this recursively as follows. If $u$ is a leaf node that contains a point $p$, then $\text{rep}(u) = \{p\}$. If $u$ is a leaf node that contains no point, then $\text{rep}(u) = \emptyset$. Otherwise, if $u$ is an internal node, then it must have at least one child $v$ that is not an empty leaf. (If there are multiple nonempty children, we may select any one. In some applications, there may be a reason for preferring one over another, for example, we may want the representative to be the one that is nearest to the center of the cell.) Set $\text{rep}(u) = \text{rep}(v)$.

Given a node $u$ in the tree, let $P_u$ denote the points that lie within the subtree rooted at $u$. We will assume that each node $u$ is associated with its *level* in the tree, or $\text{level}(u)$. This is an integer. The root has the highest level number and levels decrease by one as we go from parent to child. (This assumes an uncompressed quadtree. In a compressed quadtree, the level increases according to the length of the path that was compressed.) The key feature of level is that $\text{level}(u) \geq \text{level}(v)$ if and only if $u$’s cell is of larger or equal size to $v$’s cell.
**Constructing a WSPD:** We now have the tools needed to to show that, given an \( n \)-element point set \( P \) in \( \mathbb{R}^d \) and any \( s > 0 \), there exists a \( s \)-WSPD of size \( s^d n \), and furthermore, this WSPD can be computed in time that is roughly proportional to its size. In particular, the construction will take \( O(n \log n + s^d n) \) time and the final WSPD can be encoded in \( O(s^d n) \) total space.

The construction operates as follows. Recall the conditions (1)–(4) given above for a WSPD. We will maintain a collection of sets that satisfy properties (1) and (3), but in general they may violate conditions (2) and (4) since they will not be disjoint and will not be well separated. When the algorithm terminates, all the pairs will be well-separated, and this will imply that they are disjoint. Each set \( \{A_i, B_i\} \) of the pair decomposition will be encoded as a pair of nodes \( \{u, v\} \) in the quadtree. Implicitly, this pair represents the pairs \( P_u \otimes P_v \), that is, the set of pairs generated from all the points descended from \( u \) and all the points descended from \( v \). This is particularly nice, because it implies that the total storage requirement is proportional to the number of pairs in the decomposition.

The algorithm is based on a recursive subdivision process. Consider a pair of node \( \{u, v\} \) that arise in the decomposition process. First, let us assume that \( u \) occurs at an equal or higher level (that is, closer to the root) than \( v \). This means that \( u \)'s cell is at least as large as \( v \)'s cell. Consider the two smallest spheres of equal radius that enclose \( u \)'s cell and \( v \)'s cell. If these spheres are well separated, then we can report \( \{u, v\} \) as (the encoding of) a well separated pair. Otherwise, we subdivide \( u \) by considering its children, and apply the procedure recursively to the pairs \( \{u_i, v\} \), for each child of \( u_i \) of \( u \).

Here is a more formal presentation of the algorithm. The procedure is called `wsPairs(u, v, T, s)`, where \( \{u, v\} \) is the current pair of nodes being decomposed, \( T \) is the compressed quadtree for the point set, and \( s \) is the separation factor. The procedure returns a set node pairs, encoding the well separated pairs of the WSPD. The initial call is `wsPairs(u0, u0, T, s)`, where \( u_0 \) is the root of the tree.

We test whether \( u \) and \( v \) are well \( s \)-separated as follows. First, we consider the smallest sphere enclosing each cell. If the node is internal, then the sphere is taken to be the sphere enclosing this cube. If it is a leaf, we take a sphere of radius 0 centered at the point associated with this leaf. Next, we expand the smaller of the two spheres so that they are of equal size. Let \( r \) be the resulting radius. Then we check whether the closest distance between these spheres (which can be computed in \( O(1) \) time) exceeds \( s \cdot r \). If so, the pair is well separated. Note that a pair of leaf cells will always pass this test (since \( r = 0 \)), so the algorithm will eventually terminate.

**Analysis:** How many pairs are generated by this recursive procedure? To evaluate this quantity, we say that a call to `wsPairs` is trivial if it does not make it to the final “else” clause. Trivial calls only take \( O(1) \)
wsPairs\((u, v, T, s)\) \{ 
  if (rep\((u)\) or rep\((v)\)) is empty) 
    return \(\emptyset\);  // no pairs to report 
  else if \((u\) and \(v\) are \(s\)-well separated) 
    return \(\{u, v\}\);   // return the WSP \(\{P_u, P_v\}\) 
  else \{  // subdivide
    if \((\text{level}(u) < \text{level}(v))\) 
      swap \(u\) and \(v\);  // \(u\)'s cell is not smaller than \(v\)'s 
    Let \(u_1, u_2, \ldots, u_m\) denote the children of \(u\) in \(T\); 
    return \(\bigcup_{i=1}^{m} \text{wsPairs}(u_i, v, T, s)\); 
  \} 
\}

Furthermore, each nontrivial call generates \(2^d\) recursive calls. Thus, the total number of calls to \(\text{wsPairs}\) is at most \(2^d\) times the number of nontrivial calls.

To count the number of nontrivial calls, we will apply a charging argument to the nodes of the compressed quadtree. Each time we make it to the final “else” clause and split the larger cell \(u\), we assign a charge to the smaller cell \(v\). We claim that the total number of charges assigned to any node \(v\) is \(O(s^d)\). Our construction only requires that \(s > 0\), but in the analysis we need to write \(\max(2, s)\) in a number of places. To simplify things, let us just assume that \(s \geq 2\) for the remainder of the analysis.

We will prove this in the context of the simpler context of the “uncompressed” quadtree. (The general is a bit more technical.) In the uncompressed quadtree, observe that side lengths of the cells of a parent at its child differ by a factor of 2. Because the algorithm always splits the cell of larger size, the sizes of cells \(u\) and \(v\) differ by at most a factor of 2.

A charge is assessed to node \(v\) only if \(u\) and \(v\) are not \(s\)-well separated. Let \(r\) denote the radius of the smallest sphere enclosing \(v\)'s cell. Because \(u\)'s cell is at least as large, it follows that the minimum distance from \(v\)'s enclosing sphere to \(u\)'s enclosing sphere must be at most \(s \cdot r\). This implies that the cell associated with \(u\) overlaps a ball of radius \(s \cdot r + c \cdot r = (s + c)r\), for an appropriate constant \(c\) depending on the dimension. Since \(r\) is the radius of the smallest ball enclosing \(v\)'s cell, \(v\)'s side length is \(x = 2r/\sqrt{d}\). Since \(u\)'s cell is at least as large, the number of charges assessed to \(v\) is at most the number of cells of the quadtree of side length \(x\) that overlap a ball of radius \((s + c)r\).

\[\text{O}(sr)\]

Figure 18.7: WSPD analysis.

\[\text{O}(sr)\]

\[\text{O}(sr)\]

3Getting the constant right is left as an exercise. The key is that the bounding ball of \(u\) overlaps the ball of radius \(s \cdot r + r\) centered at the midpoint of \(v\)'s cell, but we need to consider \(u\)'s cell itself, not its bounding ball, in order to apply the Packing Lemma. The fact that \(u\) and \(v\) have similar sizes implies that the relative distances to the bounding sphere and cell are also similar.
By the Packing Lemma, it follows that the number of such cells is at most

\[ O\left(\max\left(2, \frac{(s + c)r}{2r/\sqrt{d}}\right)^d\right) = O(s^d), \]

where the constant factors depend on the dimension (which we have assumed to be a fixed constant).

Putting this all together, we recall that there are \( O(n) \) nodes in the compressed quadtree and \( O(s^d) \) charges assigned to any node of the tree, which implies that there are a total of \( O(s^d n) \) total nontrivial calls to \texttt{wsPairs}. Finally, from the argument made earlier, the total number of calls (trivial and nontrivial) is larger by a factor of \( 2^d \). Since \( d \) is a constant, this is absorbed in the constant factor for the big-Oh notation. Together with the \( O(n \log n) \) time to build the quadtree, this gives an overall running time of \( O(n \log n) \).

Note that since each recursive call can generate at most one new well separated pair, it follows that the total number of well-separated pairs is \( O(ns^d) \), which is \( O(n) \) if \( s \) and \( d \) are assumed to be constants. Since each well separated pair is represented as a link between two nodes in a quadtree, the total size of the WSPD representation is \( O(ns^d) \). In summary we have the following result.

**Theorem:** Given a point set \( P \) in \( \mathbb{R}^d \), and a fixed separation factor \( s \geq 2 \), in \( O(n \log n + s^d n) \) time it is possible to build an \( s \)-WSPD for \( P \) consisting of \( O(s^d n) \) pairs.

As mentioned above, all this holds for \( 0 < s < 2 \) as well, but should replace \( s^d \) with \( \max(2, s^d) \) everywhere.
19 Applications of WSPDs

Reading: This material is not covered in our text.

Review: Recall that given a parameter \( s > 0 \), we say that two sets of \( A \) and \( B \) are \( s \)-well separated if the sets can be enclosed within two spheres of radius \( r \) such that the closest distance between these spheres is at least \( sr \). Given a point set \( P \) and separation factor \( s > 0 \), recall that an \( s \)-well separated pair decomposition (s-WSPD) is a collection of pairs of subsets of \( P \) \( \{\{A_1, B_1\}, \{A_2, B_2\}, \ldots, \{A_m, B_m\}\} \) such that

1. \( A_i, B_i \subset P \), for \( 1 \leq i \leq m \)
2. \( A_i \cap B_i = \emptyset \), for \( 1 \leq i \leq m \)
3. \( \bigcup_{i=1}^{m} A_i \otimes B_i = P \otimes P \)
4. \( A_i \) and \( B_i \) are \( s \)-well separated, for \( 1 \leq i \leq m \)

where \( A \otimes B \) denotes the set of all unordered from \( A \) and \( B \).

Last time we showed that, given \( s \geq 2 \), there exists an \( s \)-WSPD of size \( O(s^d n) \), which can be constructed in time \( O(n \log n + s^d n) \). (The algorithm works for any \( s > 0 \), and the \( s^d \) term is more accurately stated as \( \max(2, s)^d \).) The WSPD is represented as a set of unordered pairs of nodes of a quadtree decomposition of \( P \).

Today we discuss a number of applications of WSPDs.

Approximating the Diameter: Recall that the diameter of a point set is defined to be the maximum distance between any pair of points of the set. The diameter can be computed exactly by brute force in \( O(n^2) \) time. For points in the plane, it is possible to compute the diameter in \( O(h) \) time, where \( h \) is the number of points on the convex hull. Recall that the convex hull can be computed in \( O(n \log n) \) time. This is done by a method called rotating calipers, which involves simulating the sweeping of two parallel supporting lines around the boundary of the hull, and determining the farthest that these lines come from one another. However, in higher dimensions, this approach leads to an \( O(h^2) \) running time in the worst case, which is just as bad as brute force if \( h = \Omega(n) \).

Using the WSPD construction, we can easily compute an \( \varepsilon \)-approximation to the diameter of a point set \( P \) in linear time. Given \( \varepsilon \), we let \( s = 4/\varepsilon \), and construct an \( s \)-WSPD. Recall that in our construction of the WSPD of a point set \( P \), every half of each well-separated pair \( (A_i, B_i) \) using the notation above consists of the set of points that are descended from some node in the compressed quadtree of \( P \). Also recall that as part of the quadtree construction, each node \( u \) of the quadtree can be associated with a representative point, denoted \( \text{rep}(u) \), which was chosen the be an arbitrary descendant point of \( u \). For each well separated pair derived from a pair of nodes \( u \) and \( v \), let \( P_u \) and \( P_v \) denote the corresponding points. For every well separated pair \( \{P_u, P_v\} \), we compute the distance between \( \text{rep}(u) \) and \( \text{rep}(v) \), and return the largest such distance.

To prove correctness, let \( x \) and \( y \) be the points of \( P \) that realize the maximum distance. Let \( \{P_u, P_v\} \) be the well separated pair containing these points, and \( p_u \) and \( p_v \) denote their respective representatives. By definition of well separatedness, we know that \( P_u \) and \( P_v \) can be enclosed in balls of radius \( r \) that are separated by distance at least \( s \cdot r \). (See the figure below.) Therefore, by the triangle inequality we have

\[
\|xy\| \leq \|p_u p_v\| + 2r + 2r = \|p_u p_v\| + 4r.
\]
Also, by the WSPD separation properties, we have $\|p_u p_v\| \geq s \cdot r$ implying that $r \leq \|p_u p_v\| / s$. Combining these we have

$$\|xy\| \leq \|p_u p_v\| + 4r \leq \|p_u p_v\| + \frac{4}{s}\|p_u p_v\|$$

$$= \left(1 + \frac{4}{s}\right)\|p_u p_v\|$$

$$= (1 + \varepsilon)\|p_u p_v\|,$$

Clearly $\|p_u p_v\| \leq \|xy\|$, and therefore we have

$$\frac{\|xy\|}{1 + \varepsilon} \leq \|p_u p_v\| \leq \|xy\|,$$

which implies that the output is an $\varepsilon$-approximation. The running time is dominated by the size of the WSPD, which is $O(sd n) = O(n)$, assuming that $\varepsilon$ and $d$ are constants.

**Closest Pair:** The same sort of approach could be used to produce an $\varepsilon$-approximation to the closest pair as well, but surprisingly, there is a much better solution. If we were to generalize the above algorithm, we would first compute an $s$-WSPD for an appropriate value of $s$, and for each well separated pair $\{P_u, P_v\}$ we would compute the distance $\|\text{rep}(u) \text{rep}(v)\|$, and return the smallest such distance. As before, we would like to argue that (assuming $s$ is chosen properly) this will yield an approximation to the closest pair. It is rather surprising to note that if $s$ is chosen properly, this approach yields the true closest pair, not an approximation.

To see why, consider a point set $P$, let $x$ and $y$ be the closest pair of points and let $p_u$ and $p_v$ be the representatives from their associated well separated pair. If it were the case that $x = p_u$ and $y = p_v$, then the representative-based distance would be exact. Suppose therefore that either $x \neq p_u$ or $y \neq p_v$. But wait. If the separation factor is high enough, this would imply that either $\|xp_u\| < \|xy\|$ or $\|yp_v\| < \|xy\|$, either of which contradicts the fact that $x$ and $y$ are the closest pair.

To make this more formal, let us assume that $\{x, y\}$ is the closest pair and that $s > 2$. We know that $P_u$ and $P_v$ lie within balls of radius $r$ that are separated by a distance of at least $s \cdot r > 2r$. If $p_u \neq x$, then we have

$$\|p_u x\| \leq 2r < s \cdot r \leq \|xy\|,$$

yielding a contradiction. Therefore $p_u = \text{rep}(u) = x$. By a symmetrical argument $p_v = \text{rep}(v) = y$. Since the representative was chosen arbitrarily, it follows that the $P_u = \{x\}$ and $P_v = \{y\}$. Therefore, the closest representatives are in fact, the closest pair.

Since $s$ can be chosen to be arbitrarily close to 2, the running time is $O(n \log n + 2^d n) = O(n \log n)$, since we assume that $d$ is a constant. Although this is not a real improvement over our existing closest-pair algorithm, it is interesting to note that there is yet another way to solve this problem.
Spanner Graphs: Recall that a set $P$ of $n$ points in $\mathbb{R}^d$ defines a complete weighted graph, called the Euclidean graph, in which each point is a vertex, and every pair of vertices is connected by an edge whose weight is the Euclidean distance between these points. This graph is dense, meaning that it has $\Theta(n^2)$ edges. It would be nice to have a sparse graph having only $O(n)$ edges that approximates the Euclidean graph in some sense.

One such notion is to approximate the distances (length of the shortest path) between all pairs of vertices. A subgraph of a graph that approximates all shortest paths is called a spanner. In the geometric context, suppose that we are given a set $P$ and a parameter $t \geq 1$, called the stretch factor. We define a $t$-spanner to be a weighted graph $G$ whose vertex set is $P$ and, given any pair of points $x, y \in P$ we have
\[
\|xy\| \leq \delta_G(x, y) \leq t \cdot \|xy\|,
\]
where $\delta_G(x, y)$ denotes the length of the shortest path between $x$ and $y$ in $G$.

WSPD-based Spanner Construction: Do sparse geometric spanners exist? We have actually already seen one. It can be proved that the planar Delaunay triangulation is a $t$-spanner, for some $t$, where $2 \leq t \leq 2.43$. The tightest value of $t$ is not known. (The proof is a bit involved, and we will not present it.)

There is a straightforward method for computing a geometric spanner from a WSPD on the point set. This seems intuitively reasonable. Consider each well-separated pair $\{A_i, B_i\}$. We should only need to create a single edge joining some point $a \in A_i$ to some point $b \in B_i$. Which points? It turns out that it does not matter.

Here is the algorithm in greater detail. Given $P$ and $t > 1$, we begin by computing a WSPD for an appropriate separation factor $s$ depending on $t$ (to be derived later) and the associated quadtree. Recall that each well-separated pair is associated with a pair $\{u, v\}$ from the compressed quadtree of $P$, and these are in turn associated with representative points, $\text{rep}(u)$ and $\text{rep}(v)$, respectively. We create an edge $\{\text{rep}(u), \text{rep}(v)\}$ and add it to the spanner, and assign its weight to be the distance between these points. Let $G$ be the resulting undirected weighted graph. (See the figure below.) Clearly the number of edges of $G$ is equal to the number of well-separated pairs, which is $O(s^d n)$, and can be built in the same $O(n \log n + s^d n)$ running time as the WSPD construction.

![Figure 19.2: A WSPD and its associated spanner.](image)

Correctness: To establish the correctness of this algorithm, it suffices to show that for all pairs $x, y \in P$, we have
\[
\|xy\| \leq \delta_G(x, y) \leq t \cdot \|xy\|.
\]
Clearly, the first inequality holds because we never create an edge between two vertices whose weight is less than the Euclidean distance between them, and by the triangle inequality, an indirect path between two points can never be shorter than the direct path.

To prove the second inequality, we apply an induction based on the number of edges of the shortest path in the spanner. (Actually, we will consider not true shortest path, but paths of a particular structure. This is sufficient because, if we can show that there exists a path satisfying the spanner bound, then clearly the shortest path will as well.) As a basis, if the two points are joined by an edge, then clearly \( \delta_G(x, y) = \|xy\| < t \cdot \|xy\| \) for all \( t > 1 \).

If, on the other hand, there is no direct edge between \( x \) and \( y \), we know that \( x \) and \( y \) must lie in some well-separated pair \( \{P_u, P_v\} \) defined by the pair of nodes \( \{u, v\} \) in the quadtree. let \( p_u = \text{rep}(u) \) and \( p_v = \text{rep}(v) \) be the respective representative representative. (It might be that \( p_u = x \) or \( p_v = y \), but not both.) Let us consider the length of the path from \( x \) to \( p_u \) to \( p_v \) to \( y \). Since the edge \( \{p_u, p_v\} \) is in the graph, we have

\[
\delta_G(x, y) \leq \delta_G(x, p_u) + \delta_G(p_u, p_v) + \delta_G(p_v, y) \leq \delta_G(x, p_u) + \|p_u p_v\| + \delta_G(p_v, y).
\]

(See the figure below.)

![Figure 19.3: Proof of the spanner bound.](image)

Since the paths from \( x \) to \( p_u \) and \( p_v \) to \( y \) are subpaths, and hence shorter than the overall path, we may apply the induction hypothesis, which yields \( \delta_G(x, p_u) \leq t \|xp_u\| \) and \( \delta_G(p_v, y) \leq t \|pv\| \), yielding

\[
\delta_G(x, y) \leq t(\|xp_u\| + \|pv\|) + \|p_u p_v\|.
\]

(1)

Let \( s \) denote the (as yet, unspecified) separation factor for the WSPD. Since \( P_u \) and \( P_v \) are \( s \)-well separated, we know that each of these point sets can be enclosed within a ball of radius \( r \) such that the two balls are separated by distance at least \( s \cdot r \). Thus, we have \( \max(\|xp_u\|, \|pv\|) \leq 2r \), and \( \|xy\| \geq sr \). From the second inequality we have \( r \leq \|xy\| / s \). By the triangle inequality, we have

\[
\|p_u p_v\| \leq \|p_u x\| + \|xy\| + \|yp_v\| \leq 2r + \|xy\| + 2r \leq 4r + \|xy\|.
\]

Combining these observations with Eq. (1) we obtain

\[
\delta_G(x, y) \leq t(2r + 2r) + (4r + \|xy\|) \leq 4r(t + 1) + \|xy\|.
\]

From the fact that \( r \leq \|xy\| / s \) we have

\[
\delta_G(x, y) \leq \frac{4(t + 1)}{s} \|xy\| + \|xy\| \leq \left(1 + \frac{4(t + 1)}{s}\right) \|xy\|.
\]
To complete the proof, all we need is to select \( s \) so that \( 1 + 4(t + 1)/s \leq t \). We easily see that this is true if \( s \) is chosen so that
\[
s = \frac{4t + 1}{t - 1}.
\]
Since we assume that \( t > 1 \), this is possible for any \( t \).

The number of edges in the spanner is \( s^d n \). We assume that \( t > 1 \) and spanners are most interesting with small stretch factors, say \( t \leq 2 \). If we express \( t \) as \( t = 1 + \varepsilon \), then we see that the size of the spanner is \( O(s^d n) = O((1/\varepsilon)^d n) \). In conclusion, we have the following theorem:

**Theorem:** Given a point set \( P \) in \( \mathbb{R}^d \), a \((1 + \varepsilon)\)-spanner for \( P \) containing \( O((1/\varepsilon)^d n) \) edges can be computed in \( O(n \log n + (1/\varepsilon)^d n) \) time.

**Approximating the Euclidean MST:** We will now show that with the above spanner result, we can compute an \( \varepsilon \)-approximation to the minimum spanning tree. Suppose we are given a set \( P \) of \( n \) points in \( \mathbb{R}^d \), and we wish to compute the Euclidean minimum spanning tree (MST) of \( P \). Given a graph with \( v \) vertices and \( e \) edges, it is well known that the MST can be computed in time \( O(e + v \log v) \). It follows that we can compute the MST of a set of points in any dimension by first constructing the Euclidean graph and then computing its MST, which takes \( O(n^2) \) time. To compute the approximation to the MST, we first construct a \((1 + \varepsilon)\)-spanner, call it \( G \), and then compute and return the MST of \( G \). (See the figure below.) This approach has an overall running time of \( O(n \log n + s^d n) \).

![Figure 19.4: Approximating the Euclidean MST.](image)

To see why this works, let \( T \) denote the edges of the Euclidean minimum spanning tree, and \( w(T) \) denote the total weight of its edges. For each edge \( \{x, y\} \in T \), let \( \pi(x, y) \) denote the shortest path (as a set of edges) between \( x \) and \( y \) in \( G \). The shortest path from \( x \) to \( y \) in the Euclidean graph is the edge between them, whose length is \( \|xy\| \). Since \( G \) is a spanner, we know that \( w(\pi(x, y)) \leq (1 + \varepsilon)\|xy\| \).

Now, consider the subgraph \( G' \subseteq G \) formed by taking the union of all the edges of \( \pi(x, y) \) for all \( \{x, y\} \in T \). That is, \( G \) and \( G' \) have the same vertices, but each edge of the MST is replaced by its spanner path. Clearly, \( G' \) is connected. We can bound the weight of \( G' \) in terms of the weight of the MST:
\[
\begin{align*}
  w(G') &= \sum_{\{x, y\} \in T} w(\pi(x, y)) \\
         &\leq \sum_{\{x, y\} \in T} (1 + \varepsilon)\|xy\| \\
         &= (1 + \varepsilon) \sum_{\{x, y\} \in T} \|xy\| \\
         &= (1 + \varepsilon) w(T).
\end{align*}
\]
However, because $G$ and $G'$ share the same vertices, and $G$’s edges are a superset of the edges of $G'$, it follows that the MST of $G$ is free to use any edge that $G'$ can, and so the $w(MST(G)) \leq w(MST(G'))$. Combining everything we have

\[ w(MST(G)) \leq w(MST(G')) \leq w(G') \leq (1 + \varepsilon)w(T), \]

yielding the desired approximation bound.
20  Range Spaces, Nets, and Approximations


Range Counting and Approximation: Today we consider a number of interesting issues that naturally arise from the study of range searching. In range searching, we are given a set $P$ of points in space, and we are interested in subsets formed by the intersection of $P$ with shapes, called ranges, that have been drawn from some class of shapes (e.g., axis-aligned rectangles, halfspaces, simplices, circular discs, ellipsoids). Range searching is but one example of a rich collection of computational and mathematical problems in which these concepts apply. For example, we have seen that the same concepts are very useful in the study of discrepancy. Today, we will see examples of how these concepts can be applied to geometric sampling and approximation. Our focus will be on mathematical issues, but there are numerous applications of these concepts.

This topic is also close to the core of two important topics in computational geometry, randomized algorithms and approximation algorithms. In the last couple of lectures we introduced the notion of a well-separated pair decompositions (WSPD). For WSPDs we are approximating Euclidean distances. Some problems, however, the natural notion of approximation involves counts, not distances. For example, in a range counting query, we might be quite happy with a data structure that returns an approximation to the total number of points appearing within the query range. (That is, the query range contains 45% of the points, plus or minus a 2% error.)

It is not hard to see that the answer to this question is closely related to randomized algorithms. For example, one approach would be to randomly sample of a relatively small number of points, and then build an exact range counting structure for this smaller set. But how large should this sample be? A natural question along these lines might be, given a range counting problem involving a set $P$ of $n$-points in $\mathbb{R}^d$, and a user-supplied approximation parameter $\varepsilon$, how large a random sample do we expect to need if we want to be able to approximate the fraction of points lying within a given range to within error $\varepsilon$? How does the size of this fraction depend on $n$? on the dimension $d$? on the sorts of ranges in our range space? Today, we will discuss an elegant mathematical theory for answering these sorts of problems.

Range Spaces: In order to provide a formal mathematical framework from which to base our analysis, we define a *range space* (also called a *set system*) to be a pair $\Sigma = (X, R)$, where $X$ is a (finite or infinite) set, called *points*, and $R$ is a (finite or infinite) family of subsets of $X$, called *ranges*. We will consider the problem in a geometric setting, where the elements of $X$ are points in $\mathbb{R}^d$ and each element of $R$ is the intersection of $X$ with some geometric shape from some class of shapes (axis-aligned rectangles, halfspaces, etc.)

We have allowed $X$ to be infinite (e.g., the set of all points in the plane) but when we apply this theory in the context of a computational problem, we will be considering just a finite set of points (e.g., the $(x, y)$ coordinates of all Starbucks’s coffee shops in the United States). Given a range space $(X, R)$, for any $Y \subseteq X$ (where $Y$ will be our finite set), define $P_R(Y)$, called the *projection* (or *restriction*) of $R$ on $Y$, to be $\{r \cap Y \mid r \in R\}$.

VC-dimension: In typical geometrical contexts, the sorts of ranges $R$ that we are interested in will involve simple shapes of constant description complexity. How can we characterize this concept formally?
An elegant method for doing this was developed by Vladimir Vapnik and Alexey Chervonenkis in the late 60’s and early 70’s in the general area that came to be known as statistical learning theory. This went on become a fundamental concept in many different areas of mathematics and computer science ranging from pattern recognition, classification and clustering, and information retrieval. This work was applied in the context of computational geometry and range searching by Haussler and Welzl in the late 80’s, where it made a huge impact.

The idea that Vapnik and Chervonenkis used as a basis for characterizing range spaces involves the question of whether it is possible to form all possible subsets of a given set by intersecting it with ranges, and for how large a set is this possible. To make this more formal, we say that a set $Y$ is shattered by $R$ if $P_R(Y)$ contains all the subsets of $Y$. (If $Y$ is finite, this is equivalent to saying that $|P_R(Y)| = 2^{|Y|}$.) The Vapnik-Chervonenkis dimension (or VC-dimension) of $\Sigma$, denoted $VC(\Sigma)$, is the maximum cardinality of a shattered subset of $X$. If there are arbitrarily large shattered subsets then the VC-dimension is defined to be $\infty$.

The figure below shows that if the range space consists of halfplanes, it is possible to shatter a set of three points, and therefore the VC-dimension of the range space defined by halfspaces is at least 3. (See Fig. 20.1(a).) On the other hand, observe that no 4-element set in the plane can be shattered by halfplane queries. This either because one of the points lies within the convex hull of the other three, implying that we cannot realize the subset consisting of this single point or else all four points are on the convex hull in which case it is not possible to generate a subset consisting of points that are not consecutive on the hull. (See Fig. 20.1(b).) Therefore, the VC-dimension of the range space of halfplanes in the plane is 3.

![Figure 20.1: (a) There exists a 3-element set that is shattered by halfplane ranges, but (b) no 4-element set is shattered by halfplane ranges.](image)

Note that the definition of VC-dimension does not rely on the fact that the ranges are defined by geometric shapes. It is defined in purely set-theoretic terms. This is one of the reasons that the theory is widely applicable.

**Range Spaces of Finite VC-Dimension:** An immediate consequence of a range space having finite VC-dimension is that it implies a bound on the size of the range space as a function of the number of points. The fact that a 4-element set cannot be shattered by halfplanes follows from a fundamental result from the theory of convex sets called Radon’s Theorem. It states that any set of $d + 2$ points in $\mathbb{R}^d$ can be partitioned into two subsets whose convex hulls intersect either other. No halfplane can separate these two subsets.
points. The result is sometimes called **Sauer’s Lemma** (although it was proved in slightly different forms by a number of people including Vapnik and Chervonenkis). Before stating this lemma we give a definition. For nonnegative integers \( n \) and \( d \) define

\[
g(d, n) = \sum_{i=0}^{d} \binom{n}{i}.
\]

This is just the total number of subsets of size \( d \) or smaller taken from a universe of size \( n \). It is easy to see that \( g(d, n) = O(n^d) \). From basic properties of binomial coefficients, for \( n \geq 1 \) and \( d \geq 1 \), it can be shown that \( g(d, n) = g(d, n-1) + g(d-1, n-1) \).\(^5\)

**Lemma:** (Sauer’s Lemma) If \((X, R)\) is a range space of VC-dimension \( d \) with \(|X| = n\) points, then \(|R| \leq g(d, n)\).

**Proof:** The proof is by induction on \( n \) and \( d \). The basis case \((n = d = 0)\) is trivial. For the induction step let us assume that the assertion holds for \((d, n-1)\) and \((d-1, n-1)\) and we prove it for the pair \((d, n)\).

The general strategy will be to eliminate an arbitrary point \( x \in X \). We classify the sets of \( R \) into two groups: (a) those that arise as nearly identical twins, one containing \( x \) and one that does not, and (b) those that do not have such a near twin. To formalize this, define

\[
R_x = \{r \setminus \{x\} \mid x \notin r, r \cup \{x\} \in R\}.
\]

This is bit tricky to parse at first, but observe that it consists of the ranges of \( R \) that do not contain \( x \) but there exists another range of \( R \) that is identical except it does contain \( x \). Call these sets **twins**. Since every element of \( R_x \) corresponds to a pair of ranges of \( R \), one that contains \( x \) and one that does not, it follows that the VC-dimension of \( R_x \) is \( d - 1 \). (Observe that if \( R_x \) can shatter any set \( X' \), then the original range space can shatter \( X' \cup \{x\} \).) Since \( R_x \) does not contain the element \( x \), it follows that \(|R_x| \leq g(d-1, n-1)\). The remaining sets of \( R \), are called **singles**. Consider the ranges that result by simply eliminating \( x \):

\[
R \setminus x = \{r \setminus \{x\} \mid x \in R\}.
\]

Clearly \( R \setminus x \) has one fewer point, but we cannot say that it has a smaller a VC-dimension than \( d \). (Of course, it is certainly not larger!) Therefore we have \(|R \setminus x| \leq g(d, n-1)\).

Observe that the singles are counted exactly once in \( R \setminus x \). For each twin, the one that contains \( x \) is counted in \( R \setminus x \) and the one that does not is counted in \( R_x \). (See Fig. 20.2 for an example.) Thus between these two sets we count all the elements of \( R \). In summary we have

\[
|R| = |R \setminus x| + |R_x| \leq g(d, n-1) + g(d-1, n-1) = g(d, n),
\]

where the last step follows from our previous observation about \( g(d, n) \).

\(^5\)You may remember this famous recurrence if you took a course in discrete math. It has a natural interpretation in terms of generating all subsets of size \( d \) from a universe of size \( n \). Consider any element \( x \). If we decide to include \( x \) in our subset, then there \( g(d-1, n-1) \) ways to choose the remaining \( d-1 \) elements from the remaining \( n-1 \) elements of the universe. If we decide not to include \( x \), then there are \( g(d, n-1) \) ways to select a \( d \) element subset from the remaining \( n-1 \) elements of the universe.
Note that there is some slack here. This theorem predicts that the range space of halfplanes contains $g(3, n) = O(n^3)$ elements. In fact, the number of distinct ranges is $\Theta(n^2)$, since there are essentially $\Theta(n^2)$ canonical lines determined by all the pairs of points of the set. But we are not too far off. For example, if we had altered the definition slightly, and considered instead the range space consisting all the lower halfplanes (the points lying below a nonvertical line), then it can be shown that its VC-dimension is only 2, and the size of resulting range space is $\Theta(n^2)$.

**ε-Nets and ε-Approximations:** One of the most useful aspects of having bounded VC-dimension is that it implies that the range space can be well approximated by a small well chosen set of sample points. What do we mean by “well chosen?” Consider a range space $(X, R)$, and let $Y$ be any finite subset of $X$. Let $Z \subseteq Y$ be a subset of $Y$, which will think of as our sample. (For the purposes of sampling, we expect that $|Z| \ll |Y|$.) In order for the sample to be well chosen, we would like it to be the case that for any range $r \in R$, we expect that the fraction of points of $Y$ that lie within $r$ should be similar to the fraction of points of $Z$ that lie within $r$.

To capture this idea more formally, for any $0 < \varepsilon < 1$, we say that a subset $Z \subseteq Y$ is an **ε-approximation** for $Y$ (also called an **ε-sample**) if for any $r \in R$,

$$\left| \frac{|Y \cap r|}{|Y|} - \frac{|Z \cap r|}{|Z|} \right| \leq \varepsilon.$$ 

That is, the fractions of points of the original and subset covered by the range differ only by $\varepsilon$.

Often a weaker concept is sufficient. Rather than requiring that the fractions of covered points be similar, we impose the more limited condition that if the range covers at least an $\varepsilon$ fraction $Y$, then at least one point of the sample should be in the range. To formalize this, we say that a subset $Z \subseteq Y$ is an **ε-net** for $Y$ if for any $r \in R$,

$$\frac{|Y \cap r|}{|Y|} \geq \varepsilon \Rightarrow Z \cap r \neq \emptyset.$$ 

Figure 20.2: An illustration of the proof of Sauer’s Lemma.
Computational Geometry Notes  
Nets and Approximation

(In Fig. 20.3(b) the black points form a 0.1-net for a set of points Y for halfplane ranges.) It is easy to see that if Z is an \( \varepsilon \)-approximation, it is also an \( \varepsilon \)-net, but the converse need not hold. (You should verify this yourself.)

![Figure 20.3: (a) A point set Y, (b) an \( \varepsilon \)-net Z (in black) for halfplane ranges, and (c) a counterexample that small \( \varepsilon \)-nets do not exist for ranges of unbounded VC-dimension.](image)

**The Size of \( \varepsilon \)-Nets and \( \varepsilon \)-Approximations:** You might be tempted to think that any sufficiently large random subset will very likely be a \( \varepsilon \)-approximation (and hence an \( \varepsilon \)-net). We might reason as follows. Given any fixed range that contains, say, 25% of the points of Y, the probability that a random subset of points of Y lies within the range is 0.25, and so the expected percentage of random subset in the range is also 25%. (Note that this holds irrespective of the range’s shape). Although this reasoning is correct, it applies to just a single range. In order for a set to be an \( \varepsilon \)-net or \( \varepsilon \)-approximation, the defining condition must hold for all ranges, not just one range or most ranges. For example, if you allow for arbitrarily complex ranges, it is easy to see that any relatively small random subset will fail to be an \( \varepsilon \)-net or \( \varepsilon \)-approximation. (See Fig. 20.3(c), for example.) Nonetheless, this intuition is correct for spaces of finite VC-dimension (although the proof is not quite this simple!) We will show that a sufficiently large random subset will likely be an \( \varepsilon \)-approximation with high probability, where the size of the subset depends on the VC-dimension and the desired probability of success. Remarkably, the number of random points does not depend on the size of the original point set. In other words, it does not matter whether you are sampling from the students of this University or the inhabitants of China—the same number of random points works for both!

The next two theorems establish these facts formally. The first shows that a sufficiently large random subset is likely to be an \( \varepsilon \)-approximation and was proved by Vapnik and Chervonenkis. The second does this for \( \varepsilon \)-nets, and was proved by Haussler and Welzl. Ignoring the relative small log factors, the bottom line is that we need roughly \( O(1/\varepsilon^2) \) random points to form an \( \varepsilon \)-approximation and \( O(1/\varepsilon) \) random points to form an \( \varepsilon \)-net. Henceforth, let log denote the natural logarithm (usually abbreviated ln).

**Theorem 1:** There is a positive constant \( c \) such that if \((X, R)\) is any range space of VC-dimension at most \( d \), and \( Y \) is any finite subset of \( X \) and \( \varepsilon, \delta > 0 \), then a random subset \( Z \subseteq Y \) of cardinality \( m \) where,

\[
m \geq \frac{c}{\varepsilon^2} \left( d \log \frac{d}{\varepsilon} + \log \frac{1}{\delta} \right)
\]

is an \( \varepsilon \)-approximation for \( Y \) with probability at least \( 1 - \delta \). (If \( m \) exceeds \( |Y| \), then just let \( Z = Y \).)

**Theorem 2:** Let \((X, R)\) be any range space of VC-dimension at most \( d \), let \( Y \) be any finite subset of \( X \), and let \( 0 < \varepsilon, \delta < 1 \). Let \( Z \subseteq Y \) be a set obtained by \( m \) random independent draws from \( Y \)
where
\[ m \geq \max \left( \frac{4}{\varepsilon} \log \frac{2}{\delta}, \frac{8d \log 8d}{\varepsilon} \right). \]

Then \( Z \) is an \( \varepsilon \)-net for \( Y \) with probability at least \( 1 - \delta \).

You might have noticed that the phrasing of the two theorems is slightly different. (For example, one involves a random subset of a given size and the other is based on making a series of random draws.) I believe that these are just minor technical issues arising from the fact that the theorems come from different sources.

It is remarkable to note that there is no exponential dependence on \( d \). (That is, unlike \( g(d, n) \) the size bounds on the \( \varepsilon \)-net and \( \varepsilon \)-approximation do not contain \( d \) in the exponent.) Are the messy log factors necessary, or are they just an artifact of some slack in the analysis? The answer is not completely known, but there are (rather contrived) counterexamples that show that the sample size cannot be strictly linear in \( 1/\varepsilon \).

These theorems state that a random sample is likely to be an \( \varepsilon \)-approximation (or \( \varepsilon \)-net), but does not provide a computational guarantee of this. There are both randomized and deterministic algorithms whose output is guaranteed, but they are quite a bit more complex, either in running time or in structure. If you don’t mind living dangerously, you could just compute a sufficiently large random sample and “trust” that it works.

The two theorems are proved by essentially the same methods. We will present a sketch of the proof of Theorem 2 only, since it is the simpler of the two.

**Proof of Theorem 2:** Let \( (X, R) \) be a range space of VC-dimension \( d \), let \( Y \) be any finite subset of \( X \) of size \( n \), and let \( 0 < \varepsilon, \delta < 1 \). Let \( Z = (z_1, z_2, \ldots, z_m) \) be a sequence of \( m \) elements of \( Y \) sampled randomly and independently, where \( m \) satisfies the bound given in the statement of the theorem. (Note that the sampled elements of \( Z \) need not be distinct, which is why we present them as a sequence.) Let \( E_1 \) be the event that \( Z \) fails to be an \( \varepsilon \)-net. That is,
\[ E_1 = \{ \exists r \in R : |r \cap Y| \geq \varepsilon n \text{ and } r \cap Z = \emptyset \}. \]

Intuitively, since \( r \) covers at least \( \varepsilon n \) of the elements \( Y \), we would expect it to cover at least \( \varepsilon m \geq 4 \log(2/\delta) \) elements of any random sample of size \( m \). As \( \delta \) gets smaller our confidence that at least one point is covered will grow. It suffices to show that, given the choice of \( m \), the probability of \( E_1 \) is at most \( \delta \).

Unfortunately, \( E_1 \) depends on \( Y \), and we do not want any dependence on \( n \) in our bound. To this end, we would like to express probability in terms of a quantity that does not depend on the size of \( Y \). To do this, consider another random sample \( W = (w_1, w_2, \ldots, w_m) \), also of size \( m \). Let \( E_2 \) denote the event that \( Z \) fails to be an \( \varepsilon \)-net for some range \( r \), but \( W \) succeeds in netting at least half the number of elements we would expect it to.
\[ E_2 = \left\{ \exists r \in R : |r \cap Y| \geq \varepsilon n, r \cap Z = \emptyset, \text{ and } |r \cap W| \geq \frac{\varepsilon m}{2} \right\}. \]

(Since \( Z \) and \( W \) may contain repetitions, the notation \( |r \cap W| \) in this context is taken to mean that we multiply count duplicated elements.) Observe that since \( E_2 \subseteq E_1 \), its probability is not greater than \( E_1 \). We show next that the probability of \( E_1 \) is not greater than twice that of \( E_2 \).
**Claim:** \( \Pr(E_1) \leq 2 \Pr(E_2) \).

**Proof:** Consider the conditional probability of \( E_2 \) occurring, given that \( E_1 \) occurs. This is denoted \( \Pr(E_2 \mid E_1) \). From standard definitions from probability theory and the fact that \( E_2 \subseteq E_1 \), we have

\[
\Pr(E_2 \mid E_1) = \frac{\Pr(E_2 \cap E_1)}{\Pr(E_1)} = \frac{\Pr(E_2)}{\Pr(E_1)}.
\]

Thus, it suffices to show that this conditional probability is at least 1/2. Each sample of \( W \) has a probability of at least \( \varepsilon \) of lying within \( r \). Since the samples are independent, it follows from standard results in probability theory that the number of points of \( W \) that lie within \( r \) is a random variable from a binomial distribution whose mean value is \( \mu = \varepsilon m \) and whose standard deviation is \( \sigma = \sqrt{m \varepsilon (1 - \varepsilon)} \leq \sqrt{\varepsilon m} \). A useful result at this point from probability theory is Chebyshev’s inequality, which states that the probability that a random variable is more than \( t \) standard deviations from its mean is at most \( 1/t^2 \). Using this we obtain

\[
1 - \Pr(E_2 \mid E_1) = \Pr\left( |r \cap W| < \frac{\varepsilon m}{2} \right) \leq \Pr\left( |r \cap W| - \varepsilon m > \frac{\varepsilon m}{2} \right)
\]

\[
= \Pr\left( |r \cap W| - \mu > \sigma \sqrt{\varepsilon m} \right) \leq \frac{4}{\varepsilon m} \quad \text{(by Chebyshev)}
\]

\[
\leq \frac{1}{2}.
\]

Therefore, \( \Pr(E_2 \mid E_1) \geq 1/2 \), which completes the proof.

To complete the proof, it suffices to show that \( \Pr(E_2) \leq \delta/2 \). To do this, consider first the following event

\[ E_3 = \left\{ \exists r \in R : r \cap Z = \emptyset \text{ and } |r \cap W| \geq \frac{\varepsilon m}{2} \right\} . \]

Clearly \( E_2 \subseteq E_3 \). Thus, bounding the probability of \( E_3 \) provides a bound on the probability of \( E_2 \), which is our goal. Note, however, that \( E_3 \) does not depend on \( Y \). In other words, if we can succeed in bounding \( E_3 \), we will have achieved our desired goal of eliminating any dependence on \( Y \) and \( n \).

(This is a crucial element of the proof.)

Next, we provide a bound on the probability of \( E_3 \). Here is a quick overview. Recall that the quantity \( g(d, n) \leq n^d \) introduced earlier on the size of a range space of VC-dimension \( d \). We have eliminated \( Y \) and are only left with \( Z \cup W \), whose combined size is \( 2m \). If we restrict ourselves to this range space of \( 2m \) points, there are roughly \( g(d, 2m) = O(m^d) \) distinct ranges to be considered. Assuming that the VC-dimension is bounded, this quantity grows polynomially in \( m \). We shall show below that the probability of something going wrong on any given range decreases exponentially with \( m \). Thus, as \( m \) increases, the exponential term quickly dominates to control the overall probability of a bad event. (This is another key element of the proof.) Indeed, the rather arcane choice for the value of \( m \) given in the statement is engineered exactly to obtain this tradeoff value.

**Claim:** \( \Pr(E_2) \leq \Pr(E_3) \leq g(d, 2m) \cdot 2^{-\varepsilon m/2} \).

**Proof:** As mentioned above \( E_2 \subseteq E_3 \), and therefore it suffices to prove the upper bound on \( \Pr(E_3) \).

We can imagine that we sample the elements of \( Z \cup W \) by first sampling \( 2m \) elements \( U = (u_1, u_2, \ldots, u_{2m}) \) independently from \( Y \) and then randomly deciding which \( m \) elements of \( U \)
Proof: We will assume that $\Pr(E_1) \leq \delta$. By combining the previous two claims, we have

$$\Pr(E_1) \leq 2\Pr(E_2) \leq 2\Pr(E_3) \leq 2g(d, 2m) \cdot 2^{-\varepsilon m/2}.$$ 

Therefore, all that remains to complete the proof of Theorem 2 is to show that our choice of $m$ guarantees that $2g(d, 2m) \cdot 2^{-\varepsilon m/2} \leq \delta$.

Claim: Let $m \geq \max((4/\varepsilon) \log(2/\delta), (8d/\varepsilon) \log(8d/\varepsilon))$. Then $2g(d, 2m) \cdot 2^{-\varepsilon m/2} \leq \delta$.

Proof: We will assume that $d \geq 2$. The case $d = 1$ is simpler. Recall that $g(d, 2m) \leq (2m)^d$. It suffices to show that

$$2g(d, 2m) \leq \delta 2^{\varepsilon m/2}.$$
Taking logs of both sides, we see that this is equivalent to showing that

\[ \frac{\varepsilon m}{2} \geq d \log(2m) + \log \frac{2}{\delta}. \]

This will follow by showing that our choice of \( m \) guarantees that

\[ \frac{\varepsilon m}{4} \geq \log \frac{2}{\delta} \quad \text{and} \quad \frac{\varepsilon m}{4} \geq d \log 2m. \]

The first inequality follows immediately from the fact that \( m \geq (4/\varepsilon) \log(2/\delta) \). To prove the second inequality, observe first that, as a function of \( m \), the left side grows faster than the right, so if we prove the inequality for some value of \( m \) it holds for all larger values. Consider \( m = (8d/\varepsilon) \log(8d/\varepsilon) \). Plugging this into the second inequality we have

\[ 2d \log \frac{8d}{\varepsilon} \geq d \log \left( \frac{16d}{\varepsilon} \log \frac{8d}{\varepsilon} \right). \]

It is easy to verify that this is equivalent to showing that

\[ \frac{4d}{\varepsilon} \geq \log \frac{8d}{\varepsilon}. \]

Since \( d \geq 2 \) and \( \varepsilon < 1 \), this is easily checked by direct evaluation.
21 DCELs and Subdivision Intersection

Reading: Chapter 2 in the 4M’s.

Doubly-connected Edge List: We consider the question of how to represent plane straight-line graphs (or PSLG). The DCEL is a common edge-based representation. Vertex and face information is also included for whatever geometric application is using the data structure. There are three sets of records one for each element in the PSLG: vertex records, edge records, and face records. For the purposes of unambiguously defining left and right, each undirected edge is represented by two directed half-edges.

We will make a simplifying assumption that faces do not have holes inside of them. This assumption can be satisfied by introducing some number of dummy edges joining each hole either to the outer boundary of the face, or to some other hole that has been connected to the outer boundary in this way. With this assumption, it may be assumed that the edges bounding each face form a single cyclic list.

Vertex: Each vertex stores its coordinates, along with a pointer to any incident directed edge that has this vertex as its origin, v.inc_edge.

Edge: Each undirected edge is represented as two directed edges. Each edge has a pointer to the oppositely directed edge, called its twin. Each directed edge has an origin and destination vertex. Each directed edge is associate with two faces, one to its left and one to its right. We store a pointer to the origin vertex e.org. (We do not need to define the destination, e.dest, since it may be defined to be e.twin.org.) We store a pointer to the face to the left of the edge e.left (we can access the face to the right from the twin edge). This is called the dent face. We also store the next and previous directed edges in counterclockwise order about the incident face, e.next and e.prev, respectively.

Face: Each face f stores a pointer to a single edge for which this face is the incident face, f.inc_edge. (See the text for the more general case of dealing with holes.)

Figure 21.1: Doubly-connected edge list.

The figure shows two ways of visualizing the DCEL. One is in terms of a collection of doubled-up directed edges. An alternative way of viewing the data structure that gives a better sense of the connectivity structure is based on covering each edge with a two element block, one for e and the other for its twin. The next and prev pointers provide links around each face of the polygon. The next pointers are directed counterclockwise around each face and the prev pointers are directed clockwise.
Of course, in addition the data structure may be enhanced with whatever application data is relevant. In some applications, it is not necessary to know either the face or vertex information (or both) at all, and if so these records may be deleted. See the book for a complete example.

For example, suppose that we wanted to enumerate the vertices that lie on some face \( f \). Here is the code:

```c
Vertex enumeration using DCEL

enumerate_vertices(Face f) {
    Edge start = f.inc_edge;
    Edge e = start;
    do {
        output e.org;
        e = e.next;
    } while (e != start);
}
```

Merging subdivisions: Let us return to the applications problem that lead to the segment intersection problem. Suppose that we have two planar subdivisions, \( S_1 \) and \( S_2 \), and we want to compute their overlay. In particular, this is a subdivision whose vertices are the union of the vertices of each subdivision and the points of intersection of the line segments in the subdivision. (Because we assume that each subdivision is a planar graph, the only new vertices that could arise will come from the intersection of two edges, one from \( S_1 \) and the other from \( S_2 \).) Suppose that each subdivision is represented using a DCEL. Can we adapt the plane-sweep algorithm to generate the DCEL of the overlaid subdivision?

The answer is yes. The algorithm will destroy the original subdivisions, so it may be desirable to copy them before beginning this process. The first part of the process is straightforward, but perhaps a little tedious. This part consists of building the edge and vertex records for the new subdivision. The second part involves building the face records. It is more complicated because it is generally not possible to know the face structure at the moment that the sweep is advancing, without looking “into the future” of the sweep to see whether regions will merge. (You might try to convince yourself of this.) The entire subdivision is built first, and then the face information is constructed and added later. We will skip the part of updating the face information (see the text).

For the first part, the most illustrative case arises when the sweep is processing an intersection event. In this case the two segments arise as two edges \( a_1 \) and \( b_1 \) from the two subdivisions. We will assume that we select the half-edges that are directed from left to right across the sweep-line. The process is described below (and is illustrated in the figure below). It makes use of two auxiliary procedures. \( \text{Split}(a_1, a_2) \) splits an edge \( a_1 \) at its midpoint into two consecutive edges \( a_1 \) followed by \( a_2 \), and links \( a_2 \) into the structure. \( \text{Splice}(a_1, a_2, b_1, b_2) \) takes two such split edges and links them all together.

```c
Merge two edges into a common subdivision

Merge(a_1, b_1) :

1. Create a new vertex \( v \) at the intersection point.
2. Split each of the two intersecting edges, by adding a vertex at the common intersection point. Let \( a_2 \) and \( b_2 \) be the new edge pieces. They are created by the calls \( a_2 = \text{Split}(a_1) \) and \( b_2 = \text{Split}(b_1) \) given below.
3. Link the four edges together by invoking \( \text{Splice}(a_1, a_2, b_1, b_2) \), given below.
```
The splitting procedure creates the new edge, links it into place. After this the edges have been split, but they are not linked to each other. The edge constructor is given the origin and destination of the new edge and creates a new edge and its twin. The procedure below initializes all the other fields. Also note that the destination of \( a_1 \), that is the origin of \( a_1 \)'s twin must be updated, which we have omitted. The splice procedure interlinks four edges around a common vertex in the counterclockwise order \( a_1 \) (entering), \( b_1 \) (entering), \( a_2 \) (leaving), \( b_2 \) (leaving).

```cpp
Split(edge &a1, edge &a2) { // a2 is returned
    a2 = new edge(v, a1.dest()); // create edge (v,a1.dest)
    a2.next = a1.next; a1.next.prev = a2;
    a1.next = a2; a2.prev = a1;
    a1t = a1.twin; a2t = a2.twin; // the twins
    a2t.prev = a1t.prev; a1t.prev.next = a2t;
    a1t.prev = a2t; a2t.next = a1t;
}
```

```cpp
Splice(edge &a1, edge &a2, edge &b1, edge &b2) {
    a1t = a1.twin; a2t = a2.twin; // get the twins
    b1t = b1.twin; b2t = b2.twin;
    a1.next = b2; b2.prev = a1; // link the edges together
    b2t.next = a2; a2.prev = b2t;
    a2t.next = b1t; b1t.prev = a2t;
    b1.next = alt; alt.prev = b1;
}
```
Figure 21.2: Updating the DCEL.
22  Smallest Enclosing Disk

Reading: Chapter 4 in the 4M’s.

Smallest Enclosing Disk: Although the vast majority of applications of linear programming are in relatively high dimensions, there are a number of interesting applications in low dimensions. We will present one such example, called the smallest enclosing disk problem. We are given \( n \) points in the plane and we are asked to find the closed circular disk of minimum radius that encloses all of these points. We will present a randomized algorithm for this problem that runs in \( O(n) \) expected time.

We should say a bit about terminology. A circle is the set of points that are equidistant from some center point. A disk is the set of points lying within a circle. We can talk about open or closed disks to distinguish whether the bounding circle itself is part of the disk. In higher dimensions the generalization of a circle is a sphere in 3-space, or hypersphere in higher dimensions. The set of points lying within a sphere or hypersphere is called a ball.

Before discussing algorithms, we first observe that any circle is uniquely determined by three points (as the circumcenter of the triangle they define). We will not prove this, but it follows as an easy consequence of linearization, which we will discuss later in the lecture.

Claim: For any finite set of points in general position (no four cocircular), the smallest enclosing disk either has at least three points on its boundary, or it has two points, and these points form the diameter of the circle. If there are three points then they subdivide the circle bounding the disk into arcs of angle at most \( \pi \).

Proof: Clearly if there are no points on the boundary the disk’s radius could be decreased. If there is only one point on the boundary then this is also clearly true. If there are two points on the boundary, and they are separated by an arc of length strictly less than \( \pi \), then observe that we can find a disk that passes through both points and has a slightly smaller radius. (By considering a disk whose center point is only the perpendicular bisector of the two points and lies a small distance closer to the line segment joining the points.)

Thus, none of these configurations could be a candidate for the minimum enclosing disk. Also observe that if there are three points that define the smallest enclosing disk they subdivide the circle into three arcs each of angle at most \( \pi \) (for otherwise we could apply the same operation above). Because points are in general position we may assume there cannot be four or more cocircular points.

This immediately suggests a simple \( O(n^4) \) time algorithm. In \( O(n^3) \) time we can enumerate all triples of points and then for each we generate the resulting circle and test whether it encloses all the points in \( O(n) \) additional time, for an \( O(n^4) \) time algorithm. You might make a few observations to improve
this a bit (e.g. by using only triples of points on the convex hull). But even so a reduction from $O(n^4)$ to $O(n)$ is quite dramatic.

**Linearization:** We can “almost” reduce this problem to a linear programming problem in 3-space. Although the method does not work, it does illustrate the similarity between this problem and LP.

Recall that a point $p = (p_x, p_y)$ lies within a circle with center point $c = (c_x, c_y)$ and radius $r$ if

$$(p_x - c_x)^2 + (p_y - c_y)^2 \leq r^2.$$

In our case we are given $n$ such points $p_i$ and are asked to determine whether there exists $c_x$, $c_y$ and $r$ satisfying the resulting $n$ inequalities, with $r$ as small as possible. The problem is that these inequalities clearly involve quantities like $c_x^2$ and $r^2$ and so are not linear inequalities in the parameters of interest.

The technique of linearization can be used to fix this. First let us expand the inequality above and rearrange the terms

$$p_x^2 - 2p_x c_x + c_x^2 + p_y^2 - 2p_y c_y + c_y^2 \leq r^2$$

$$2p_x c_x + 2p_y c_y + (r^2 - c_x^2 - c_y^2) \geq p_x^2 + p_y^2.$$  

Now, let us introduce a new parameter $R = r^2 - c_x^2 - c_y^2$. Now we have

$$(2p_x) c_x + (2p_y) c_y + R \geq (p_x^2 + p_y^2).$$

Observe that this is a linear inequality in $c_x$, $c_y$ and $R$. If we let $p_x$ and $p_y$ range over all the coordinates of all the $n$ points we generate $n$ linear inequalities in 3-space, and so we can apply linear programming to find the solution, right? The only problem is that the previous objective function was to minimize $r$. However $r$ is no longer a parameter in the new version of the problem. Since we $r^2 = R + c_x^2 + c_y^2$, and minimizing $r$ is equivalent to minimizing $r^2$ (since we are only interested in positive $r$), we could say that the objective is to minimize $R + c_x^2 + c_y^2$. Unfortunately, this is not a linear function of the parameters $c_x$, $c_y$ and $R$. Thus we are left with an optimization problem in 3-space with linear constraints and a nonlinear objective function.

This shows that LP is closely related, and so perhaps the same techniques can be applied.

**Randomized Incremental Algorithm:** Let us consider how we can modify the randomized incremental algorithm for LP directly to solve this problem. The algorithm will mimic each step of the randomized LP algorithm.

To start we randomly permute the points. We select any two points and compute the unique circle with these points as diameter. (We could have started with three just as easily.) Let $D_{i-1}$ denote the minimum disk after the insertion of the first $i-1$ points. For point $p_i$ we determine in constant time whether the point lies within $D_{i-1}$. If so, then we set $D_i = D_{i-1}$ and go on to the next stage. If not, then we need to update the current disk to contain $p_i$, letting $D_i$ denote the result. When the last point is inserted we output $D_n$.

How do we compute this updated disk? It might be tempting at first to say that we just need to compute the smallest disk that encloses $p_i$ and the three points that define the current disk. However, it is not hard to construct examples in which doing so will cause previously interior points to fall outside the
current disk. As with the LP problem we need to take all the existing points into consideration. But as in the LP algorithm we want some way to reduce the “dimensionality” of the problem. How do we do this?

The important claim is that if \( p_i \) is not in the minimum disk of the first \( i-1 \) points, then \( p_i \) does help constrain the problem, which we establish below.

**Claim:** If \( p_i \notin D_{i-1} \) then \( p_i \) is on the boundary of the minimum enclosing disk for the first \( i \) points, \( D_i \).

**Proof:** The proof makes use of the following geometric observation. Given a disk of radius \( r_1 \) and a circle of radius \( r_2 \), where \( r_1 < r_2 \), the intersection of the disk with the circle is an arc of angle less than \( \pi \). This is because an arc of angle \( \pi \) or more contains two (diametrically opposite) points whose distance from each other is \( 2r_2 \), but the disk of radius \( r_1 \) has diameter only \( 2r_1 \) and hence could not simultaneously cover two such points.

Now, suppose to the contrary that \( p_i \) is not on the boundary of \( D_i \). It is easy to see that because \( D_i \) covers a point not covered by \( D_{i-1} \) that \( D_i \) must have larger radius than \( D_{i-1} \). If we let \( r_1 \) denote the radius of \( D_{i-1} \) and \( r_2 \) denote the radius of \( D_i \), then by the above argument, the disk \( D_{i-1} \) intersects the circle bounding \( D_i \) in an arc of angle less than \( \pi \). (Shown in a heavy line in the figure below.)

![Figure 22.2: Why \( p_i \) must lie on the boundary of \( D_i \).](image)

Since \( p_i \) is not on the boundary of \( D_i \), the points defining \( D_i \) must be chosen from among the first \( i-1 \) points, from which it follows that they all lie within this arc. However, this would imply that between two of the points is an arc of angle greater than \( \pi \) (the arc not shown with a heavy line) which, by the earlier claim could not be a minimum enclosing disk.

The algorithm is identical in structure to the LP algorithm. We will randomly permute the points and insert them one by one. For each new point \( p_i \), if it lies within the current disk then there is nothing to update. Otherwise, we need to update the disk. We do this by computing the smallest enclosing disk that contains all the points \( \{p_1, \ldots, p_{i-1}\} \) and is constrained to have \( p_i \) on its boundary. (The requirement that \( p_i \) be on the boundary is analogous to the constraint used in linear programming that optimum vertex lie on the line supporting the current halfplane.)

This will involve a slightly different recursion. In this recursion, when we encounter a point that lies outside the current disk, we will then recurse on a subproblem in which two points are constrained to lie on the boundary of the disk. Finally, if this subproblem requires a recursion, we will have a problem in which there are three points constrained to lie on a the boundary of the disk. But this problem is trivial, since there is only one circle passing through three points.
MinDisk\((P)\) :

1. If \(|P| \leq 3\), then return the disk passing through these points. Otherwise, randomly permute the points in \(P\) yielding the sequence \((p_1, p_2, \ldots, p_n)\).
2. Let \(D_2\) be the minimum disk enclosing \(\{p_1, p_2\}\).
3. for \(i = 3\) to \(|P|\) do
   - (a) if \(p_i \in D_{i-1}\) then \(D_i = D_{i-1}\).
   - (a) else \(D_i = \text{MinDiskWith1Pt}(P[1..i-1], p_i)\).

MinDiskWith1Pt\((P, q)\) :

1. Randomly permute the points in \(P\). Let \(D_1\) be the minimum disk enclosing \(\{q, p_1\}\).
2. for \(i = 2\) to \(|P|\) do
   - (a) if \(p_i \in D_{i-1}\) then \(D_i = D_{i-1}\).
   - (a) else \(D_i = \text{MinDiskWith2Pts}(P[1..i-1], q, p_i)\).

MinDiskWith2Pts\((P, q_1, q_2)\) :

1. Randomly permute the points in \(P\). Let \(D_0\) be the minimum disk enclosing \(\{q_1, q_2\}\).
2. for \(i = 1\) to \(|P|\) do
   - (a) if \(p_i \in D_{i-1}\) then \(D_i = D_{i-1}\).
   - (a) else \(D_i = \text{Disk}(q_1, q_2, p_i)\).
23 Interval Trees

Reading: Chapter 10 in the 4M’s.

Segment Data: So far we have considered geometric data structures for storing points. However, there are many others types of geometric data that we may want to store in a data structure. Today we consider how to store orthogonal (horizontal and vertical) line segments in the plane. We assume that a line segment is represented by giving its pair of endpoints. The segments are allowed to intersect one another.

As a basic motivating query, we consider the following window query. Given a set of orthogonal line segments \( S \), which have been preprocessed, and given an orthogonal query rectangle \( W \), count or report all the line segments of \( S \) that intersect \( W \). We will assume that \( W \) is closed and solid rectangle, so that even if a line segment lies entirely inside of \( W \) or intersects only the boundary of \( W \), it is still reported. For example, given the window below, the query would report the segments that are shown with solid lines, and segments with broken lines would not be reported.

![Figure 23.1: Window Query.](image)

Window Queries for Orthogonal Segments: We will present a data structure, called the interval tree, which (combined with a range tree) can answer window counting queries for orthogonal line segments in \( O(\log^2 n) \) time, where \( n \) is the number line segments. It can report these segments in \( O(k + \log^2 n) \) time, where and \( k \) is the total number of segments reported. The interval tree uses \( O(n \log n) \) storage and can be built in \( O(n \log n) \) time.

We will consider the case of range reporting queries. (There are some subtleties in making this work for counting queries.) We will derive our solution in steps, starting with easier subproblems and working up to the final solution. To begin with, observe that the set of segments that intersect the window can be partitioned into three types: those that have no endpoint in \( W \), those that have one endpoint in \( W \), and those that have two endpoints in \( W \).

We already have a way to report segments of the second and third types. In particular, we may build a range tree just for the \( 2n \) endpoints of the segments. We assume that each endpoint has a cross-link indicating the line segment with which it is associated. Now, by applying a range reporting query to \( W \) we can report all these endpoints, and follow the cross-links to report the associated segments. Note that segments that have both endpoints in the window will be reported twice, which is somewhat unpleasant. We could fix this either by sorting the segments in some manner and removing duplicates, or by marking each segment as it is reported and ignoring segments that have already been marked. (If we use marking, after the query is finished we will need to go back an “unmark” all the reported segments in preparation for the next query.)
All that remains is how to report the segments that have no endpoint inside the rectangular window. We will do this by building two separate data structures, one for horizontal and one for vertical segments. A horizontal segment that intersects the window but neither of its endpoints intersects the window must pass entirely through the window. Observe that such a segment intersects any vertical line passing from the top of the window to the bottom. In particular, we could simply ask to report all horizontal segments that intersect the left side of \( W \). This is called a \textit{vertical segment stabbing query}. In summary, it suffices to solve the following subproblems (and remove duplicates):

**Endpoint inside:** Report all the segments of \( S \) that have at least one endpoint inside \( W \). (This can be done using a range query.)

**Horizontal through segments:** Report all the horizontal segments of \( S \) that intersect the left side of \( W \). (This reduces to a vertical segment stabbing query.)

**Vertical through segments:** Report all the vertical segments of \( S \) that intersect the bottom side of \( W \). (This reduces to a horizontal segment stabbing query.)

We will present a solution to the problem of vertical segment stabbing queries. Before dealing with this, we will first consider a somewhat simpler problem, and then modify this simple solution to deal with the general problem.

**Vertical Line Stabbing Queries:** Let us consider how to answer the following query, which is interesting in its own right. Suppose that we are given a collection of horizontal line segments \( S \) in the plane and are given an (infinite) vertical query line \( \ell_q : x = x_q \). We want to report all the line segments of \( S \) that intersect \( \ell_q \). Notice that for the purposes of this query, the \( y \)-coordinates are really irrelevant, and may be ignored. We can think of each horizontal line segment as being a closed \textit{interval} along the \( x \)-axis. We show an example in the figure below on the left.

![Figure 23.2: Line Stabbing Query.](image)

As is true for all our data structures, we want some balanced way to decompose the set of intervals into subsets. Since it is difficult to define some notion of order on intervals, we instead will order the endpoints. Sort the interval endpoints along the \( x \)-axis. Let \( \langle x_1, x_2, \ldots, x_{2n} \rangle \) be the resulting sorted sequence. Let \( x_{\text{med}} \) be the median of these \( 2n \) endpoints. Split the intervals into three groups, \( L \) those that lie strictly to the left of \( x_{\text{med}} \), \( R \) those that lie strictly to the right of \( x_{\text{med}} \), and \( M \) those that contain the point \( x_{\text{med}} \). We can then define a binary tree by putting the intervals of \( L \) in the left subtree and recursing, putting the intervals of \( R \) in the right subtree and recursing. Note that if \( x_q < x_{\text{med}} \) we can eliminate the right subtree and if \( x_q > x_{\text{med}} \) we can eliminate the left subtree. See the figure right.

But how do we handle the intervals of \( M \) that contain \( x_{\text{med}} \)? We want to know which of these intervals intersects the vertical line \( \ell_q \). At first it may seem that we have made no progress, since it appears that...
we are back to the same problem that we started with. However, we have gained the information that all these intervals intersect the vertical line \( x = x_{\text{med}} \). How can we use this to our advantage?

Let us suppose for now that \( x_q \leq x_{\text{med}} \). How can we store the intervals of \( M \) to make it easier to report those that intersect \( \ell_q \)? The simple trick is to sort these lines in increasing order of their left endpoint. Let \( M_L \) denote the resulting sorted list. Observe that if some interval in \( M_L \) does not intersect \( \ell_q \), then its left endpoint must be to the right of \( x_q \), and hence none of the subsequent intervals intersects \( \ell_q \). Thus, to report all the segments of \( M_L \) that intersect \( \ell_q \), we simply traverse the sorted list and list elements until we find one that does not intersect \( \ell_q \), that is, whose left endpoint lies to the right of \( x_q \). As soon as this happens we terminate. If \( k' \) denotes the total number of segments of \( M \) that intersect \( \ell_q \), then clearly this can be done in \( O(k' + 1) \) time.

On the other hand, what do we do if \( x_q > x_{\text{med}} \)? This case is symmetrical. We simply sort all the segments of \( M \) in a sequence, \( M_R \), which is sorted from right to left based on the right endpoint of each segment. Thus each element of \( M \) is stored twice, but this will not affect the size of the final data structure by more than a constant factor. The resulting data structure is called an interval tree.

**Interval Trees:** The general structure of the interval tree was derived above. Each node of the interval tree has a left child, right child, and itself contains the median \( x \)-value used to split the set, \( x_{\text{med}} \), and the two sorted sets \( M_L \) and \( M_R \) (represented either as arrays or as linked lists) of intervals that overlap \( x_{\text{med}} \). We assume that there is a constructor that builds a node given these three entities. The following high-level pseudocode describes the basic recursive step in the construction of the interval tree. The initial call is \( \text{root} = \text{IntTree}(S) \), where \( S \) is the initial set of intervals. Unlike most of the data structures we have seen so far, this one is not built by the successive insertion of intervals (although it would be possible to do so). Rather we assume that a set of intervals \( S \) is given as part of the constructor, and the entire structure is built all at once. We assume that each interval in \( S \) is represented as a pair \((x_{\text{lo}}, x_{\text{hi}})\). An example is shown in the following figure.

```java
IntTreeNode IntTree(IntervalSet S) {
    if (|S| == 0) return null // no more
    xMed = median endpoint of intervals in S // median endpoint
    L = \{[xlo, xhi] in S | xhi < xMed\} // left of median
    R = \{[xlo, xhi] in S | xlo > xMed\} // right of median
    M = \{[xlo, xhi] in S | xlo <= xMed <= xhi\} // contains median
    ML = sort M in increasing order of xlo // sort M
    MR = sort M in decreasing order of xhi
    t = new IntTreeNode(xMed, ML, MR) // this node
    t.left = IntTree(L) // left subtree
    t.right = IntTree(R) // right subtree
    return t
}
```

We assert that the height of the tree is \( O(\log n) \). To see this observe that there are \( 2n \) endpoints. Each time through the recursion we split this into two subsets \( L \) and \( R \) of sizes at most half the original size.
Computational Geometry Notes

Interval Trees

(minus the elements of \( M \)). Thus after at most \( \lg(2n) \) levels we will reduce the set sizes to 1, after which the recursion bottoms out. Thus the height of the tree is \( O(\log n) \).

Implementing this constructor efficiently is a bit subtle. We need to compute the median of the set of all endpoints, and we also need to sort intervals by left endpoint and right endpoint. The fastest way to do this is to pre-sort all these values and store them in three separate lists. Then as the sets \( L, R, \) and \( M \) are computed, we simply copy items from these sorted lists to the appropriate sorted lists, maintaining their order as we go. If we do so, it can be shown that this procedure builds the entire tree in \( O(n \log n) \) time.

The algorithm for answering a stabbing query was derived above. We summarize this algorithm below.

Let \( x_q \) denote the \( x \)-coordinate of the query line.

---

**Line Stabbing Queries for an Interval Tree**

```java
int stab(IntTreeNode t, Scalar xq) {
    if (t == null) return // fell out of tree
    if (xq < t.xMed) { // left of median?
        for (i = 0; i < t.ML.length; i++) { // traverse ML
            if (t.ML[i].lo <= xq) print(t.ML[i]) // ..report if in range
                else break // ..else done
        }
        stab(t.left, xq) // recurse on left
    }
    else { // right of median
        for (i = 0; i < t.MR.length; i++) { // traverse MR
            if (t.MR[i].hi >= xq) print(t.MR[i]) // ..report if in range
                else break // ..else done
        }
        stab(t.right, xq) // recurse on right
    }
}
```

---

This procedure actually has one small source of inefficiency, which was intentionally included to make code look more symmetric. Can you spot it? Suppose that \( x_q = t.x_{\text{med}} \)? In this case we will recursively search the right subtree. However this subtree contains only intervals that are strictly to the right of \( x_{\text{med}} \) and so is a waste of effort. However it does not affect the asymptotic running time.

As mentioned earlier, the time spent processing each node is \( O(1+k') \) where \( k' \) is the total number of points that were recorded at this node. Summing over all nodes, the total reporting time is \( O(k + v) \), where \( k \) is the total number of intervals reported, and \( v \) is the total number of nodes visited. Since at
each node we recurse on only one child or the other, the total number of nodes visited \( v \) is \( O(log n) \), the height of the tree. Thus the total reporting time is \( O(k + log n) \).

**Vertical Segment Stabbing Queries:** Now let us return to the question that brought us here. Given a set of horizontal line segments in the plane, we want to know how many of these segments intersect a vertical line segment. Our approach will be exactly the same as in the interval tree, except for how the elements of \( M \) (those that intersect the splitting line \( x = x_{med} \)) are handled.

Going back to our interval tree solution, let us consider the set \( M \) of horizontal line segments that intersect the splitting line \( x = x_{med} \) and as before let us consider the case where the query segment \( q \) with endpoints \((x_q, y_{lo})\) and \((x_q, y_{hi})\) lies to the left of the splitting line. The simple trick of sorting the segments of \( M \) by their left endpoints is not sufficient here, because we need to consider the \( y \)-coordinates as well. Observe that a segment of \( M \) stabs the query segment \( q \) if and only if the left endpoint of a segment lies in the following semi-infinite rectangular region.

\[
\{(x, y) \mid x \leq x_q \text{ and } y_{lo} \leq y \leq y_{hi}\}
\]

This is illustrated in the figure below. Observe that this is just an orthogonal range query. (It is easy to generalize the procedure given last time to handle semi-infinite rectangles.) The case where \( q \) lies to the right of \( x_{med} \) is symmetrical.

![Figure 23.4: The segments that stab \( q \) lie within the shaded semi-infinite rectangle.](image)

So the solution is that rather than storing \( M_L \) as a list sorted by the left endpoint, instead we store the left endpoints in a 2-dimensional range tree (with cross-links to the associated segments). Similarly, we create a range tree for the right endpoints and represent \( M_R \) using this structure.

The segment stabbing queries are answered exactly as above for line stabbing queries, except that part that searches \( M_L \) and \( M_R \) (the for-loops) are replaced by searches to the appropriate range tree, using the semi-infinite range given above.

We will not discuss construction time for the tree. (It can be done in \( O(n \log n) \) time, but this involves some thought as to how to build all the range trees efficiently). The space needed is \( O(n \log n) \), dominated primarily from the \( O(n \log n) \) space needed for the range trees. The query time is \( O(k + \log^3 n) \), since we need to answer \( O(\log n) \) range queries and each takes \( O(\log^2 n) \) time plus the time for reporting. If we use the spiffy version of range trees (which we mentioned but never discussed) that can answer queries in \( O(k + \log n) \) time, then we can reduce the total time to \( O(k + \log^2 n) \).
24 Hereditary Segment Trees and Red-Blue Intersection

**Reading:** Segment trees are presented in Chapt 10 of the 4M’s. However, most of today’s material has been taken from the paper, “Algorithms for Bichromatic Line-Segment Problems and Polyhedral Terrains,” by B. Chazelle, H. Edelsbrunner, L. J. Guibas, and M. Sharir, *Algorithmica*, 11, 1994 116–132.

**Red-Blue Segment Intersection:** We have been talking about the use of geometric data structures for solving query problems. Often data structures are used as intermediate structures for solving traditional input/output problems, which do not involve preprocessing and queries. (Another famous example of this is HeapSort, which introduces the heap data structure for sorting a list of numbers.) Today we will discuss a variant of a useful data structure, the segment tree. The particular variant is called a hereditary segment tree. It will be used to solve the following problem.

**Red-Blue Segment Intersection:** Given a set $B$ of $m$ pairwise disjoint “blue” segments in the plane and a set $R$ of $n$ pairwise disjoint “red” segments, count (or report) all bichromatic pairs of intersecting line segments (that is, intersections between red and blue segments).

It will make things simpler to think of the segments as being open (not including their endpoints). In this way, the pairwise disjoint segments might be the edges of a planar straight line graph (PSLG). Indeed, one of the most important application of red-blue segment intersection involves computing the overlay of two PSLG’s (one red and the other blue) This is also called the map overlay problem, and is often used in geographic information systems. The most time consuming part of the map overlay problem is determining which pairs of segments overlap. See the figure below.

![Figure 24.1: Red-blue line segment intersection. The algorithm outputs the white intersection points between segments of different colors. The segments of each color are pairwise disjoint (except possibly at their endpoints).](image)

Let $N = n + m$ denote the total input size and let $k$ denote the total number of bichromatic intersecting pairs. We will present an algorithm for this problem that runs in $O(k + N \log^2 N)$ time for the reporting problem and $O(N \log^2 N)$ time for the counting problem. Both algorithms use $O(N \log N)$ space. Although we will not discuss it (but the original paper does) it is possible to remove a factor of $\log n$ from both the running time and space, using a somewhat more sophisticated variant of the algorithm that we will present.

Because the set of red segments are each pairwise disjoint as are the blue segments, it follows that we could solve the reporting problem by our plane sweep algorithm for segment intersection (as discussed
in an earlier lecture) in $O((N + k) \log N)$ time and $O(N)$ space. Thus, the more sophisticated algorithm is an improvement on this. However, plane sweep will not allow us to solve the counting problem.

**The Hereditary Segment Tree:** Recall that we are given two sets $B$ and $R$, consisting of, respectively, $m$ and $n$ line segments in the plane, and let $N = m + n$. Let us make the general position assumption that the $2N$ endpoints of these line segments have distinct $x$-coordinates. The $x$-coordinates of these endpoints subdivide the $x$-axis into $2N + 1$ intervals, called *atomic intervals*. We construct a balanced binary tree whose leaves are in 1–1 correspondence with these intervals, ordered from left to right. Each internal node $u$ of this tree is associated with an interval $I_u$ of the $x$-axis, consisting of the union of the intervals of its descendent leaves. We can think of each such interval as a vertical slab $S_u$ whose intersection with the $x$-axis is $I_u$. (See the figure below, left.)

![Figure 24.2: Hereditary Segment Tree: Intervals, slabs and the nodes associated with a segment.](image1)

We associate a segment $s$ with a set of nodes of the tree. A segment is said to span interval $I_u$ if its projection covers this interval. We associate a segment $s$ with a node $u$ if $s$ spans $I_u$ but $s$ does not span $I_p$, where $p$ is $u$’s parent. (See the figure above, right.)

Each node (internal or leaf) of this tree is associated with a list, called the *blue standard list*, $B_u$ of all blue line segments whose vertical projection contains $I_u$ but does not contain $I_p$, where $p$ is the parent of $u$. Alternately, if we consider the nodes in whose standard list a segment is stored, the intervals corresponding to these nodes constitute a disjoint cover of the segment’s vertical projection. The node is also associated with a red standard list, denoted $R_u$, which is defined analogously for the red segments. (See the figure below, left.)

![Figure 24.3: Hereditary Segment Tree with standard lists (left) and hereditary lists (right).](image2)

Each node $u$ is also associated with a list $B^*_u$, called the *blue hereditary list*, which is the union of the $B_v$ for all proper descendents $v$ or $u$. The red hereditary list $R^*_u$ is defined analogously. (Even
though a segment may occur in the standard list for many descendents, there is only one copy of each segment in the hereditary lists.) The segments of \( R_u \) and \( B_u \) are called the long segments, since they span the entire interval. The segments of \( R^*_u \) and \( B^*_u \) are called the short segments, since they do not span the entire interval.

By the way, if we ignored the fact that we have two colors of segments and just considered the standard lists, the resulting tree is called a segment tree. The addition of the hereditary lists makes this a hereditary segment tree. Our particular data structure differs from the standard hereditary segment tree in that we have partitioned the various segment lists according to whether the segment is red or blue.

**Time and Space Analysis:** We claim that the total size of the hereditary segment tree is \( O(N \log N) \). To see this observe that each segment is stored in the standard list of at most \( 2 \log N \) nodes. The argument is very similar to the analysis of the 1-dimensional range tree. If you locate the left and right endpoints of the segment among the atomic intervals, these define two paths in the tree. In the same manner as canonical sets for the 1-dimensional range tree, the segment will be stored in all the “inner” nodes between these two paths. (See the figure below.) The segment will also be stored in the hereditary lists for all the ancestors of these nodes. These ancestors lie along the two paths to the left and right, and hence there are at most \( 2 \log N \) of them. Thus, each segment appears in at most \( 4 \log N \) lists, for a total size of \( O(N \log N) \).

The tree can be built in \( O(N \log N) \) time. In \( O(N \log N) \) time we can sort the \( 2N \) segment endpoints. Then for each segment, we search for its left and right endpoints and insert the segment into the standard and hereditary lists for the appropriate nodes and we descend each path in \( O(1) \) time for each node visited. Since each segment appears in \( O(\log N) \) lists, this will take \( O(\log N) \) time per segment and \( O(N \log N) \) time overall.

**Computing Intersections:** Let us consider how to use the hereditary segment tree to count and report bichromatic intersections. We will do this on a node-by-node basis. Consider any node \( u \). We classify the intersections into two types, long-long intersections are those between a segment of \( B_u \) and \( R_u \), and long-short intersections are those between a segment of \( B^*_u \) and \( R_u \) or between \( R^*_u \) and \( B_u \). Later we will show that by considering just these intersection cases, we will consider every intersection exactly once.

**Long-long intersections:** Sort each of the lists \( B_u \) and \( R_u \) of long segments in ascending order by \( y \)-coordinate. (Since the segments of each set are disjoint, this order is constant throughout the interval for each set.) Let \( \langle b_1, b_2, \ldots, b_m \rangle \) and \( \langle r_1, r_2, \ldots, r_n \rangle \) denote these ordered lists. Merge these lists twice, once according to their order along the left side of the slab and one according to their order along the right side of the slab.
Observe that for each blue segment $b \in B_u$, this allows us to determine two indices $i$ and $j$, such that $b$ lies between $r_i$ and $r_{i+1}$ along the left boundary and between $r_j$ and $r_{j+1}$ along the right boundary. (For convenience, we can think of segment 0 as an imaginary segment at $y = -\infty$.) It follows that if $i < j$ then $b$ intersects the red segments $r_{i+1}, \ldots, r_j$. (See the figure below, (a)). On the other hand, if $i \geq j$ then $b$ intersects the red segments $r_{j+1}, \ldots, r_i$. (See the figure below, (b)). We can count these intersections in $O(1)$ time or report them in time proportional to the number of intersections.

For example, consider the segment $b = b_2$ in the figure below, (c). On the left boundary it lies between $r_3$ and $r_4$, and hence $i = 3$. On the right boundary it lies between $r_0$ and $r_1$, and hence $j = 0$. (Recall that $r_0$ is at $y = -\infty$.) Thus, since $i \geq j$ it follows that $b$ intersects the three red segments $\{r_1, r_2, r_3\}$.

![Figure 24.5: Red-blue intersection counting/reporting. Long-long intersections.](image)

The total time to do this is dominated by the $O(m_u \log m_u + n_u \log n_u)$ time needed to sort both lists. The merging and counting only requires linear time.

**Long-short intersections:** There are two types of long-short intersections to consider. Long red and short blue, and long blue and short red. Let us consider the first one, since the other one is symmetrical.

As before, sort the long segments of $R_u$ in ascending order according to $y$-coordinate, letting $\langle r_1, r_2, \ldots, r_{n_u}\rangle$ denote this ordered list. These segments naturally subdivide the slab into $n_u + 1$ trapezoids. For each short segment $b \in B^*_u$, perform two binary searches among the segments of $R_u$ to find the lowest segment $r_i$ and the highest segment $r_j$ that $b$ intersects. (See the figure above, right.) Then $b$ intersects all the red segments $r_i, r_{i+1}, \ldots, r_j$.

![Figure 24.6: Red-blue intersection counting/reporting: Long-short intersections.](image)

Thus, after $O(\log n_u)$ time for the binary searches, the segments of $R_u$ intersecting $b$ can be counted in $O(1)$ time, for a total time of $O(m_u \log n_u)$. Reporting can be done in time propor-
tional to the number of intersections reported. Adding this to the time for the long blue and short red case, we have a total time complexity of $O(m_u^* \log n_u + n_u^* \log m_u)$.

If we let $N_u = m_u + n_u + m_u^* + n_u^*$, then observe that the total time to process vertex $u$ is $O(N_u \log N_u)$ time. Summing this over all nodes of the tree, and recalling that $\sum_u N_u = O(N \log N)$ we have a total time complexity of

$$T(N) = \sum_u N_u \log N_u \leq \left( \sum_u N_u \right) \log N = O(N \log^2 N).$$

**Correctness:** To show that the algorithm is correct, we assert that each bichromatic intersection is counted exactly once. For any bichromatic intersection between $b_i$ and $r_j$ consider the leaf associated with the atomic interval containing this intersection point. As we move up to the ancestors of this leaf, we will encounter $b_i$ in the standard list of one of these ancestors, denoted $u_i$, and will encounter $r_j$ at some node, denoted $u_j$. If $u_i = u_j$ then this intersection will be detected as a long-long intersection at this node. Otherwise, one is a proper ancestor of the other, and this will be detected as a long-short intersection (with the ancestor long and descendent short).
25  Kirkpatrick’s Planar Point Location

Today’s material is not covered in our text. See the book by Preparata and Shamos, *Computational Geometry*, Section 2.2.2.3.

**Point Location:** The point location problem (in 2-space) is: given a polygonal subdivision of the plane (that is, a PSLG) with \( n \) vertices, preprocess this subdivision so that given a query point \( q \), we can efficiently determine which face of the subdivision contains \( q \). We may assume that each face has some identifying label, which is to be returned. We also assume that the subdivision is represented in any “reasonable” form (e.g. as a DCEL). In general \( q \) may coincide with an edge or vertex. To simplify matters, we will assume that \( q \) does not lie on an edge or vertex, but these special cases are not hard to handle.

It is remarkable that although this seems like such a simple and natural problem, it took quite a long time to discover a method that is optimal with respect to both query time and space. It has long been known that there are data structures that can perform these searches reasonably well (e.g. quad-trees and kd-trees), but for which no good theoretical bounds could be proved. There were data structures of with \( O(\log n) \) query time but \( O(n \log n) \) space, and \( O(n) \) space but \( O(\log^2 n) \) query time.

The first construction to achieve both \( O(n) \) space and \( O(\log n) \) query time was a remarkably clever construction due to Kirkpatrick. It turns out that Kirkpatrick’s idea has some large embedded constant factors that make it less attractive practically, but the idea is so clever that it is worth discussing, nonetheless. Later we will discuss a more practical randomized method that is presented in our text.

**Kirkpatrick’s Algorithm:** Kirkpatrick’s idea starts with the assumption that the planar subdivision is a triangulation, and further that the outer face is a triangle. If this assumption is not met, then we begin by triangulating all the faces of the subdivision. The label associated with each triangular face is the same as a label for the original face that contained it. For the outer face is not a triangle, first compute the convex hull of the polygonal subdivision, triangulate everything inside the convex hull. Then surround this convex polygon with a large triangle (call the vertices \( a, b, \) and \( c \)), and then add edges from the convex hull to the vertices of the convex hull. It may sound like we are adding a lot of new edges to the subdivision, but recall from earlier in the semester that the number of edges and faces in any straight-line planar subdivision is proportional to \( n \), the number of vertices. Thus the addition only increases the size of the structure by a constant factor.

Note that once we find the triangle containing the query point in the augmented graph, then we will know the original face that contains the query point. The triangulation process can be performed in \( O(n \log n) \) time by a plane sweep of the graph, or in \( O(n) \) time if you want to use sophisticated methods like the linear time polygon triangulation algorithm. In practice, many straight-line subdivisions, may already have convex faces and these can be triangulated easily in \( O(n) \) time.

![Triangulation of a planar subdivision.](image)
Let $T_0$ denote the initial triangulation. What Kirkpatrick’s method does is to produce a sequence of triangulations, $T_0, T_1, T_2, \ldots, T_k$, where $k = O(\log n)$, such that $T_k$ consists only of a single triangle (the exterior face of $T_0$), and each triangle in $T_{i+1}$ overlaps a constant number of triangles in $T_i$.

We will see how to use such a structure for point location queries later, but for now let us concentrate on how to build such a sequence of triangulations. Assuming that we have $T_i$, we wish to compute $T_{i+1}$. In order to guarantee that this process will terminate after $O(\log n)$ stages, we will want to make sure that the number of vertices in $T_{i+1}$ decreases by some constant factor from the number of vertices in $T_i$. In particular, this will be done by carefully selecting a subset of vertices of $T_i$ and deleting them (and along with them, all the edges attached to them). After these vertices have been deleted, we need retriangulate the resulting graph to form $T_{i+1}$. The question is: How do we select the vertices of $T_i$ to delete, so that each triangle of $T_{i+1}$ overlaps only a constant number of triangles in $T_i$?

There are two things that Kirkpatrick observed at this point, that make the whole scheme work.

**Constant degree:** We will make sure that each of the vertices that we delete have constant ($\leq d$) degree (that is, each is adjacent to at most $d$ edges). Note that the when we delete such a vertex, the resulting hole will consist of at most $d - 2$ triangles. When we retriangulate, each of the new triangles, can overlap at most $d$ triangles in the previous triangulation.

**Independent set:** We will make sure that no two of the vertices that are deleted are adjacent to each other, that is, the vertices to be deleted form an independent set in the current planar graph $T_i$. This will make retriangulation easier, because when we remove $m$ independent vertices (and their incident edges), we create $m$ independent holes (non triangular faces) in the subdivision, which we will have to retriangulate. However, each of these holes can be triangulated independently of one another. (Since each hole contains a constant number of vertices, we can use any triangulation algorithm, even brute force, since the running time will be $O(1)$ in any case.)

An important question to the success of this idea is whether we can always find a sufficiently large independent set of vertices with bounded degree. We want the size of this set to be at least a constant fraction of the current number of vertices. Fortunately, the answer is “yes,” and in fact it is quite easy to find such a subset. Part of the trick is to pick the value of $d$ to be large enough (too small and there may not be enough of them). It turns out that $d = 8$ is good enough.

**Lemma:** Given a planar graph with $n$ vertices, there is an independent set consisting of vertices of degree at most 8, with at least $n/18$ vertices. This independent set can be constructed in $O(n)$ time.

We will present the proof of this lemma later. The number 18 seems rather large. The number is probably smaller in practice, but this is the best bound that this proof generates. However, the size of this constant is one of the reasons that Kirkpatrick’s algorithm is not used in practice. But the construction is quite clever, nonetheless, and once a optimal solution is known to a problem it is often not long before a practical optimal solution follows.

**Kirkpatrick Structure:** Assuming the above lemma, let us give the description of how the point location data structure, the Kirkpatrick structure, is constructed. We start with $T_0$, and repeatedly select an independent set of vertices of degree at most 8. We never include the three vertices $a$, $b$, and $c$ (forming the outer face) in such an independent set. We delete the vertices from the independent set from the
graph, and retriangulate the resulting holes. Observe that each triangle in the new triangulation can overlap at most 8 triangles in the previous triangulation. Since we can eliminate a constant fraction of vertices with each stage, after $O(\log n)$ stages, we will be down to the last 3 vertices.

The constant factors here are not so great. With each stage, the number of vertices falls by a factor of $17/18$. To reduce to the final three vertices, implies that $(18/17)^k = n$ or that

$$k = \log_{18/17} n \approx 12 \log n.$$

It can be shown that by always selecting the vertex of smallest degree, this can be reduced to a more palatable $4.5 \log n$.

The data structure is based on this decomposition. The root of the structure corresponds to the single triangle of $T_k$. The nodes at the next lower level are the triangles of $T_{k-1}$, followed by $T_{k-2}$, until we reach the leaves, which are the triangles of our initial triangulation, $T_0$. Each node for a triangle in triangulation $T_{i+1}$, stores pointers to all the triangles it overlaps in $T_i$ (there are at most 8 of these). Note that this structure is a directed acyclic graph (DAG) and not a tree, because one triangle may have many parents in the data structure. This is shown in the following figure.

![Kirkpatrick’s point location structure](image)

Figure 25.2: Kirkpatrick’s point location structure.

To locate a point, we start with the root, $T_k$. If the query point does not lie within this single triangle, then we are done (it lies in the exterior face). Otherwise, we search each of the (at most 8) triangles in $T_{k-1}$ that overlap this triangle. When we find the correct one, we search each of the triangles in $T_{k-2}$ that overlap this triangles, and so forth. Eventually we will find the triangle containing the query point in the last triangulation, $T_0$, and this is the desired output. See the figure below for an example.
Computational Geometry Notes  
Kirkpatrick’s Algorithm

**Construction and Analysis:** The structure has $O(\log n)$ levels (one for each triangulation), it takes a constant amount of time to move from one level to the next (at most 8 point-in-triangle tests), thus the total query time is $O(\log n)$. The size of the data structure is the sum of sizes of the triangulations. Since the number of triangles in a triangulation is proportional to the number of vertices, it follows that the size is proportional to

$$n(1 + 17/18 + (17/18)^2 + (17/18)^3 + \ldots) \leq 18n,$$

(using standard formulas for geometric series). Thus the data structure size is $O(n)$ (again, with a pretty hefty constant).

The last thing that remains is to show how to construct the independent set of the appropriate size. We first present the algorithm for finding the independent set, and then prove the bound on its size.

1. Mark all nodes of degree $\geq 9$.
2. While there exists an unmarked node do the following:
   a. Choose an unmarked vertex $v$.
   b. Add $v$ to the independent set.
   c. Mark $v$ and all of its neighbors.

It is easy to see that the algorithm runs in $O(n)$ time (e.g., by keeping unmarked vertices in a stack and representing the triangulation so that neighbors can be found quickly.)

Intuitively, the argument that there exists a large independent set of low degree is based on the following simple observations. First, because the average degree in a planar graph is less than 6, there must be a lot of vertices of degree at most 8 (otherwise the average would be unattainable). Second, whenever we add one of these vertices to our independent set, only 8 other vertices become ineligible for inclusion in the independent set.

Here is the rigorous argument. Recall from Euler’s formula, that if a planar graph is fully triangulated, then the number of edges $e$ satisfies $e = 3n - 6$. If we sum the degrees of all the vertices, then each
edge is counted twice. Thus the average degree of the graph is
\[ \sum_{v} \deg(v) = 2e = 6n - 12 < 6n. \]

Next, we claim that there must be at least \( n/2 \) vertices of degree 8 or less. To see why, suppose to the contrary that there were more than \( n/2 \) vertices of degree 9 or greater. The remaining vertices must have degree at least 3 (with the possible exception of the 3 vertices on the outer face), and thus the sum of all degrees in the graph would have to be at least as large as
\[ 9 \frac{n}{2} + 3 \frac{n}{2} = 6n, \]
which contradicts the equation above.

Now, when the above algorithm starts execution, at least \( n/2 \) vertices are initially unmarked. Whenever we select such a vertex, because its degree is 8 or fewer, we mark at most 9 new vertices (this node and at most 8 of its neighbors). Thus, this step can be repeated at least \( (n/2)/9 = n/18 \) times before we run out of unmarked vertices. This completes the proof.
26 Divide-and-Conquer Algorithm for Voronoi Diagrams

**Reading:** O’Rourke, Chapt 5, Mulmuley, Sect. 2.8.4.

**Planar Voronoi Diagrams:** Recall that, given \( n \) points \( P = \{p_1, p_2, \ldots, p_n\} \) in the plane, the Voronoi polygon of a point \( p_i \), \( V(p_i) \), is defined to be the set of all points \( q \) in the plane for which \( p_i \) is among the closest points to \( q \) in \( P \). That is,

\[
V(p_i) = \{ q : |p_i - q| \leq |p_j - q|, \forall j \neq i \}.
\]

The union of the boundaries of the Voronoi polygons is called the Voronoi diagram of \( P \), denoted \( VD(P) \). The dual of the Voronoi diagram is a triangulation of the point set, called the Delaunay triangulation. Recall from our discussion of quad-edge data structure, that given a good representation of any planar graph, the dual is easy to construct. Hence, it suffices to show how to compute either one of these structures, from which the other can be derived easily in \( O(n) \) time.

There are four fairly well-known algorithms for computing Voronoi diagrams and Delaunay triangulations in the plane. They are

- **Divide-and-Conquer:** (For both VD and DT.) The first \( O(n \log n) \) algorithm for this problem. Not widely used because it is somewhat hard to implement. Can be generalized to higher dimensions with some difficulty. Can be generalized to computing Voronoi diagrams of line segments with some difficulty.

- **Randomized Incremental:** (For DT and VD.) The simplest. \( O(n \log n) \) time with high probability. Can be generalized to higher dimensions as with the randomized algorithm for convex hulls. Can be generalized to computing Voronoi diagrams of line segments fairly easily.

- **Fortune’s Plane Sweep:** (For VD.) A very clever and fairly simple algorithm. It computes a “deformed” Voronoi diagram by plane sweep in \( O(n \log n) \) time, from which the true diagram can be extracted easily. Can be generalized to computing Voronoi diagrams of line segments fairly easily.

- **Reduction to convex hulls:** (For DT.) Computing a Delaunay triangulation of \( n \) points in dimension \( d \) can be reduced to computing a convex hull of \( n \) points in dimension \( d + 1 \). Use your favorite convex hull algorithm. Unclear how to generalize to compute Voronoi diagrams of line segments.

We will cover all of these approaches, except Fortune’s algorithm. O’Rourke does not give detailed explanations of any of these algorithms, but he does discuss the idea behind Fortune’s algorithm. Today we will discuss the divide-and-conquer algorithm. This algorithm is presented in Mulmuley, Section 2.8.4.

**Divide-and-conquer algorithm:** The divide-and-conquer approach works like most standard geometric divide-and-conquer algorithms. We split the points according to \( x \)-coordinates into 2 roughly equal sized groups (e.g. by presorting the points by \( x \)-coordinate and selecting medians). We compute the Voronoi diagram of the left side, and the Voronoi diagram of the right side. Note that since each diagram alone covers the entire plane, these two diagrams overlap. We then merge the resulting diagrams into a single diagram.

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The merging step is where all the work is done. Observe that every point in the plane lies within two Voronoi polygons, one in $VD(L)$ and one in $VD(R)$. We need to resolve this overlap, by separating overlapping polygons. Let $V(l_0)$ be the Voronoi polygon for a point from the left side, and let $V(r_0)$ be the Voronoi polygon for a point on the right side, and suppose these polygons overlap one another. Observe that if we insert the bisector between $l_0$ and $r_0$, and through away the portions of the polygons that lie on the “wrong” side of the bisector, we resolve the overlap. If we do this for every pair of overlapping Voronoi polygons, we get the final Voronoi diagram. This is illustrated in the figure below.

![Diagram](image)

Figure 26.1: Merging Voronoi diagrams.

The union of these bisectors that separate the left Voronoi diagram from the right Voronoi diagram is called the contour. A point is on the contour if and only if it is equidistant from 2 points in $S$, one in $L$ and one in $R$.

1. Presort the points by $x$-coordinate (this is done once).
2. Split the point set $S$ by a vertical line into two subsets $L$ and $R$ of roughly equal size.
3. Compute $VD(L)$ and $VD(R)$ recursively. (These diagrams overlap one another.)
4. Merge the two diagrams into a single diagram, by computing the contour and discarding the portion of the $VD(L)$ that is to the right of the contour, and the portion of $VD(R)$ that is to the left of the contour.

Assuming we can implement step (3) in $O(n)$ time (where $n$ is the size of the remaining point set) then the running time will be defined by the familiar recurrence

$$T(n) = 2T(n/2) + n,$$

which we know solves to $O(n \log n)$.
Computing the contour: What makes the divide-and-conquer algorithm somewhat tricky is the task of computing the contour. Before giving an algorithm to compute the contour, let us make some observations about its geometric structure. Let us make the usual simplifying assumptions that no 4 points are cocircular.

Lemma: The contour consists of a single polygonal curve (whose first and last edges are semiinfinite) which is monotone with respect to the $y$-axis.

Proof: A detailed proof is a real hassle. Here are the main ideas, though. The contour separates the plane into two regions, those points whose nearest neighbor lies in $L$ from those points whose nearest neighbor lies in $R$. Because the contour locally consists of points that are equidistant from 2 points, it is formed from pieces that are perpendicular bisectors, with one point from $L$ and the other point from $R$. Thus, it is a piecewise polygonal curve. Because no 4 points are cocircular, it follows that all vertices in the Voronoi diagram can have degree at most 3. However, because the contour separates the plane into only 2 types of regions, it can contain only vertices of degree 2. Thus it can consist only of the disjoint union of closed curves (actually this never happens, as we will see) and unbounded curves. Observe that if we orient the contour counterclockwise with respect to each point in $R$ (clockwise with respect to each point in $L$), then each segment must be directed in the $-y$ directions, because $L$ and $R$ are separated by a vertical line. Thus, the contour contains no horizontal cusps. This implies that the contour cannot contain any closed curves, and hence contains only vertically monotone unbounded curves. Also, this orientability also implies that there is only one such curve.

Lemma: The topmost (bottommost) edge of the contour is the perpendicular bisector for the two points forming the upper (lower) tangent of the left hull and the right hull.

Proof: This follows from the fact that the vertices of the hull correspond to unbounded Voronoi polygons, and hence upper and lower tangents correspond to unbounded edges of the contour.

These last two theorem suggest the general approach. We start by computing the upper tangent, which we know can be done in linear time (once we know the left and right hulls, or by prune and search). Then, we start tracing the contour from top to bottom. When we are in Voronoi polygons $V(l_0)$ and $V(r_0)$ we trace the bisector between $l_0$ and $r_0$ in the negative $y$-direction until its first contact with the boundaries of one of these polygons. Suppose that we hit the boundary of $V(l_0)$ first. Assuming that we use a good data structure for the Voronoi diagram (e.g. quad-edge data structure) we can determine the point $l_1$ lying on the other side of this edge in the left Voronoi diagram. We continue following the contour by tracing the bisector of $l_0$ and $r_0$.

However, in order to insure efficiency, we must be careful in how we determine where the bisector hits the edge of the polygon. Consider the figure shown below. We start tracing the contour between $l_0$ and $r_0$. By walking along the boundary of $V(l_0)$ we can determine the edge that the contour would hit first. This can be done in time proportional to the number of edges in $V(l_0)$ (which can be as large as $O(n)$). However, we discover that before the contour hits the boundary of $V(l_0)$ it hits the boundary of $V(r_0)$. We find the new point $r_1$ and now trace the bisector between $l_0$ and $r_1$. Again we can compute the intersection with the boundary of $V(l_0)$ in time proportional to its size. However the contour hits the boundary of $V(r_1)$ first, and so we go on to $r_2$. As can be seen, if we are not smart, we can rescan the boundary of $V(l_0)$ over and over again, until the contour finally hits the boundary. If we do this $O(n)$ times, and the boundary of $V(l_0)$ is $O(n)$, then we are stuck with $O(n^2)$ time to trace the contour.
We have to avoid this repeated rescanning. However, there is a way to scan the boundary of each Voronoi polygon at most once. Observe that as we walk along the contour, each time we stay in the same polygon $V(l_0)$, we are adding another edge onto its Voronoi polygon. Because the Voronoi polygon is convex, we know that the edges we are creating turn consistently in the same direction (clockwise for points on the left, and counterclockwise for points on the right). To test for intersections between the contour and the current Voronoi polygon, we trace the boundary of the polygon clockwise for polygons on the left side, and counterclockwise for polygons on the right side. Whenever the contour changes direction, we continue the scan from the point that we left off. In this way, we know that we will never need to rescan the same edge of any Voronoi polygon more than once.
27 Delaunay Triangulations and Convex Hulls

Reading: O’Rourke 5.7 and 5.8.

Delaunay Triangulations and Convex Hulls: At first, Delaunay triangulations and convex hulls appear to be quite different structures, one is based on metric properties (distances) and the other on affine properties (collinearity, coplanarity). Today we show that it is possible to convert the problem of computing a Delaunay triangulation in dimension \(d\) to that of computing a convex hull in dimension \(d + 1\). Thus, there is a remarkable relationship between these two structures.

We will demonstrate the connection in dimension 2 (by computing a convex hull in dimension 3). Some of this may be hard to visualize, but see O’Rourke for illustrations. (You can also reason by analogy in one lower dimension of Delaunay triangulations in 1-d and convex hulls in 2-d, but the real complexities of the structures are not really apparent in this case.)

The connection between the two structures is the *paraboloid* \(z = x^2 + y^2\). Observe that this equation defines a surface whose vertical cross sections (constant \(x\) or constant \(y\)) are parabolas, and whose horizontal cross sections (constant \(z\)) are circles. For each point in the plane, \((x, y)\), the *vertical projection* of this point onto this paraboloid is \((x, y, x^2 + y^2)\) in 3-space. Given a set of points \(S\) in the plane, let \(S'\) denote the projection of every point in \(S\) onto this paraboloid. Consider the *lower convex hull* of \(S'\). This is the portion of the convex hull of \(S'\) which is visible to a viewer standing at \(z = -\infty\). We claim that if we take the lower convex hull of \(S'\), and project it back onto the plane, then we get the Delaunay triangulation of \(S\). In particular, let \(p, q, r \in S\), and let \(p', q', r'\) denote the projections of these points onto the paraboloid. Then \(p'q'r'\) define a *face* of the lower convex hull of \(S'\) if and only if \(\triangle pqr\) is a triangle of the Delaunay triangulation of \(S\). The process is illustrated in the following figure.

![Figure 27.1: Delaunay triangulations and convex hull.](image)

The question is, why does this work? To see why, we need to establish the connection between the triangles of the Delaunay triangulation and the faces of the convex hull of transformed points. In particular, recall that

**Delaunay condition:** Three points \(p, q, r \in S\) form a Delaunay triangle if and only if the circumcircle of these points contains no other point of \(S\).

**Convex hull condition:** Three points \(p', q', r' \in S'\) form a face of the convex hull of \(S'\) if and only if the plane passing through \(p', q',\) and \(r'\) has all the points of \(S'\) lying to one side.
Clearly, the connection we need to establish is between the emptiness of circumcircles in the plane and the emptiness of halfspaces in 3-space. We will prove the following claim.

**Lemma:** Consider 4 distinct points $p, q, r, s$ in the plane, and let $p', q', r', s'$ be their respective projections onto the paraboloid, $z = x^2 + y^2$. The point $s$ lies within the circumcircle of $p, q, r$ if and only if $s'$ lies on the lower side of the plane passing through $p', q', r'$.

To prove the lemma, first consider an arbitrary (nonvertical) plane in 3-space, which we assume is tangent to the paraboloid above some point $(a, b)$ in the plane. To determine the equation of this tangent plane, we take derivatives of the equation $z = x^2 + y^2$ with respect to $x$ and $y$ giving

$$\frac{\partial z}{\partial x} = 2x, \quad \frac{\partial z}{\partial y} = 2y.$$ 

At the point $(a, b, a^2 + b^2)$ these evaluate to $2a$ and $2b$. It follows that the plane passing through these point has the form

$$z = 2ax + 2by + \gamma.$$ 

To solve for $\gamma$ we know that the plane passes through $(a, b, a^2 + b^2)$ so we solve giving

$$a^2 + b^2 = 2a \cdot a + 2b \cdot b + \gamma,$$

Implying that $\gamma = -(a^2 + b^2)$. Thus the plane equation is

$$z = 2ax + 2by - (a^2 + b^2).$$

If we shift the plane upwards by some positive amount $r^2$ we get the plane

$$z = 2ax + 2by - (a^2 + b^2) + r^2.$$ 

How does this plane intersect the paraboloid? Since the paraboloid is defined by $z = x^2 + y^2$ we can eliminate $z$ giving

$$x^2 + y^2 = 2ax + 2by - (a^2 + b^2) + r^2,$$

which after some simple rearrangements is equal to

$$(x - a)^2 + (y - b)^2 = r^2.$$ 

This is just a circle. Thus, we have shown that the intersection of a plane with the paraboloid produces a space curve (which turns out to be an ellipse), which when projected back onto the $(x, y)$-coordinate plane is a circle centered at $(a, b)$.

Thus, we conclude that the intersection of an arbitrary lower halfspace with the paraboloid, when projected onto the $(x, y)$-plane is the interior of a circle. Going back to the lemma, when we project the points $p, q, r$ onto the paraboloid, the projected points $p', q', r'$ define a plane. Since $p', q', r'$, lie at the intersection of the plane and paraboloid, the original points $p, q, r$ lie on the projected circle. Thus this circle is the (unique) circumcircle passing through these $p, q, r$. Thus, the point $s$ lies within this circumcircle, if and only if its projection $s'$ onto the paraboloid lies within the lower halfspace of the plane passing through $p, q, r$.

Now we can prove the main result.
Theorem: Given a set of points $S$ in the plane (assume no 4 are cocircular), and given 3 points $p, q, r \in S$, the triangle $\triangle pqr$ is a triangle of the Delaunay triangulation of $S$ if and only if triangle $\triangle p'q'r'$ is a face of the lower convex hull of the projected set $S'$.

From the definition of Delaunay triangulations we know that $\triangle pqr$ is in the Delaunay triangulation if and only if there is no point $s \in S$ that lies within the circumcircle of $pqr$. From the previous lemma this is equivalent to saying that there is no point $s'$ that lies in the lower convex hull of $S'$, which is equivalent to saying that $p'q'r'$ is a face of the lower convex hull. This completes the proof.

In order to test whether a point $s$ lies within the circumcircle defined by $p, q, r$, it suffices to test whether $s'$ lies within the lower halfspace of the plane passing through $p', q', r'$. If we assume that $p, q, r$ are oriented counterclockwise in the plane this reduces to determining whether the quadruple $p', q', r', s'$ is positively oriented, or equivalently whether $s$ lies to the left of the oriented circle passing through $p, q, r$.

This leads to the incircle test we presented last time.

$$
in(p, q, r, s) = \det \begin{pmatrix} p_x & p_y & p_x^2 + p_y^2 & 1 \\ q_x & q_y & q_x^2 + q_y^2 & 1 \\ r_x & r_y & r_x^2 + r_y^2 & 1 \\ s_x & s_y & s_x^2 + s_y^2 & 1 \end{pmatrix} > 0.$$

Voronoi Diagrams and Upper Envelopes: We know that Voronoi diagrams and Delaunay triangulations are dual geometric structures. We have also seen (informally) that there is a dual relationship between points and lines in the plane, and in general, points and planes in 3-space. From this latter connection we argued that the problems of computing convex hulls of point sets and computing the intersection of halfspaces are somehow “dual” to one another. It turns out that these two notions of duality, are (not surprisingly) interrelated. In particular, in the same way that the Delaunay triangulation of points in the plane can be transformed to computing a convex hull in 3-space, it turns out that the Voronoi diagram of points in the plane can be transformed into computing the intersection of halfspaces in 3-space.

Here is how we do this. For each point $p = (a, b)$ in the plane, recall the tangent plane to the paraboloid above this point, given by the equation

$$z = 2ax + 2by - (a^2 + b^2).$$

Define $H^+(p)$ to be the set of points that are above this halfplane, that is, $H^+(p) = \{(x, y, z) \mid z \geq 2ax + 2by - (a^2 + b^2)\}$. Let $S = \{p_1, p_2, \ldots, p_n\}$ be a set of points. Consider the intersection of the
halfspaces $H^+(p_i)$. This is also called the upper envelope of these halfspaces. The upper envelope is an (unbounded) convex polyhedron. If you project the edges of this upper envelope down into the plane, it turns out that you get the Voronoi diagram of the points.

**Theorem:** Given a set of points $S$ in the plane (assume no 4 are cocircular), let $H$ denote the set of upper halfspaces defined by the previous transformation. Then the Voronoi diagram of $H$ is equal to the projection onto the $(x, y)$-plane of the 1-skeleton of the convex polyhedron which is formed from the intersection of halfspaces of $S'$.

![Figure 27.3: Intersection of halfspaces.](image)

It is hard to visualize this surface, but it is not hard to show why this is so. Suppose we have 2 points in the plane, $p = (a, b)$ and $q = (c, d)$. The corresponding planes are:

$$z = 2ax + 2by - (a^2 + b^2) \quad \text{and} \quad z = 2cx + 2dy - (c^2 + d^2).$$

If we determine the intersection of the corresponding planes and project onto the $(x, y)$-coordinate plane (by eliminating $z$ from these equations) we get

$$x(2a - 2c) + y(2b - 2d) = (a^2 - c^2) + (b^2 - d^2).$$

We claim that this is the perpendicular bisector between $(a, b)$ and $(c, d)$. To see this, observe that it passes through the midpoint $((a + c)/2, (b + d)/2)$ between the two points since

$$\frac{a + c}{2}(2a - 2c) + \frac{b + d}{2}(2b - 2d) = (a^2 - c^2) + (b^2 - d^2).$$

and, its slope is $-(a - c)/(b - d)$, which is the negative reciprocal of the line segment from $(a, b)$ to $(c, d)$. From this it can be shown that the intersection of the upper halfspaces defines a polyhedron whose edges project onto the Voronoi diagram edges.
28 Topological Plane Sweep


Topological Plane Sweep: In the last two lectures we have introduced arrangements of lines and geometric duality as important tools in solving geometric problems on lines and points. Today give an efficient algorithm for sweeping an arrangement of lines.

As we will see, many problems in computational geometry can be solved by applying line-sweep to an arrangement of lines. Since the arrangement has size $O(n^2)$, and since there are $O(n^2)$ events to be processed, each involving an $O(\log n)$ heap deletion, this typically leads to algorithms running in $O(n^2 \log n)$ time, using $O(n^2)$ space. It is natural to ask whether we can dispense with the additional $O(\log n)$ factor in running time, and whether we need all of $O(n^2)$ space (since in theory we only need access to the current $O(n)$ contents of the sweep line).

We discuss a variation of plane sweep called topological plane sweep. This method runs in $O(n^2)$ time, and uses only $O(n)$ space (by essentially constructing only the portion of the arrangement that we need at any point). Although it may appear to be somewhat sophisticated, it can be implemented quite efficiently, and is claimed to outperform conventional plane sweep on arrangements of any significant size (e.g. over 20 lines).

Cuts and topological lines: The algorithm is called topological plane sweep because we do not sweep a straight vertical line through the arrangement, but rather we sweep a curved topological line that has the essential properties of a vertical sweep line in the sense that this line intersects each line of the arrangement exactly once. The notion of a topological line is an intuitive one, but it can be made formal in the form of something called a cut. Recall that the faces of the arrangement are convex polygons (possibly unbounded). (Assuming no vertical lines) the edges incident to each face can naturally be partitioned into the edges that are above the face, and those that are below the face. Define a cut in an arrangement to be a sequence of edges $c_1, c_2, \ldots, c_n$, in the arrangement, one taken from each line of the arrangement, such that for $1 \leq i \leq n - 1$, $c_i$ and $c_{i+1}$ are incident to the same face of the arrangement, and $c_i$ is above the face and $c_{i+1}$ is below the face. An example of a topological line and the associated cut is shown below.

![Figure 28.1: Topological line and associated cut.](image)

The topological plane sweep starts at the leftmost cut of the arrangement. This consists of all the left-unbounded edges of the arrangement. Observe that this cut can be computed in $O(n \log n)$ time, because the lines intersect the cut in inverse order of slope. The topological sweep line will sweep to the right until we come to the rightmost cut, which consists all of the right-unbounded edges of
the arrangement. The sweep line advances by a series of what are called \textit{elementary steps}. In an elementary steps, we find two consecutive edges on the cut that meet at a vertex of the arrangement (we will discuss later how to determine this), and push the topological sweep line through this vertex. Observe that on doing so these two lines swap in their order along the sweep line. This is shown below.

![Elementary step](image)

\textbf{Figure 28.2: Elementary step.}

It is not hard to show that an elementary step is always possible, since for any cut (other than the rightmost cut) there must be two consecutive edges with a common right endpoint. In particular, consider the edge of the cut whose right endpoint has the smallest \( x \)-coordinate. It is not hard to show that this endpoint will always allow an elementary step. Unfortunately, determining this vertex would require at least \( O(\log n) \) time (if we stored these endpoints in a heap, sorted by \( x \)-coordinate), and we want to perform each elementary step in \( O(1) \) time. Hence, we will need to find some other method for finding elementary steps.

**Upper and Lower Horizon Trees:** To find elementary steps, we introduce two simple data structures, the \textit{upper horizon tree} (UHT) and the \textit{lower horizon tree} (LHT). To construct the upper horizon tree, trace each edge of the cut to the right. When two edges meet, keep only the one with the higher slope, and continue tracing it to the right. The lower horizon tree is defined symmetrically. There is one little problem in these definitions in the sense that these trees need not be connected (forming a forest of trees) but this can be fixed conceptually at least by the addition of a vertical line at \( x = +\infty \). For the upper horizon we think of its slope as being \( +\infty \) and for the lower horizon we think of its slope as being \(-\infty\). Note that we consider the left endpoints of the edges of the cut as not belonging to the trees, since otherwise they would not be trees. It is not hard to show that with these modifications, these are indeed trees. Each edge of the cut defines exactly one line segment in each tree. An example is shown below.

![Upper and lower horizon trees](image)

\textbf{Figure 28.3: Upper and lower horizon trees.}

The important things about the UHT and LHT is that they give us an easy way to determine the right endpoints of the edges on the cut. Observe that for each edge in the cut, its right endpoint results from a line of smaller slope intersecting it from above (as we trace it from left to right) or from a line of larger slope intersecting it from below. It is easy to verify that the UHT and LHT determine the first such intersecting line of each type, respectively. It follows that if we intersect the two trees, then the segments they share in common correspond exactly to the edges of the cut. Thus, by knowing the UHT and LHT, we know where are the right endpoints are, and from this we can determine easily
which pairs of consecutive edges share a common right endpoint, and from this we can determine all
the elementary steps that are legal. We store all the legal steps in a stack (or queue, or any list is fine),
and extract them one by one.

The sweep algorithm: Here is an overview of the topological plane sweep.

(1) Input the lines and sort by slope. Let \( C \) be the initial (leftmost) cut, a list of lines in decreasing
order of slope.

(2) Create the initial UHT incrementally by inserting lines in decreasing order of slope. Create the
initial LHT incrementally by inserting line in increasing order of slope. (More on this later.)

(3) By consulting the LHT and UHT, determine the right endpoints of all the edges of the initial cut,
and for all pairs of consecutive lines \((l_i, l_{i+1})\) sharing a common right endpoint, store this pair
in stack \( S \).

(4) Repeat the following elementary step until the stack is empty (implying that we have arrived at
the rightmost cut).

(a) Pop the pair \((l_i, l_{i+1})\) from the top of the stack \( S \).

(b) Swap these lines within \( C \), the cut (we assume that each line keeps track of its position in
the cut).

(c) Update the horizon trees. (More on this later.)

(d) Consulting the changed entries in the horizon tree, determine whether there are any new cut
edges sharing right endpoints, and if so push them on the stack \( S \).

The important unfinished business is to show that we can build the initial UHT and LHT in \( O(n) \) time,
and to show that, for each elementary step, we can update these trees and all other relevant information
in \( O(1) \) amortized time. By amortized time we mean that, even though a single elementary step can
take more than \( O(1) \) time, the total time needed to perform all \( O(n^2) \) elementary steps is \( O(n^2) \), and
hence the average time for each step is \( O(1) \).

This is done by an adaptation of the same incremental “face walking” technique we used in the incre-
mental construction of line arrangements. Let’s consider just the UHT, since the LHT is symmetric.
To create the initial (leftmost) UHT we insert the lines one by one in decreasing order of slope. Ob-
serve that as each new line is inserted it will start above all of the current lines. The uppermost face
of the current UHT consists of a convex polygonal chain, see the figure below left. As we trace the
newly inserted line from left to right, there will be some point at which it first hits this upper chain
of the current UHT. By walking along the chain from left to right, we can determine this intersection
point. Each segment that is walked over is never visited again by this initialization process (because
it is no longer part of the upper chain), and since the initial UHT has a total of \( O(n) \) segments, this
implies that the total time spent in walking is \( O(n) \). Thus, after the \( O(n \log n) \) time for sorting the
segments, the initial UHT tree can be built in \( O(n) \) additional time.

Next we show how to update the UHT after an elementary step. The process is quite similar, as shown
in the figure right. Let \( v \) be the vertex of the arrangement which is passed over in the sweep step. As
we pass over \( v \), the two edges swap positions along the sweep line. The new lower edge, call it \( l \),
which had been cut off of the UHT by the previous lower edge, now must be reentered into the tree.
We extend \( l \) to the left until it contacts an edge of the UHT. At its first contact, it will terminate (and
this is the only change to be made to the UHT). In order to find this contact, we start with the edge
immediately below \( l \) the current cut. We traverse the face of the UHT in counterclockwise order, until finding the edge that this line intersects. Observe that we must eventually find such an edge because \( l \) has a lower slope than the other edge intersecting at \( v \), and this edge lies in the same face.

**Analysis:** A careful analysis of the running time can be performed using the same amortization proof (based on pebble counting) that was used in the analysis of the incremental algorithm. We will not give the proof in full detail. Observe that because we maintain the set of legal elementary steps in a stack (as opposed to a heap as would be needed for standard plane sweep), we can advance to the next elementary step in \( O(1) \) time. The only part of the elementary step that requires more than constant time is the update operations for the UHT and LHT. However, we claim that the total time spent updating these trees is \( O(n^2) \). The argument is that when we are tracing the edges (as shown in the previous figure) we are “essentially” traversing the edges in the zone for \( L \) in the arrangement. (This is not quite true, because there are edges above \( l \) in the arrangement, which have been cut out of the upper tree, but the claim is that their absence cannot increase the complexity of this operation, only decrease it. However, a careful proof needs to take this into account.) Since the zone of each line in the arrangement has complexity \( O(n) \), all \( n \) zones have total complexity \( O(n^2) \). Thus, the total time spent in updating the UHT and LHT trees is \( O(n^2) \).
29 Ham-Sandwich Cuts

Reading: This material is not covered in our book.

Ham Sandwich Cuts of Linearly Separated Point Sets: We are given \(n\) red points \(A\), and \(m\) blue points \(B\), and we want to compute a single line that simultaneously bisects both sets. (If the cardinality of either set is odd, then the line passes through one of the points of the set.) We make the simplifying assumption that the sets are separated by a line. (This assumption makes the problem much simpler to solve, but the general case can still be solved in \(O(n^2)\) time using arrangements.)

To make matters even simpler we assume that the points have been translated and rotated so this line is the \(y\)-axis. Thus all the red points (set \(A\)) have positive \(x\)-coordinates, and hence their dual lines have positive slopes, whereas all the blue points (set \(B\)) have negative \(x\)-coordinates, and hence their dual lines have negative slopes. As long as we are simplifying things, let’s make one last simplification, that both sets have an odd number of points. This is not difficult to get around, but makes the pictures a little easier to understand.

Consider one of the sets, say \(A\). Observe that for each slope there exists one way to bisect the points. In particular, if we start a line with this slope at positive infinity, so that all the points lie beneath it, and drop in downwards, eventually we will arrive at a unique placement where there are exactly \((n - 1)/2\) points above the line, one point lying on the line, and \((n - 1)/2\) points below the line (assuming no two points share this slope). This line is called the median line for this slope.

What is the dual of this median line? If we dualize the points using the standard dual transformation: \(D(a, b) : y = ax - b\), then we get \(n\) lines in the plane. By starting a line with a given slope above the points and translating it downwards, in the dual plane we moving a point from \(-\infty\) upwards in a vertical line. Each time the line passes a point in the primal plane, the vertically moving point crosses a line in the dual plane. When the translating line hits the median point, in the dual plane the moving point will hit a dual line such that there are exactly \((n - 1)/2\) dual lines above this point and \((n - 1)/2\) dual lines below this point. We define a point to be at level \(k\), \(L_k\), in an arrangement if there are at most \(k\) – 1 lines above this point and at most \(n - k\) lines below this point. The median level in an arrangement of \(n\) lines is defined to be the \([ (n - 1)/2 ]\)-th level in the arrangement. This is shown as \(M(A)\) in the following figure on the left.

![Figure 29.1: Ham sandwich: Dual formulation.](image)

Thus, the set of bisecting lines for set \(A\) in dual form consists of a polygonal curve. Because this curve is formed from edges of the dual lines in \(A\), and because all lines in \(A\) have positive slope, this curve is monotonically increasing. Similarly, the median for \(B\), \(M(B)\), is a polygonal curve which is monotonically decreasing. It follows that \(A\) and \(B\) must intersect at a unique point. The dual of this point is a line that bisects both sets.
We could compute the intersection of these two curves by a simultaneous topological plane sweep of both arrangements. However, it turns out that it is possible to do much better, and in fact the problem can be solved in $O(n + m)$ time. Since the algorithm is rather complicated, I will not describe the details, but here are the essential ideas. The algorithm operates by prune and search. In $O(n + m)$ time we will generate a hypothesis for where the ham sandwich point is in the dual plane, and if we are wrong, we will succeed in throwing away a constant fraction of the lines from future consideration.

First observe that for any vertical line in the dual plane, it is possible to determine in $O(n + m)$ time whether this line lies to the left or the right of the intersection point of the median levels, $M(A)$ and $M(B)$. This can be done by computing the intersection of the dual lines of $A$ with this line, and computing their median in $O(n)$ time, and computing the intersection of the dual lines of $B$ with this line and computing their median in $O(m)$ time. If $A$’s median lies below $B$’s median, then we are to the left of the ham sandwich dual point, and otherwise we are to the right of the ham sandwich dual point. It turns out that with a little more work, it is possible to determine in $O(n + m)$ time whether the ham sandwich point lies to the right or left of a line of arbitrary slope. The trick is to use prune and search. We find two lines $L_1$ and $L_2$ in the dual plane (by a careful procedure that I will not describe). These two lines define four quadrants in the plane. By determining which side of each line the ham sandwich point lies, we know that we can throw away any line that does not intersect this quadrant from further consideration. It turns out that by a judicious choice of $L_1$ and $L_2$, we can guarantee that a fraction of at least $(n + m)/8$ lines can be thrown away by this process. We recurse on the remaining lines. By the same sort of analysis we made in the Kirkpatrick and Seidel prune and search algorithm for upper tangents, it follows that in $O(n + m)$ time we will find the ham sandwich point.
Shortest Paths and Visibility Graphs

Reading: The material on visibility graphs is taken roughly from Chapter 15, but we will present slightly more efficient variant of the one that appears in this chapter.

Shortest paths: We are given a set of \( n \) disjoint polygonal obstacles in the plane, and two points \( s \) and \( t \) that lie outside of the obstacles. The problem is to determine the shortest path from \( s \) to \( t \) that avoids the interiors of the obstacles. (It may travel along the edges or pass through the vertices of the obstacles.) The complement of the interior of the obstacles is called free space. We want to find the shortest path that is constrained to lie entirely in free space.

Today we consider a simple (but perhaps not the most efficient) way to solve this problem. We assume that we measure lengths in terms of Euclidean distances. How do we measure paths lengths for curved paths? Luckily, we do not have to, because we claim that the shortest path will always be a polygonal curve.

Claim: The shortest path between any two points that avoids a set of polygonal obstacles is a polygonal curve, whose vertices are either vertices of the obstacles or the points \( s \) and \( t \).

Proof: We show that any path \( \pi \) that violates these conditions can be replaced by a slightly shorter path from \( s \) to \( t \). Since the obstacles are polygonal, if the path were not a polygonal curve, then there must be some point \( p \) in the interior of free space, such that the path passing through \( p \) is not locally a line segment. If we consider any small neighborhood about \( p \) (small enough to not contain \( s \) or \( t \) or any part of any obstacle), then since the shortest path is not locally straight, we can shorten it slightly by replacing this curved segment by a straight line segment jointing one end to the other. Thus, \( \pi \) is not shortest, a contradiction.

Thus \( \pi \) is a polygonal path. Suppose that it contained a vertex \( v \) that was not an obstacle vertex. Again we consider a small neighborhood about \( v \) that contains no part of any obstacle. We can shorten the path, as above, implying that \( \pi \) is not a shortest path.

From this it follows that the edges that constitute the shortest path must travel between \( s \) and \( t \) and vertices of the obstacles. Each of these edges must have the property that it does not intersect the interior of any obstacle, implying that the endpoints must be visible to each other. More formally, we say that two points \( p \) and \( q \) are mutually visible if the open line segment joining them does not intersect the interior of any obstacle. By this definition, the two endpoints of an obstacle edge are not mutually visible, so we will explicitly allow for this case in the definition below.

Definition: The visibility graph of \( s \) and \( t \) and the obstacle set is a graph whose vertices are \( s \) and \( t \) the obstacle vertices, and vertices \( v \) and \( w \) are joined by an edge if \( v \) and \( w \) are either mutually visible or if \( (v, w) \) is an edge of some obstacle.

It follows from the above claim that the shortest path can be computed by first computing the visibility graph and labeling each edge with its Euclidean length, and then computing the shortest path by, say, Dijkstra’s algorithm (see CLR). Note that the visibility graph is not planar, and hence may consist of \( \Omega(n^2) \) edges. Also note that, even if the input points have integer coordinates, in order to compute distances we need to compute square roots, and then sums of square roots. This can be approximated by floating point computations. (If exactness is important, this can really be a problem, because
there is no known polynomial time procedure for performing arithmetic with arbitrary square roots of integers.)

Computing the Visibility Graph: We give an $O(n^2)$ procedure for constructing the visibility graph of $n$ line segments in the plane. The more general task of computing the visibility graph of an arbitrary set of polygonal obstacles is a very easy generalization. In this context, two vertices are visible if the line segment joining them does not intersect any of the obstacle line segments. However, we allow each line segment to contribute itself as an edge in the visibility graph. We will make the general position assumption that no three vertices are collinear, but this is not hard to handle with some care. The algorithm is not output sensitive. If $k$ denotes the number of edges in the visibility graph, then an $O(n \log n + k)$ algorithm does exist, but it is quite complicated.

The text gives an $O(n^2 \log n)$ time algorithm. We will give an $O(n^2)$ time algorithm. Both algorithms are based on the same concept, namely that of performing an angular sweep around each vertex. The text’s algorithm operates by doing this sweep one vertex at a time. Our algorithm does the sweep for all vertices simultaneously. We use the fact (given in the lecture on arrangements) that this angular sort can be performed for all vertices in $O(n^2)$ time. If we build the entire arrangement, this sorting algorithm will involve $O(n^2)$ space. However it can be implemented in $O(n)$ space using an algorithm called topological plane sweep. Topological plane sweep provides a way to sweep an arrangement of lines using a “flexible” sweeping line. Because events do not need to sorted, we can avoid the $O(\log n)$ factor, which would otherwise be needed to maintain the priority queue.

Here is a high-level intuitive view of the algorithm. First, recall the algorithm for computing trapezoidal maps. We shoot a bullet up and down from every vertex until it hits its first line segment. This implicitly gives us the vertical visibility relationships between vertices and segments. Now, we imagine that angle $\theta$ continuously sweeps out all slopes from $-\infty$ to $+\infty$. Imagine that all the bullet lines attached to all the vertices begin to turn slowly counterclockwise. If we play the mind experiment of visualizing the rotation of these bullet paths, the question is what are the significant event points, and what happens with each event? As the sweep proceeds, we will eventually determine everything that is visible from every vertex in every direction. Thus, it should be an easy matter to piece together the edges of the visibility graph as we go.

Let us consider this “multiple angular sweep” in greater detail.

It is useful to view the problem both in its primal and dual form. For each of the $2n$ segment endpoints $v = (v_a, v_b)$, we consider its dual line $v^* : y = v_a x - v_b$. Observe that a significant event occurs whenever a bullet path in the primal plane jumps from one line segment to another. This occurs when $\theta$ reaches the slope of the line joining two visible endpoints $v$ and $w$. Unfortunately, it is somewhat complicated to keep track of which endpoints are visible and which are not (although if we could do so it would lead to a more efficient algorithm). Instead we will take events to be all angles $\theta$ between two endpoints, whether they are visible or not. By duality, the slope of such an event will correspond
to the $a$-coordinate of the intersection of dual lines $v^*$ and $w^*$ in the dual arrangement. (Convince yourself of this.) Thus, by sweeping the arrangement of the $2n$ dual lines from left-to-right, we will enumerate all the slope events in angular order.

Next let’s consider what happens at each event point. Consider the state of the angular sweep algorithm for some slope $\theta$. For each vertex $v$, there are two bullet paths emanating from $v$ along the line with slope $\theta$. Call one the forward bullet path and the other the backward bullet path. Let $f(v)$ and $b(v)$ denote the line segments that these bullet paths hit, respectively. If either path does not hit any segment then we store a special null value. As $\theta$ varies the following events can occur. Assuming (through symbolic perturbation) that each slope is determined by exactly two lines, whenever we arrive at an events slope $\theta$ there are exactly two vertices $v$ and $w$ that are involved. Here are the possible scenarios:

**Same segment**: If $v$ and $w$ are endpoints of the same segment, then they are visible, and we add the edge $(v, w)$ to the visibility graph.

**Invisible**: Consider the distance from $v$ to $w$. First, determine whether $w$ lies on the same side as $f(v)$ or $b(v)$. For the remainder, assume that it is $f(v)$. (The case of $b(v)$ is symmetrical). Compute the contact point of the bullet path shot from $v$ in direction $\theta$ with segment $f(v)$. If this path hits $f(v)$ strictly before $w$, then we know that $w$ is not visible to $v$, and so this is a “non-event”.

**Segment entry**: Consider the segment that is incident to $w$. Either the sweep is just about to enter this segment or is just leaving it. If we are entering the segment, then we set $f(v)$ to this segment.

**Segment exit**: If we are just leaving this segment, then the bullet path will need to shoot out and find the next segment that it hits. Normally this would require some searching. (In particular,
this is one of the reasons that the text’s algorithm has the extra $O(\log n)$ factor—to perform this search.) However, we claim that the answer is available to us in $O(1)$ time.

In particular, since we are sweeping over $w$ at the same time that we are sweeping over $v$. Thus we know that the bullet extension from $w$ hits $f(w)$. All we need to do is to set $f(v) = f(w)$.

This is a pretty simple algorithm (although there are a number of cases). The only information that we need to keep track of is (1) a priority queue for the events, and (2) the $f(v)$ and $b(v)$ pointers for each vertex $v$. The priority queue is not stored explicitly. Instead it is available from the line arrangement of the duals of the line segment vertices. By performing a topological sweep of the arrangement, we can process all of these events in $O(n^2)$ time.
31 Motion Planning

Reading: Chapt 13 in the 4M’s.

Motion planning: Last time we considered the problem of computing the shortest path of a point in space around a set of obstacles. Today we will study a much more general approach to the more general problem of how to plan the motion of one or more robots, each with potentially many degrees of freedom in terms of its movement and perhaps having articulated joints.

Work Space and Configuration Space: The environment in which the robot operates is called its work space, which consists of a set of obstacles that the robot is not allowed to intersect. We assume that the work space is static, that is, the obstacles do not move. We also assume that a complete geometric description of the work space is available to us.

For our purposes, a robot will be modeled by two main elements. The first is a configuration, which is a finite sequence of values that fully specifies the position of the robot. The second element is the robot’s geometric shape description. Combined these two element fully define the robot’s exact position and shape in space. For example, suppose that the robot is a 2-dimensional polygon that can translate and rotate in the plane. Its configuration may be described by the \((x, y)\) coordinates of some reference point for the robot, and an angle \(\theta\) that describes its orientation. Its geometric information would include its shape (say at some canonical position), given, say, as a simple polygon. Given its geometric description and a configuration \((x, y, \theta)\), this uniquely determines the exact position \(R(x, y, \theta)\) of this robot in the plane. Thus, the position of the robot can be identified with a point in the robot’s configuration space.

A more complex example would be an articulated arm consisting of a set of links, connected to one another by a set of revolute joints. The configuration of such a robot would consist of a vector of joint angles. The geometric description would probably consist of a geometric representation of the links. Given a sequence of joint angles, the exact shape of the robot could be derived by combining this configuration information with its geometric description. For example, a typical 3-dimensional industrial robot has six joints, and hence its configuration can be thought of as a point in a 6-dimensional space. Why six? Generally, there are three degrees of freedom needed to specify a location in 3-space, and 3 more degrees of freedom needed to specify the direction and orientation of the robot’s end manipulator.

Given a point \(p\) in the robot’s configuration space, let \(R(p)\) denote the placement of the robot at this configuration. The figure below illustrates this in the case of the planar robot defined above.

Because of limitations on the robot’s physical structure and the obstacles, not every point in configuration space corresponds to a legal placement of the robot. Any configuration which is illegal in that it
causes the robot to intersect one of the obstacles is called a *forbidden configuration*. The set of all forbidden configurations is denoted $C_{\text{forb}}(R, S)$, and all other placements are called *free configurations*, and the set of these configurations is denoted $C_{\text{free}}(R, S)$, or *free space*.

Now consider the *motion planning* problem in robotics. Given a robot $R$, an work space $S$, and initial configuration $s$ and final configuration $t$ (both points in the robot’s free configuration space), determine (if possible) a way to move the robot from one configuration to the other without intersecting any of the obstacles. This reduces to the problem of determining whether there is a path from $s$ to $t$ that lies entirely within the robot’s free configuration space. Thus, we map the task of computing a robot’s motion to the problem of finding a path for a single point through a collection of obstacles.

Configuration spaces are typically higher dimensional spaces, and can be bounded by curved surfaces (especially when rotational elements are involved). Perhaps the simplest case to visualize is that of translating a convex polygonal robot in the plane amidst a collection of polygonal obstacles. In this case both the work space and configuration space are two dimensional. Consider a reference point placed in the center of the robot. As shown in the figure above, the process of mapping to configuration space involves replacing the robot with a single point (its reference point) and “growing” the obstacles by a compensating amount. These grown obstacles are called *configuration obstacles* or $C$-obstacles.

This approach while very general, ignores many important practical issues. It assumes that we have complete knowledge of the robot’s environment and have perfect knowledge and control of its placement. As stated we place no requirements on the nature of the path, but in reality physical objects can not be brought to move and stop instantaneously. Nonetheless, this abstract view is very powerful, since it allows us to abstract the motion planning problem into a very general framework.

For the rest of the lecture we will consider a very simple case of a convex polygonal robot that is translating among a convex set of obstacles. Even this very simple problem has a number of interesting algorithmic issues.

**Planning the Motion of a Point Robot:** As mentioned above, we can reduce complex motion planning problems to the problem of planning the motion of a point in free configuration space. First we will consider the question of how to plan the motion of a point amidst a set of polygonal obstacles in the plane, and then we will consider the question of how to construct configuration spaces.

To determine whether there is a path from one point to another of free configuration space, we will subdivide free space into simple convex regions. In the plane, we already know how to do this by computing a trapezoidal map. We can construct a trapezoidal map for all of the line segments bounding the obstacles, then throw away any faces that lie in the forbidden space. We also assume that we have a point location data structure for the trapezoidal map.
Next, we create a planar graph, called a \textit{road map}, based on the trapezoidal map. To do this we create a vertex in the center of each trapezoid and a vertex at the midpoint of each vertical edge. We create edges joining each center vertex to the vertices on its (at most four) edges.

![Figure 31.3: Motion planning using road maps.](image)

Now to answer the motion planning problem, we assume we are given the start point \( s \) and destination point \( t \). We locate the trapezoids containing these two points, and connect them to the corresponding center vertices. We can join them by a straight line segment, because the cells of the subdivision are convex. Then we determine whether there is a path in the road map graph between these two vertices, say by breadth-first search. Note that this will not necessarily produce the shortest path, but if there is a path from one position to the other, it will find it.

This description ignores many practical issues that arise in motion planning, but it is the basis for many practical motion planning problems. More realistic configuration spaces will contain more information (for example, encodings of the current joint rotation velocities) and will usually refine the road map to a much finer extent, so that short paths can be approximated well, as well as handling other elements such as guaranteeing minimal clearances around obstacles.

**Configuration Obstacles and Minkowski Sums:** Let us consider how to build a configuration space for a set of polygonal obstacles. We consider the simplest case of translating a convex polygonal robot amidst a collection of convex obstacles. If the obstacles are not convex, then we may subdivide them into convex pieces.

Consider a robot \( \mathcal{R} \), whose placement is defined by a translation \( \vec{p} = (x, y) \). Let \( \mathcal{R}(x, y) \) (also denoted \( \mathcal{R}(\vec{p}) \)) be the placement of the robot with its reference point at \( \vec{p} \). Given an obstacle \( P \), the \textit{configuration obstacle} is defined as all the placements of \( \mathcal{R} \) that intersect \( P \), that is

\[
\mathcal{C}P = \{ \vec{p} | \mathcal{R}(\vec{p}) \cap P \neq \emptyset \}.
\]

One way to visualize \( \mathcal{C}P \) is to imagine “scraping” \( \mathcal{R} \) along the boundary of \( P \) and seeing the region traced out by \( \mathcal{R} \)’s reference point.

The problem we consider next is, given \( \mathcal{R} \) and \( P \), compute the configuration obstacle \( \mathcal{C}P \). To do this, we first introduce the notion of a \textit{Minkowski sum}. Let us violate our notions of affine geometry for a while, and think of points \((x, y)\) in the plane as vectors. Given any two sets \( S_1 \) and \( S_2 \) in the plane, define their \textit{Minkowski sum} to be the set of all pairwise sums of points taken from each set:

\[
S_1 \oplus S_2 = \{ \vec{p} + \vec{q} | \vec{p} \in S_1, \ \vec{q} \in S_2 \}.
\]

Also, define \(-S = \{-\vec{p} | \vec{p} \in S\}\). Observe that for the case of a translating robot, we can define \( \mathcal{R}(\vec{p}) \) as \( \mathcal{R} \oplus \vec{p} \).
Claim: Given a translating robot \( R \) and an obstacle \( P \), \( CP = P \oplus (-R) \).

Proof: We show that \( R(\vec{q}) \) intersects \( P \) if and only if \( \vec{q} \in P \oplus (-R) \). Note that \( \vec{q} \in P \oplus (-R) \) if and only if there exist \( \vec{p} \in P \) and \( \vec{r} \in R \) such that \( \vec{q} = \vec{p} - \vec{r} \). Similarly, \( R(\vec{q}) = R \oplus \vec{q} \) intersects \( P \) if and only if there exists points \( \vec{r} \in R \) and \( \vec{p} \in P \) such that \( \vec{r} + \vec{q} = \vec{p} \). These two conditions are clearly equivalent.

Note that the proof made no use of the convexity of \( R \) or \( P \). It works for any shapes and in any dimension. However, computation of the Minkowski sums is most efficient for convex polygons.

It is an easy matter to compute \(-R\) in linear time (by simply negating all of its vertices) the problem of computing the C-obstacle \( CP \) reduces to the problem of computing a Minkowski sum of two convex polygons. We claim that this can be done in \( O(m + n) \) time, where \( m \) is the number of vertices in \( R \) and \( n \) is the number of vertices in \( P \). We will leave the construction as an exercise.

The algorithm is based on the following observation. Given a vector \( \vec{d} \), we say that a point \( p \) is extreme in direction \( \vec{d} \) if it maximizes the dot product \( p \cdot \vec{d} \).

Observation: Given two polygons \( P \) and \( R \), then the set of extreme points of \( P \oplus R \) in direction \( \vec{d} \) is the set of sums of points \( p \) and \( r \) that are extreme in direction \( \vec{d} \) for \( P \) and \( R \), respectively.

The book leaves the proof as an exercise. It follows easily by the linearity of the dot product.

From this observation, it follows that there is a simple algorithm for computing \( P \oplus R \), when both are convex polygons. In particular, we perform an angular sweep by considering a unit vector \( \vec{d} \) rotating counterclockwise around a circle. As \( \vec{d} \) rotates, it is an easy matter to keep track of the vertex or edge of \( P \) and \( R \) that is extreme in this direction. Whenever \( \vec{d} \) is perpendicular to an edge of either \( P \) or \( R \), we add this edge to the vertex of the other polygon. The algorithm is given in the text, and is illustrated in the figure below. The technique of applying one or more angular sweeps to a convex polygon is called the method of rotating calipers.

Assuming \( P \) and \( R \) are convex, observe that each edge of \( P \) and each edge of \( R \) contributes exactly one edge to \( P + R \). (If two edges are parallel and on the same side of the polygons, then these edges will be combined into one edge, which is as long as their sum.) Thus we have the following.

Claim: Given two convex polygons, \( P \) and \( R \), with \( n \) and \( m \) edges respectively, their Minkowski sum \( P \oplus R \) can be computed in \( O(n + m) \) time, and consist of at most \( n + m \) edges.
Complexity of Minkowski Sums: We have shown that free space for a translating robot is the complement of a union of C-obstacles \( \cup P_i \), each of which is a Minkowski sum of the form \( P_i \oplus R \), where \( P_i \) ranges over all the obstacles in the environment. If \( P_i \) and \( R \) are polygons, then the resulting region will be a union of polygons. How complex might this union be, that is, how many edges and vertices might it have?

To begin with, let’s see just how bad things might be. Suppose you are given a robot \( R \) with \( m \) sides and a set of work-space obstacle \( P \) with \( n \) sides. How many sides might the Minkowski sum \( P \oplus R \) have in the worst case? \( O(n + m) \)? \( O(nm) \), even more? The complexity generally depends on what special properties if any \( P \) and \( R \) have.

Nonconvex Robot and Nonconvex Obstacles: Suppose that both \( R \) and \( P \) are nonconvex simple polygons. Let \( m \) be the number of sides of \( R \) and \( n \) be the number of sides of \( P \). How many sides might there be in the Minkowski sum \( P \oplus R \) in the worst case? We can derive a quick upper bound as follows. First observe that if we triangulate \( P \), we can break it into the union of at most \( n \) - 2 triangles. That is:

\[
P = \bigcup_{i=1}^{n-2} T_i,
\]

\[
R = \bigcup_{j=1}^{m-2} S_j.
\]

It follows that

\[
P \oplus R = \bigcup_{i=1}^{n-2} \bigcup_{j=1}^{m-2} (T_i \oplus S_j).
\]

Thus, the Minkowski sum is the union of \( O(nm) \) polygons, each of constant complexity. Thus, there are \( O(nm) \) sides in all of these polygons. The arrangement of all of these line segments can have at most \( O(n^2m^2) \) intersection points (if each side intersects with each other), and hence this is an upper bound on the number of vertices in the final result.

Could things really be this bad? Yes they could. Consider the two polygons in the figure below left. There are \( O(n^2m^2) \) ways that these two polygons can be “docked”, as shown on the right. The Minkowski sum \( P \oplus -R \) is shown in the text. Notice that the large size is caused by the number of holes. (It might be argued that this is not fair, since we are not really interested in the entire Minkowski sum, just a single face of the Minkowski sum. Proving bounds on the complexity of a single face is an interesting problem, and the analysis is quite complex.)

As a final observation, notice that the upper bound holds even if \( P \) (and \( R \) for that matter) is not a single simple polygon, but any union of \( n \) triangles.

Convex Robot and Nonconvex Obstacles: We have seen that the worst-case complexity of the Minkowski sum might range from \( O(n + m) \) to as high as \( O(n^2m^2) \), which is quite a gap. Let us consider an
intermediate but realistic situation. Suppose that we assume that \( P \) is an arbitrary \( n \)-sided simple polygon, and \( R \) is a convex \( m \)-sided polygon. Typically \( m \) is much smaller than \( n \). What is the combinatorial complexity of \( P \oplus R \) in the worst case? As before we can observe that \( P \) can be decomposed into the union of \( n - 2 \) triangles \( T_i \), implying that
\[
P \oplus R = \bigcup_{i=1}^{n-2} (T_i \oplus R).
\]
Each Minkowski sum in the union is of complexity \( m + 3 \). So the question is how many sides might there be in the union of \( O(n) \) convex polygons each with \( O(m) \) sides? We could derive a bound on this quantity, but it will give a rather poor bound on the worst-case complexity. To see why, consider the limiting case of \( m = 3 \). We have the union of \( n \) convex objects, each of complexity \( O(1) \). This could have complexity as high as \( \Omega(n^2) \), as seen by generating a criss-crossing pattern of very skinny triangles. But, if you try to construct such a counter example, you won’t be able to do it.

To see why such a counterexample is impossible, suppose that you start with nonintersecting triangles, and then take the Minkowski sum with some convex polygon. The claim is that it is impossible to generate this sort of criss-cross arrangement. So how complex an arrangement can you construct? We will show the following.

**Theorem:** Let \( R \) be a convex \( m \)-gon and \( P \) and simple \( n \)-gon, then the Minkowski sum \( P \oplus R \) has total complexity \( O(nm) \).

Is \( O(nm) \) an attainable bound? The idea is to go back to our analogy of “scraping” \( R \) around the boundary of \( P \). Can we arrange \( P \) such that most of the edges of \( R \) scrape over most of the \( n \) vertices of \( P \)? Suppose that \( R \) is a regular convex polygon with \( m \) sides, and that \( P \) has the comb structure shown in the figure below, where the teeth of the comb are separated by a distance at least as large as the diameter of \( R \). In this case \( R \) will have many sides scrape across each of the pointy ends of the teeth, implying that the final Minkowski sum will have total complexity \( \Omega(nm) \).

The **Union of Pseudodisks:** Consider a translating robot given as an \( m \)-sided convex polygon and a collection of polygonal obstacles having a total of \( n \) vertices. We may assume that the polygonal obstacles have been triangulated into at most \( n \) triangles, and so, without any loss of generality, let us consider an instance of an \( m \)-sided robot translating among a set of \( n \) triangles. As argued earlier, each \( C \)-obstacle has \( O(3 + m) = O(m) \) sides, for a total of \( O(nm) \) line segments. A naive analysis suggests
that this many line segments might generate as many as $O(n^2m^2)$ intersections, and so the complexity of the free space can be no larger. However, we assert that the complexity of the space will be much smaller, in fact its complexity will be $O(nm)$.

To show that $O(nm)$ is an upper bound, we need some way of extracting the special geometric structure of the union of Minkowski sums. Recall that we are computing the union of $T_i \oplus R$, where the $T_i$’s have disjoint interiors. A set of convex objects $\{o_1, o_2, \ldots, o_n\}$ is called a collection of pseudodisks if for any two distinct objects $o_i$ and $o_j$ both of the set-theoretic differences $o_i \setminus o_j$ and $o_j \setminus o_i$ are connected. That is, if the objects intersect then they do not “cross through” one another. Note that the pseudodisk property is not a property of a single object, but a property that holds among a set of objects.

**Figure 31.8: Pseudodisks.**

**Lemma 1:** Given a set convex objects $T_1, T_2, \ldots, T_n$ with disjoint interiors, and convex $R$, the set

$$\{T_i \oplus R \mid 1 \leq i \leq n\}$$

is a collection of pseudodisks.

**Proof:** Consider two polygons $T_1$ and $T_2$ with disjoint interiors. We want to show that $T_1 \oplus R$ and $T_2 \oplus R$ do not cross over one another.

Given any directional unit vector $\vec{d}$, the most extreme point of $R$ in direction $\vec{d}$ is the point $r \in R$ that maximizes the dot product $(\vec{d} \cdot r)$. (Recall that we treat the “points” of the polygons as if they were vectors.) The point of $T_1 \oplus R$ that is most extreme in direction $\vec{d}$ is the sum of the points $t$ and $r$ that are most extreme for $T_1$ and $R$, respectively.

Given two convex polygons $T_1$ and $T_2$ with disjoint interiors, they define two outer tangents, as shown in the figure below. Let $d_1$ and $d_2$ be the outward pointing perpendicular vectors for these tangents. Because these polygons do not intersect, it follows easily that as the directional vector rotates from $d_1$ to $d_2$, $T_1$ will be the more extreme polygon, and from $d_2$ to $d_1$, $T_2$ will be the more extreme. See the figure below.
Figure 31.9: Alternation of extremes.

Now, if to the contrary $T_1 \oplus R$ and $T_2 \oplus R$ had a crossing intersection, then observe that we can find points $p_1$, $p_2$, $p_3$, and $p_4$, in cyclic order around the boundary of the convex hull of $(T_1 \oplus R) \cup (T_2 \oplus R)$ such that $p_1, p_3 \in T_1 \oplus R$ and $p_2, p_4 \in T_2 \oplus R$. First consider $p_1$. Because it is on the convex hull, consider the direction $\vec{d}_1$ perpendicular to the supporting line here. Let $r$, $t_1$, and $t_2$ be the extreme points of $R$, $T_1$ and $T_2$ in direction $\vec{d}_1$, respectively. From our basic fact about Minkowski sums we have

$$p_1 = r + t_1 \quad p_2 = r + t_2.$$ 

Since $p_1$ is on the convex hull, it follows that $t_1$ is more extreme than $t_2$ in direction $\vec{d}_1$, that is, $T_1$ is more extreme than $T_2$ in direction $\vec{d}_1$. By applying this same argument, we find that $T_1$ is more extreme than $T_2$ in directions $\vec{d}_1$ and $\vec{d}_3$, but that $T_2$ is more extreme than $T_1$ in directions $\vec{d}_2$ and $\vec{d}_4$. But this is impossible, since from the observation above, there can be at most one alternation in extreme points for nonintersecting convex polygons. See the figure below.

Figure 31.10: Proof of Lemma 1.

Lemma 2: Given a collection of pseudodisks, with a total of $n$ vertices, the complexity of their union is $O(n)$.

Proof: This is a rather cute combinatorial lemma. We are given some collection of pseudodisks, and told that altogether they have $n$ vertices. We claim that their entire union has complexity $O(n)$. (Recall that in general the union of $n$ convex polygons can have complexity $O(n^2)$, by criss-crossing.) The proof is based on a clever charging scheme. Each vertex in the union will be charged to a vertex among the original pseudodisks, such that no vertex is charged more than twice. This will imply that the total complexity is at most $2n$.

There are two types of vertices that may appear on the boundary. The first are vertices from the original polygons that appear on the union. There can be at most $n$ such vertices, and each is charged to itself. The more troublesome vertices are those that arise when two edges of two
pseudodisks intersect each other. Suppose that two edges $e_1$ and $e_2$ of pseudodisks $P_1$ and $P_2$ intersect along the union. Follow edge $e_1$ into the interior of the pseudodisk $e_2$. Two things might happen. First, we might hit the endpoint $v$ of this $e_1$ before leaving the interior $P_2$. In this case, charge the intersection to $v$. Note that $v$ can get at most two such charges, one from either incident edge. If $e_1$ passes all the way through $P_2$ before coming to the endpoint, then try to do the same with edge $e_2$. Again, if it hits its endpoint before coming out of $P_1$, then charge to this endpoint. See the figure below.

![Figure 31.11: Proof of Lemma 2.](image)

But what do we do if both $e_1$ shoots straight through $P_2$ and $e_2$ shoots straight through $P_1$? Now we have no vertex to charge. This is okay, because the pseudodisk property implies that this cannot happen. If both edges shoot completely through, then the two polygons must cross over each other.

Recall that in our application of this lemma, we have $n$ C-obstacles, each of which has at most $m + 3$ vertices, for a total input complexity of $O(nm)$. Since they are all pseudodisks, it follows from Lemma 2 that the total complexity of the free space is $O(nm)$. 
Fixed-Radius Near Neighbors

Reading: This material is taken from the paper: “The complexity of finding fixed-radius near neighbors,” by J. L. Bentley, D. F. Stanat, and E. H. Williams, Information Processing Letters, 6(6), 1977, 209–212.

Fixed-Radius Near Neighbor Problem: As a warm-up exercise for the course, we begin by considering one of the oldest results in computational geometry. This problem was considered back in the mid 70’s, and is a fundamental problem involving a set of points in dimension \(d\). We will consider the problem in the plane, but the generalization to higher dimensions will be straightforward. The solution also illustrates a common class of algorithms in CG, which are based on grouping objects into buckets that are arranged in a square grid.

We are given a set \(P\) of \(n\) points in the plane. It will be our practice throughout the course to assume that each point \(p\) is represented by its \((x, y)\) coordinates, denoted \((p_x, p_y)\). Recall that the Euclidean distance between two points \(p\) and \(q\), denoted \(\|pq\|\), is

\[
\|pq\| = \sqrt{(p_x - q_x)^2 + (p_y - q_y)^2}.
\]

Given the set \(P\) and a distance \(r > 0\), our goal is to report all pairs of distinct points \(p, q \in P\) such that \(\|pq\| \leq r\). This is called the fixed-radius near neighbor (reporting) problem.

Reporting versus Counting: We note that this is a reporting problem, which means that our objective is to report all such pairs. This is in contrast to the corresponding counting problem, in which the objective is to return a count of the number of pairs satisfying the distance condition.

It is usually easier to solve reporting problems optimally than counting problems. This may seem counterintuitive at first (after all, if you can report, then you can certainly count). The reason is that we know that any algorithm that reports some number \(k\) of pairs must take at least \(\Omega(k)\) time. Thus if \(k\) is large, a reporting algorithm has the luxury of being able to run for a longer time and still claim to be optimal. In contrast, we cannot apply such a lower bound to a counting algorithm.

The approach described here seems to work only for the reporting case. There is a more efficient solution to the counting problem, but this requires more sophisticated methods.

Simple Observations: To begin, let us make a few simple observations. This problem can easily be solved in \(O(n^2)\) time, by simply enumerating all pairs of distinct points and computing the distance between each pair. The number of distinct pairs of \(n\) points is

\[
\binom{n}{2} = \frac{n(n-1)}{2}.
\]

Letting \(k\) denote the number of pairs that reported, our goal will be to find an algorithm whose running time is (nearly) linear in \(n\) and \(k\), ideally \(O(n + k)\). This will be optimal, since any algorithm must take the time to read all the input and print all the results. (This assumes a naive representation of the output. Perhaps there are more clever ways in which to encode the output, which would require less than \(O(k)\) space.)

To gain some insight to the problem, let us consider how to solve the 1-dimensional version, where we are just given a set of \(n\) points on the line, say, \(x_1, x_2, \ldots, x_n\). In this case, one solution would be to first sort the values in increasing order. Let suppose we have already done this, and so:

\[x_1 < x_2 < \ldots < x_n.\]
Now, for $i$ running from 1 to $n$, we consider the successive points $x_{i+1}, x_{i+2}, x_{i+3}$, and so on, until we first find a point whose distance exceeds $r$. We report $x_i$ together with all succeeding points that are within distance $r$.

![Figure 32.1: Fixed radius nearest neighbor on the line.](image)

The running time of this algorithm involves the $O(n \log n)$ time needed to sort the points and the time required for distance computations. Let $k_i$ denote the number of pairs generated when we visit $p_i$. Observe that the processing of $p_i$ involves $k_i + 1$ distance computations (one additional computation for the points whose distance exceeds $r$). Thus, up to constant factors, the total running time is:

$$T(n, k) = n \log n + \sum_{i=1}^{n} (k_i + 1) = n \log n + n \sum_{i=1}^{n} k_i$$

This is close to the $O(k + n)$ time we were hoping for. It seems that any approach based on sorting is doomed to take at least $\Omega(n \log n)$ time. So, if we are to improve upon this, we cannot sort. But is sorting really necessary? Let us consider an approach based on bucketing.

**1-dimensional Solution with Bucketing:** Rather than sorting the points, suppose that we subdivide the line into intervals of length $r$. In particular, we can take the line to be composed of an infinite collection of half-open intervals:

$$\ldots, [-3r, -2r), [-2r, -r), [-r, 0), [0, r), [r, 2r), [2r, 3r), \ldots$$

We refer to these disjoint intervals as *buckets*. Given the interval $[br, (b+1)r)$, its *bucket index* is the integer $b$. Given any point $x$, it is easy to see that the index of the containing bucket is just $b(x) = \lfloor x/r \rfloor$. Thus, in $O(n)$ time we can associate the $n$ points of $P$ with a set of $n$ integer bucket indices, $b(x)$ for each $x \in P$. Although there are an infinite number of buckets, at most $n$ will be *occupied*, meaning that they contain at least one point of $P$.

There are a number of ways to organize the occupied buckets. They could be sorted, but then we are back to $O(n \log n)$ time. Since bucket indices are integers, a better approach is to store the occupied bucket indices in a *hash table*. Recall from basic data structures that a hash table is a data structure that supports the following operations in $O(1)$ expected time:

- **insert**($o, b$) : Insert object $o$ with key value $b$. We allow multiple objects to be inserted with the same key.
- **find**($b$) : Return a list $L$ of references to objects having key value $b$. This operation takes $O(1 + |L|)$ expected time. If no keys have this value, then an empty list is returned.
- **remove**($o, b$) : Remove the object indicated by reference $o$, having key value $b$ from the table.
Each point is associated with the key value given by its bucket index $b = \lfloor x/r \rfloor$. Thus in $O(1)$ expected time, we can determine which bucket contains a given point and look this bucket up in the hash table.

The fact that the running time is in the expected case, rather than worst case is a bit unpleasant. However, it can be shown that by using a good randomized hash function, the probability that the total running time is worse than $O(n)$ can be made arbitrarily small. If the algorithm performs significantly more than the expected number of computations, we can simply chose a different random hash function and try again. This will lead to a very practical solution.

How does bucketing help? Observe that if point $p$ lies in bucket $b$, then any successors that lie within distance $r$ must lie either in bucket $b$ or in $b+1$. This suggests the straightforward solution shown below.

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**Fixed-Radius Near Neighbor on the Line by Bucketing**

1. For each $p \in P$, insert $p$ in the hash table with the key value $b(p)$.
2. For each $p \in P$ do the following:
   - Let $b(p)$ be the bucket containing $p$.
   - Enumerate all the points of buckets $b(p)$ and $b(p)+1$, and for each point $q \in b(p) \cup b(p)+1$ such that $q \neq p$ and $|q-p| \leq r$, output the pair $(p, q)$.

Note that this will output duplicate pairs $(p, q)$ and $(q, p)$. If this bothers you, we could add the additional condition that $q > p$. The key question is determining the time complexity of this algorithm is how many distance computations are performed in step (2b). We compare each point in bucket $b$ with all the points in buckets $b$ and $b+1$. However, not all of these distance computations will result in a pair of points whose distance is within $r$. Might it be that we waste a great deal of time in performing computations for which we receive no benefit? The lemma below shows that we perform no more than a constant factor times as many distance computations and pairs that are produced.

It will simplify things considerably if, rather than counting distinct pairs of points, we simply count all (ordered) pairs of points that lie within distance $r$ of each other. Thus each pair of points will be counted twice, $(p, q)$ and $(q, p)$. Note that this includes reporting each point as a pair $(p, p)$ as well, since each point is within distance $r$ of itself. This does not affect the asymptotic bounds, since the number of distinct pairs is smaller by a factor of roughly $1/2$.

**Lemma**: Let $k$ denote the number of (not necessarily distinct) pairs of points of $P$ that are within distance $r$ of each other. Let $D$ denote the total number distance computations made in step (2b) of the above algorithm. Then $D \leq 2k$.

**Proof**: We will make use of the following inequality in the proof:

$$xy \leq \frac{x^2 + y^2}{2}.$$

This follows by expanding the obvious inequality $(x - y)^2 \geq 0$.

Let $B$ denote the (infinite) set of buckets. For any bucket $b \in B$, let $b+1$ denote its successor bucket on the line, and let $n_b$ denote the number of points of $P$ in $b$. Define

$$S = \sum_{b \in B} n_b^2.$$
First we bound the total number of distance computations $D$ as a function of $S$. Each point in bucket $b$ computes the distance to every other point in bucket $b$ and every point in bucket $b + 1$, and hence

$$D = \sum_{b \in B} n_b (n_b + n_{b+1}) = \sum_{b \in B} n_b^2 + n_b n_{b+1} = \sum_{b \in B} n_b^2 + \sum_{b \in B} n_b n_{b+1}$$

$\leq \sum_{b \in B} n_b^2 + \sum_{b \in B} \frac{n_b^2 + n_{b+1}^2}{2}$

$= \sum_{b \in B} n_b^2 + \sum_{b \in B} \frac{n_b^2}{2} + \sum_{b \in B} \frac{n_{b+1}^2}{2} = S + \frac{S}{2} + \frac{S}{2} = 2S.$

Next we bound the number of pairs reported from below as a function of $S$. Since each pair of points lying in bucket $b$ is within distance $r$ of every other, there are $n_b^2$ pairs in bucket $b$ alone that are within distance $r$ of each other, and hence (considering just the pairs generated within each bucket) we have $k \geq S$. Therefore we have

$$D \leq 2S \leq 2k,$$

which completes the proof.

By combining this with the $O(n)$ expected time needed to bucket the points, it follows that the total expected running time is $O(n + k)$.

A worthwhile exercise to consider at this point is the issue of the bucket width $r$. How would changing the value of $r$ affect the implementation of the algorithm and its efficiency? For example, if we used buckets of size $r/2$ or $2r$, would the above algorithm (after suitable modifications) have the same asymptotic running time? Would buckets of size any constant times $r$ work?

**Generalization to $d$ dimensions:** This bucketing algorithm is easy to extend to multiple dimensions. For example, in dimension 2, we bucket points into a square grid in which each grid square is of side length $r$. (As before, you might consider the question of what values of bucket sizes lead to a correct and efficient algorithm.) The bucket index of a point $p : (p_x, p_y)$ is a pair $B(p) = (b(p_x), b(p_y)) = (\lfloor p_x/r \rfloor, \lfloor p_y/r \rfloor)$. We apply a hash function that accepts two arguments. To generalize the algorithm, for each point we consider the points in its surrounding $3 \times 3$ subgrid of buckets. The result is shown in the following code fragment.

![Figure 32.2: Fixed radius nearest neighbor on the plane.](image-url)
(1) For each \( p = (p_x, p_y) \) of \( P \), insert \( p \) in the hash table with the (2-dimensional) bucket index \( B(p) = (b(p_x), b(p_y)) \).

(2) For each \( p \in P \) do the following:
   
   (a) Let \( B(p) = (b(p_x), b(p_y)) \) be the bucket index for \( p \).

   (b) Enumerate all the points of buckets \((b(p_x) + i, b(p_y) + j)\), for \( i, j \in \{-1, 0, +1\} \). For each such point \( q \), if \( \|pq\| \leq r \), output the pair \((p, q)\).

By generalizing the analysis used in the 1-dimensional case, it can be shown that the algorithm’s expected running time is \( O(n + k) \). The details are left as an exercise (we just have more terms to consider, but each cell is involved with at most 9 other cells which is absorbed into the constant factor hidden by the O-notation).

This example problem serves to illustrate some of the typical elements of computational geometry. Geometry itself did not play a significant role in the problem, other than the relatively easy task of computing distances. We will see examples later this semester where geometry plays a much more important role. The major emphasis was on accounting for the algorithm’s running time. Also note that, although we discussed the possibility of generalizing the algorithm to higher dimensions, we did not treat the dimension as an asymptotic quantity. In fact, a more careful analysis reveals that this algorithm’s running time increases exponentially with the dimension. (Can you see why?)
33 More Convex Hull Algorithms

Reading: Today’s material is not covered in the 4M’s book. Chan’s output sensitive algorithm can be found in T. Chan, “Optimal output-sensitive convex hull algorithms in two and three dimensions”, *Discrete and Computational Geometry*, 16, 1996, 361–368.

Output Sensitive Convex Hull Algorithms: It turns out that in the worst-case, convex hulls cannot be computed faster than in $\Omega(n \log n)$ time. One way to see this intuitively is to observe that the convex hull itself is sorted along its boundary, and hence if every point lies on the hull, then computing the hull requires sorting of some form. Yao proved the much harder result that determining which points are on the hull (without sorting them along the boundary) still requires $\Omega(n \log n)$ time. However both of these results rely on the fact that all (or at least a constant fraction) of the points lie on the convex hull. This is often not true in practice.

The Jarvis’s March algorithms that we saw last time raises the question of just how fast can convex hulls be computed assuming we allow the running time to be described in terms of both the input size $n$ and the output size $h$. Many geometric algorithms have the property that the output size can be a widely varying function of the input size, and worst-case output size may not be a good indicator of what happens typically. An algorithm which attempts to be more efficient for small output sizes is called an output sensitive algorithm, and running time is described as a asymptotic function of both input size and output size.

Chan’s Algorithm: Given than any convex hull algorithm must take at least $O(n)$ time, and given that “$\log n$” factor arises from the fact that you need to sort the at most $n$ points on the hull, then a reasonable target running time is $O(n \log h)$. (Below we will see that this is optimal.) Kirkpatrick and Seidel discovered a relatively complicated $O(n \log h)$ time algorithm, based on a clever pruning method in 1986. The problem was considered closed until around 10 years later when Timothy Chan came up with a much simpler algorithm with the same running time. One of the interesting aspects of Chan’s algorithm is that it involves cleverly combining two slower algorithms (Graham’s scan and Jarvis’s March) together to form an algorithm that is faster than either one.

The problem with Graham’s scan is that it sorts all the points, and hence is doomed to having an $\Omega(n \log n)$ running time, irrespective of the size of the hull. On the other hand, Jarvis’s march can perform better if you have few vertices on the hull, but it takes $\Omega(n)$ time for each hull vertex.

The first observation is that if we hope to achieve a running time of $O(n \log h)$, we can only afford a log factor depending on $h$. So, if we run Graham’s algorithm, we are limited to considering sets of size $h$. (Actually, any polynomial in $h$ will work as well, since for any constant $c$ we have $\log h^c = c \log h = O(\log h)$. We’ll use this observation later on.) We do not know the value of $h$ in advance, but suppose for now that we had an estimate for $h$, call it $m$, whose value is at least as large as $h$, but not too much larger (say $h \leq m \leq h^2$).

Chan’s idea is to partition the points into groups, each of size $m$. The number of groups is $r = \lceil n/m \rceil$. For each group we compute its hull using Graham’s scan, which takes $O(m \log m)$ time per group, for a total time of $O(rm \log m) = O(n \log m)$. Since $m \leq h^2$, this is $O(n \log h)$, and so we are within our desired execution time budget. (Note, by the way, that since each group is an arbitrary subset of $P$, we cannot assume that it will have $h$ points on its hull. The number of hull points per group can be as large as $m$.)
This leaves the question of how to merge these \( r \) individual hulls together into a single hull. The idea is to run Jarvis’s march, but we treat each group as if it is a “fat point”. Here we take advantage of the fact that you can compute the tangents between a point and a convex \( m \)-gon in \( O(\log m) \) time. (We will leave the proof of this fact as an exercise, but observe that finding the point of tangency can be reduced to a sort of binary search.) Note that since each “fat point” is itself a convex polygon, it is quite possible that the next hull vertex comes from the same group as the current vertex.

Let’s analyze the running time. We know that there are \( h \) points on the final hull, and therefore at most \( h \) of \( r \) group hulls can contribute to the final hull. That is, there are at most \( h \) steps in the Jarvis march, where each step involves computing \( r \) tangents, and each tangent takes \( O(\log m) \) time to compute.

Putting this together, we find that the overall time for the merging phase of the algorithm is

\[
O(hr \log m) = O \left( h \frac{n}{m} \log m \right)
\]

time. Combining these two parts, we get a total of

\[
O \left( \left( n + h \frac{n}{m} \right) \log m \right)
\]

time. By our assumption that \( h \leq m \leq h^2 \) it follows that the total time is \( O(n \log h) \), as desired.

There is still one problem here. We do not know what \( h \) is in advance, and therefore we do not know what \( m \) should be when running the algorithm. We will see how to remedy this later. For now, let’s imagine that someone tells us the value of \( m \). The following algorithm works correctly as long as \( m \geq h \). If we are wrong, it returns a special error status.

**Chan’s Partial Convex Hull Algorithm (Given \( m \))**

\[
\text{PartialHull}(P, m) : \\
(1) \text{Let } r = \lceil n/m \rceil. \text{ Partition } P \text{ into disjoint subsets } P_1, P_2, \ldots, P_r, \text{ each of size at most } m. \\
(2) \text{For } i = 1 \text{ to } r \text{ do:} \\
\hspace{1em} (a) \text{Compute } \text{Hull}(P_i) \text{ using Graham’s scan and store the vertices in an ordered array.} \\
(3) \text{Let } p_0 = (-\infty, 0) \text{ and let } p_1 \text{ be the bottommost point of } P. \\
(4) \text{For } k = 1 \text{ to } m \text{ do:} \\
\hspace{1em} (a) \text{For } i = 1 \text{ to } r \text{ do:} \\
\hspace{2em} \bullet \text{Compute point } q_i \in P_i \text{ that maximizes the angle } \angle p_{k-1} p_k q_i. \\
\hspace{2em} (b) \text{Let } p_{k+1} \text{ be the point } q \in \{q_1, \ldots, q_r\} \text{ that maximizes the angle } \angle p_{k-1} p_k q. \\
\hspace{2em} (c) \text{If } p_{k+1} = p_1 \text{ then return } \langle p_1, \ldots, p_k \rangle. \\
(5) \text{Return ”Fail: } m \text{ was too small, try again.”}
\]

We assume that we store the convex hulls from step (2a) in an ordered array so that the step inside the for-loop of step (4a) can be solved in \( O(\log m) \) time using binary search. Otherwise, the analysis follows directly from the comments made earlier.

The only question remaining is how do we know what value to give to \( m \)? We could try \( m = 1, 2, 3, \ldots \), until we luck out and have \( m \geq h \), but this would take too long. (Convince yourself that this will take a total time of \( O(nh \log h) \), which is even worse than Jarvis’s march.)

Binary search for \( m \) is a more efficient option than linear search, but if we guess to large a value for \( m \) (e.g., \( m = n/2 \)) then we are immediately stuck with \( O(n \log n) \) time, and this is too slow. In order
to avoid this, a neat trick is to start with a small value of \( m \) (e.g., \( m = 1 \)) and repeatedly increase it by doubling until we succeed. That is, we run the algorithm with \( m = 1, 2, 4, 8, \ldots, 2^t \) until it succeeds. This is called doubling search. Unfortunately, it turns out that this is still too slow. (You should do the analysis yourself and convince yourself that it will result in a running time of \( O(m \log^2 h) \). Better but still not the best.)

The dependence in running time on \( m \) is only in the log term. As we observed earlier, as long as our value of \( m \) is within a polynomial of \( h \), that is, \( m \leq h^c \) for some constant \( c \), then the running time will still be \( O(n \log h) \). Chan’s approach is to guess successively larger values of \( m \), each time squaring the previous value, until the algorithm returns a successful result. Observe that the sequence of choices is therefore

\[
m = 2, 4, 16, \ldots
\]
\[
m = 2^1, 2^2, 2^4, \ldots, 2^{2^t}.
\]

This suggests the algorithm shown below.

---

**Chan’s Complete Convex Hull Algorithm**

**Hull**(\( P \)):

1. For \( t = 1, 2, \ldots \) do:
   - (a) Let \( m = \min(2^{2^t}, n) \).
   - (b) Invoke PartialHull(\( P, m \)), returning the result in \( L \).
   - (c) If \( L \neq \text{"Fail"} \) then return \( L \).

Note that \( 2^{2^t} \) has the effect of squaring the previous value of \( m \). How long does this take? The \( t \)-th iteration takes \( O(n \log 2^{2^t}) = O(n 2^t) \) time. We know that it will stop as soon as \( 2^{2^t} \geq h \), that is if \( t = \lceil \lg \lg h \rceil \). (We will use \( \lg \) to denote logarithm base 2.) Using the fact that \( \sum_{i=0}^{k} 2^i = 2^{k+1} - 1 \), the running time (up to constant factors) is

\[
\sum_{t=1}^{\lg \lg h} n 2^t = n \sum_{t=1}^{\lg \lg h} 2^t \leq n 2^{\lg \lg h} = 2n 2^{2\lg \lg h} = 2n \lg h = O(n \log h),
\]

which is just what we want.
Lower Bound: Next we will show that Chan’s result is asymptotically optimal in the sense that any algorithm for computing the convex hull of \(n\) points with \(h\) points on the hull requires \(\Omega(n \log h)\) time. The proof is a generalization of the proof that sorting a set of \(n\) numbers requires \(\Omega(n \log n)\) comparisons.

If you recall the proof that sorting takes at least \(\Omega(n \log n)\) comparisons, it is based on the idea that any sorting algorithm can be described in terms of a decision tree. Each comparison has at most 3 outcomes (<, =, or >). Each such comparison corresponds to an internal node in the tree. The execution of an algorithm can be viewed as a traversal along a path in the resulting 3-ary tree. The height of the tree is a lower bound on the worst-case running time of the algorithm. There are at least \(n!\) different possible inputs, each of which must be reordered differently, and so you have a 3-ary tree with at least \(n\) leaves. Any such tree must have \(\Omega(\log_3 n!)\) height. Using Stirling’s approximation for \(n!\), this solves to \(\Omega(n \log n)\) height.

We will give an \(\Omega(n \log h)\) lower bound for the convex hull problem. In fact, we will give an \(\Omega(n \log h)\) lower bound on the following simpler decision problem, whose output is either yes or no.

Convex Hull Size Verification Problem (CHSV): Given a point set \(P\) and integer \(h\), does the convex hull of \(P\) have \(h\) distinct vertices?

Clearly if this takes \(\Omega(n \log h)\) time, then computing the hull must take at least as long. As with sorting, we will assume that the computation is described in the form of a decision tree. Assuming that the algorithm uses only comparisons is too restrictive for computing convex hulls, so we will generalize the model of computation to allow for more complex functions. We assume that we are allowed to compute any algebraic function of the point coordinates, and test the sign of the resulting function. The result is called an algebraic decision tree.

The input to the CHSV problem is a sequence of \(2n = N\) real numbers. We can think of these numbers as forming a vector \((z_1, z_2, \ldots, z_N) = \vec{z} \in \mathbb{R}^N\), which we will call a configuration. Each node of the decision tree is associated with a multivariate algebraic formula of degree at most \(d\). e.g.

\[
f(\vec{z}) = z_1z_4 - 2z_3z_6 + z_0^2,
\]

would be an algebraic function of degree 2. The node branches in one of three ways, depending on whether the result is negative, zero, or positive. (Computing orientations and dot-products both fall into this category.) Each leaf of the resulting tree corresponds to a possible answer that the algorithm might give.

For each input vector \(\vec{z}\) to the CHSV problem, the answer is either “yes” or “no”. The set of all “yes” points is just a subset of points \(Y \subset \mathbb{R}^N\), that is a region in this space. Given an arbitrary input \(\vec{z}\) the purpose of the decision tree is to tell us whether this point is in \(Y\) or not. This is done by walking down the tree, evaluating the functions on \(\vec{z}\) and following the appropriate branches until arriving at a leaf, which is either labeled “yes” (meaning \(\vec{z} \in Y\)) or “no”. An abstract example (not for the convex hull problem) of a region of configuration space and a possible algebraic decision tree (of degree 1) is shown in the following figure. (We have simplified it by making it a binary tree.) In this case the input is just a pair of real numbers.

We say that two points \(\vec{u}, \vec{v} \in Y\) are in the same connected component of \(Y\) if there is a path in \(\mathbb{R}^N\) from \(\vec{u}\) to \(\vec{v}\) such that all the points along the path are in the set \(Y\). (There are two connected
components in the figure.) We will make use of the following fundamental result on algebraic decision trees, due to Ben-Or. Intuitively, it states that if your set has $M$ connected components, then there must be at least $M$ leaves in any decision tree for the set, and the tree must have height at least the logarithm of the number of leaves.

**Theorem:** Let $Y \in R^N$ be any set and let $T$ be any $d$-th order algebraic decision tree that determines membership in $W$. If $W$ has $M$ disjoint connected components, then $T$ must have height at least $\Omega((\log M) - N)$.

We will begin our proof with a simpler problem.

**Multiset Size Verification Problem (MSV):** Given a multiset of $n$ real numbers and an integer $k$, confirm that the multiset has exactly $k$ distinct elements.

**Lemma:** The MSV problem requires $\Omega(n \log k)$ steps in the worst case in the $d$-th order algebraic decision tree.

**Proof:** In terms of points in $R^n$, the set of points for which the answer is “yes” is

$$Y = \{(z_1, z_2, \ldots, z_n) \in R^n \mid |\{z_1, z_2, \ldots, z_n\}| = k\}.$$  

It suffices to show that there are at least $k!k^{n-k}$ different connected components in this set, because by Ben-Or’s result it would follow that the time to test membership in $Y$ would be

$$\Omega(\log(k!k^{n-k}) - n) = \Omega(k \log k + (n - k) \log k - n) = \Omega(n \log k).$$

Consider the all the tuples $(z_1, \ldots, z_n)$ with $z_1, \ldots, z_k$ set to the distinct integers from 1 to $k$, and $z_{k+1} \ldots z_n$ each set to an arbitrary integer in the same range. Clearly there are $k!$ ways to select the first $k$ elements and $k^{n-k}$ ways to select the remaining elements. Each such tuple has exactly $k$ distinct items, but it is not hard to see that if we attempt to continuously modify one of these tuples to equal another one, we must change the number of distinct elements, implying that each of these tuples is in a different connected component of $Y$.

To finish the lower bound proof, we argue that any instance of MSV can be reduced to the convex hull size verification problem (CHSV). Thus any lower bound for MSV problem applies to CHSV as well.

**Theorem:** The CHSV problem requires $\Omega(n \log h)$ time to solve.
**Proof:** Let \( Z = (z_1, \ldots, z_n) \) and \( k \) be an instance of the MSV problem. We create a point set \( \{p_1, \ldots, p_n\} \) in the plane where \( p_i = (z_i, z_i^2) \), and set \( h = k \). (Observe that the points lie on a parabola, so that all the points are on the convex hull.) Now, if the multiset \( Z \) has exactly \( k \) distinct elements, then there are exactly \( h = k \) points in the point set (since the others are all duplicates of these) and so there are exactly \( h \) points on the hull. Conversely, if there are \( h \) points on the convex hull, then there were exactly \( h = k \) distinct numbers in the multiset to begin with in \( Z \).

Thus, we cannot solve CHSV any faster than \( \Omega(n \log h) \) time, for otherwise we could solve MSV in the same time.

The proof is rather unsatisfying, because it relies on the fact that there are many duplicate points. You might wonder, does the lower bound still hold if there are no duplicates? Kirkpatric and Seidel actually prove a stronger (but harder) result that the \( \Omega(n \log h) \) lower bound holds even you assume that the points are distinct.
34 Multidimensional Polytopes and Convex Hulls

Reading: Today’s material is not covered in the 4M’s book. See the book Algorithmic Geometry by Boissonnat and Yvinec.

Polytopes: Today we consider convex hulls in dimensions 3 and higher. Although dimensions greater than 3 may seem rather esoteric, we shall see that many geometric optimization problems can be stated as some search over a polytope in \( d \)-dimensional space, where \( d \) may be greater than 3.

Before delving into this, let us first present some basic terms. We define a polytope (or more specifically a \( d \)-polytope) to be the convex hull of a finite set of points in \( \mathbb{R}^d \). We say that a set of \( k \) points is affinely independent if no one point can be expressed as an affine combination (that is, a linear combination whose coefficients sum to 1) of the others. For example, three points are affinely independent if they are not on the same line, four points are affinely independent if they are not on the same plane, and so on. The convex hull of \( k + 1 \) affinely independent points is called a simplex or \( k \)-simplex. For example, the line segment joining two points is a 1-simplex, the triangle defined by three points is a 2-simplex, and the tetrahedron defined by four points is a 3-simplex.

Any \((d-1)\)-dimensional hyperplane \( h \) in \( d \)-dimensional space divides the space into (open) halfspaces, denoted \( h^- \) and \( h^+ \), so that \( \mathbb{R}^d = h^- \cup h \cup h^+ \). Let us define \( h^- = h^- \cup h \) and \( h^+ = h^+ \cup h \) to be the closures of these halfspaces. We say that a hyperplane supports a polytope \( P \) (and is called a supporting hyperplane of \( P \)) if \( h \cap P \) is nonempty and \( P \) is entirely contained within either \( h^- \) or \( h^+ \). The intersection of the polytope and any supporting hyperplane is called a face of \( P \). Faces are themselves convex polytopes of dimensions ranging from 0 to \( d - 1 \). The 0-dimensional faces are called vertices, the 1-dimensional faces are called edges, and the \((d-1)\)-dimensional faces are called facets. (Note: When discussing polytopes in dimension 3, people often use the term “face” when they mean “facet”. It is usually clear from context which meaning is intended.)

The faces of dimensions 0 to \( d - 1 \) are called proper faces. It will be convenient to define two additional faces. The empty set is said to be a face of dimension \(-1\) and the entire polytope is said to be a face of dimension \( d \). We will refer to all the faces, including these two additional faces as the improper faces of the polytope.

There are a number of facts that follow from these definitions.
• The boundary of a polytope is the union of its proper faces.
• A polytope has a finite number of faces. Each face is a polytope.
• A polytope is the convex hull of its vertices.
• A polytope is the intersection of a finite number of closed halfspaces. (Note that the converse need not be true, since the intersection of halfspaces may generally be unbounded. Such an unbounded convex body is either called a polyhedron or an unbounded polytope.)

Observe that a $d$-simplex has a particularly regular face structure. If we let $v_0, v_1, v_2, \ldots, v_d$ denote the vertices of the simplex, then for each 2-element set $\{v_i, v_j\}$ there is an edge of the simplex joining these vertices, for each 3-element set $\{v_i, v_j, v_k\}$ there is a 3-face joining these three vertices, and so.

Fact: The number of $j$-dimensional faces on a $d$-simplex is equal to the number $(j + 1)$-element subsets of domain of size $d + 1$, that is,

$$\binom{d+1}{j+1} = \frac{(d+1)!}{(j+1)!(d-j)!}.$$

Incidence Graph: How are polytopes represented? In addition to the geometric properties of the polytope (e.g., the coordinates of its vertices or the equation of its faces) it is useful to store discrete connectivity information, which is often referred to as the topology of the polytope. There are many representations for polytopes. In dimension 2, a simple circular list of vertices suffices. In dimension 3, we need some sort of graph structure. There are many data structures that have been proposed. They are evaluated based on the ease with which the polytope can be traversed and the amount of storage needed. (Examples include the winged-edge, quad-edge, and half-edge data structures. We may discuss these later in the semester.)

A useful structure for polytopes in arbitrary dimensions is called the incidence graph. Each node of the incidence graph corresponds to an (improper) face of the polytope. We create an edge between two faces if their dimension differs by 1, and one (of lower dimension) is contained within the other (of higher dimension). An example is shown in Fig. 34.3 below for a simplex. Note the similarity between this graph and the lattice of subsets based on inclusion relation.

Figure 34.3: The incidence graph for a tetrahedron.

Polarity: There are two natural ways to create polytopes. One is as the convex hull of a set of points and the other is as the intersection of a collection of closed halfspaces (assuming it is bounded). These
two concepts are essentially identical, and this can be observed by the polar transformation, which maps points to hyperplanes and vice versa. Fix any point \( O \) in \( d \)-dimensional space. We may think of \( O \) as the origin, and therefore, any point \( p \in \mathbb{R}^d \) can be viewed as a \( d \)-element vector. (If \( O \) is not the origin, then \( p \) can be identified with the vector \( p - O \).) The polar hyperplane of \( p \), denoted \( p^* \) is defined by

\[
p^* = \{ x \in \mathbb{R}^d \mid (p \cdot x) = 1 \},
\]

where the expression \((p \cdot x)\) is just the standard vector dot-product \((p \cdot x) = p_1x_1 + p_2x_2 + \cdots + p_dx_d\). Observe that if \( p \) is on the unit sphere centered about \( O \), then \( p^* \) is a hyperplane that passes through \( O \) and is orthogonal to the vector \( OP \). As we move \( p \) away from the origin along this vector, the dual hyperplane moves closer to the origin, and vice versa, so that the product of their distances from the origin is always 1.

Now, let \( h \) be any hyperplane that does not contain \( O \). The pole of \( h \), denoted \( h^* \) is the point that satisfies

\[
(h^* \cdot x) = 1 \quad \text{for all } x \in h.
\]

Clearly this double transformation is an involution, that is, \((p^*)^* = p \) and \((h^*)^* = h \). The polar transformation preserves important geometric relationships. Given a hyperplane \( h \), define

\[
h^+ = \{ x \in \mathbb{R}^d \mid (x \cdot h^*) < 1 \} \quad h^- = \{ x \in \mathbb{R}^d \mid (x \cdot h^*) > 1 \}.
\]

That is, \( h^+ \) is the open halfspace containing the origin and \( h^- \) is the other open halfspace for \( h \).

**Claim:** Let \( p \) be any point in \( \mathbb{R}^d \) and let \( h \) be any hyperplane in \( \mathbb{R}^d \). The polar transformation satisfies the following two properties.

**Incidence preserving:** The polarity transformation preserves incidence relationships between points and hyperplanes. That is, \( p \) belongs to \( h \) if and only if \( h^* \) belongs to \( p^* \).

**Inclusion Reversing:** The polarity transformation reverses relative position relationships in the sense that \( p \) belongs to \( h^+ \) if and only if \( h^* \) belongs to \( (p^*)^+ \), and \( p \) belongs to \( h^- \) if and only if \( h^* \) belongs to \( (p^*)^- \).

In general, any bijective transformation that preserves incidence relations is called a duality. The above claim implies that polarity is a duality.

We can now formalize the aforementioned notion of polytope equivalence. The idea will be to transform a polytope defined as the convex hull of a finite set of points to a polytope defined as the intersection of a finite set of closed halfspaces. To do this, we need a way of mapping a point to a halfspace.
Our approach will be to take the halfspace that contains the origin. For any point \( p \in \mathbb{R}^d \) define the following closed halfspace based on its polar:

\[
p^\# = p^* = \{ x \in \mathbb{R}^d \mid (x \cdot p) \leq 1 \}.
\]

(The notation is ridiculous, but this is easy to parse. First consider the polar hyperplane of \( p \), and take the closed halfspace containing the origin.) Observe that if a halfspace \( h^+ \) contains \( p \), then by the inclusion-reversing property of polarity, the polar point \( h^* \) is contained within \( p^\# \).

Now, for any set of points \( P \subseteq \mathbb{R}^d \), we define its polar image to be the intersection of these halfspaces

\[
P^\# = \{ x \in \mathbb{R}^d \mid (x \cdot p) \leq 1, \forall p \in P \}.
\]

Thus \( P^\# \) is the intersection of an (infinite) set of closed halfspaces, one for each point \( p \in P \). A halfspace is convex and the intersect of convex sets is convex, so \( P^\# \) is a convex set.

To see the connection with convex hulls, let \( S = \{ p_1, \ldots, p_n \} \) be a set of points and let \( P = \text{conv}(S) \). Let us assume that the origin \( O \) is contained within \( P \). (We can guarantee this in a number of ways, e.g., by translating \( P \) so its center of mass coincides with the origin.) By definition, the convex hull is the intersection of the set of all closed halfspaces that contain \( S \). That is, \( P \) is the intersect of an infinite set of closed halfspaces. What are these halfspaces? If \( h^+ \) is a halfspace that contains all the points of \( S \), then by the inclusion-reversing property of polarity, the polar point \( h^* \) is contained within all the hyperplanes \( p_i^* \), which implies that \( h^* \in P^\# \). This means that, through polarity, the halfspaces whose intersection is the convex hull of a set of points is essentially equivalent to the polar points that lie within the polar image of the convex hull.

**Lemma:** Let \( S = \{ p_1, \ldots, p_n \} \) be a set of points in \( \mathbb{R}^d \) and let \( P = \text{conv}(S) \). Then its polar image is the intersection of the corresponding polar halfspaces, that is,

\[
P^\# = \bigcap_{i=1}^{n} p_i^\#.
\]

Furthermore:

(i) A point \( a \in \mathbb{R}^d \) lies on the boundary of \( P \) if and only if the polar hyperplane \( a^* \) supports \( P^\# \).

(ii) Each \( k \)-face of \( P \) corresponds to a \( (d-1-k) \)-face of \( P^\# \) and given faces \( f_1, f_2 \) of \( P \) where \( f_1 \subseteq f_2 \), the corresponding faces \( f_1^\#, f_2^\# \) of \( P^\# \) satisfy \( f_1^\# \supseteq f_2^\# \). (That is, inclusion relations are reversed.)
It is not hard to prove that the polar image of a polytope is an involution, that is \((P^#)^# = P\). (See Boissonnat and Yvinec for proofs of all these facts.)

Thus, the polar image \(P^#\) of a polytope is structurally isomorphic to \(P\) and all affine relations on \(P\) map through polarity to \(P^#\). From a computational perspective, this means that we compute the polar of all the points of \(P\), consider the halfspaces that contain the origin, and take the intersection of these halfspaces. Thus, the problems of computing convex hulls and computing the intersection of halfspaces are computationally equivalent. (In fact, once you have computed the incidence graph for one, you just flip it “upside-down” to get the other.)

For example, if you know your Platonic solids (tetrahedron, cube, octahedron, dodecahedron, and icosahedron), you may remember that the square and octahedron are polar duals, the dodecahedron and icosohedron are polar duals, and the tetrahedron is self-dual.

**Simple and Simplicial Polytopes:** Observe that if a polytope is the convex hull of a set of points in general position, then for \(0 \leq j \leq d - 1\), each \(j\)-face is a \(j\)-simplex. A polytope is simplicial if all its proper faces are simplices.

If we take a dual view, consider a polytope that is the intersection of a set of \(n\) halfspaces in general position. Then each \(j\)-face is the intersection of exactly \((d - j)\) hyperplanes. A polytope is said to be simple if each \(j\)-face is the intersection of exactly \((d - j)\)-hyperplanes. In particular, this implies that each vertex is incident to exactly \(d\) facets. Further, each \(j\)-face can be uniquely identified with a subset of \(d - j\) hyperplanes, whose intersection defines the face. Following the same logic as in the previous paragraph, it follows that the number of vertices in such a polytope is naively at most \(O(n^d)\). (Again, we’ll see later that the tight bound is \(O(n^{\lfloor d/2 \rfloor})\).) It follows from the results on polarity that a polytope is simple if and only if its polar is simplicial.

An important observation about simple polytopes is that the local region around each vertex is equivalent to a vertex of a simplex. In particular, if we cut off a vertex of a simple polytope by a hyperplane that is arbitrarily close to the vertex, the piece that has been cut off is a \(d\)-simplex.

It is easy to show that among all polytopes having a fixed number of vertices, simplicial polytopes maximize the number of faces of all higher degrees. (Observe that otherwise there must be degeneracy among the vertices. Perturbing the points breaks the degeneracy, and will generally split faces of higher degree into multiple faces of lower degree.) Dually, among all polytopes having a fixed number of facets, simple polytopes maximize the number of faces of all lower degrees.

Another observation allows us to provide crude bounds on the number of faces of various dimensions. Consider first a simplicial polytope having \(n\) vertices. Each \((j - 1)\)-face can be uniquely identified with a subset of \(j\) points whose convex hull gives this face. Of course, unless the polytope is a simplex, not all of these subsets will give rise to a face. Nonetheless this yields the following naive upper bound on the numbers of faces of various dimensions. By applying the polar transformation we in fact get two bounds, one for simplicial polytopes and one for simple polytopes.

**Lemma:** (Naive bounds)

(i) The number faces of dimension \(j\) of a polytope with \(n\) vertices is at most \(\binom{n}{j+1}\).

(ii) The number of faces of dimension \(j\) of a polytope with \(n\) facets is at most \(\binom{n}{d-j}\).
These naive bounds are not tight. Tight bounds can be derived using more sophisticated relations on the numbers of faces of various dimensions, called the Dehn-Sommerville relations. We will not cover these, but see the discussion below of the Upper Bound Theorem.

The Combinatorics of Polytopes: Let $P$ be a $d$-polytope. For $-1 \leq k \leq d$, let $n_k(P)$ denote the number of $k$-faces of $P$. Clearly $n_{-1}(P) = n_d(P) = 1$. The numbers of faces of other dimensions generally satisfy a number of combinatorial relationships. The simplest of these is called Euler’s relation:

**Theorem:** (Euler’s Relation) Given any $d$-polytope $P$ we have $\sum_{k=-1}^{d} (-1)^k n_k(P) = 0$.

This says that the alternating sum of the numbers of faces sums to 0. For example, a cube has 8 vertices, 12 edges, 6 facets, and together with the faces of dimension $-1$ and $d$ we have

$$-1 + 8 - 12 + 6 - 1 = 0.$$

Although the formal proof of Euler’s relation is rather complex, there is a very easy way to see why its true. First, consider the simplest polytope, namely a $d$-simplex, as the base case. (This is easy to see if you recall that for a simplex $n_j = \binom{d+1}{j+1}$. If you take the expression $(1 - 1)^{d+1}$ and expand it symbolically (as you would for example for $(a + b)^2 = a^2 + 2ab + b^2$) you will get exactly the sum in Euler’s formula. Clearly $(1 - 1)^{d+1} = 0$. The induction part of the proof comes by observing that in order making a complex polytope out of a simple one, essentially involves a series of splitting operation. Every time you split a face of dimension $j$, you do so by adding a face of dimension $j - 1$. Thus, $n_{j-1}$ and $n_j$ each increase by one, and so the value of the alternating sum is unchanged.

Euler’s relation can be used to prove that the convex hull of a set of $n$ points in 3-space has $O(n)$ edges and $O(n)$ faces. However, what happens as dimension increases? We will prove the following theorem. The remarkably simple proof is originally due to Raimund Seidel. We will state the theorem both in its original and dual form.

**The Upper Bound Theorem:** A polytope defined by the convex hull of $n$ points in $\mathbb{R}^d$ has $O(n^{\lfloor d/2 \rfloor})$ facets.

**Upper Bound Theorem (Polar Form):** A polytope defined by the intersection of $n$ halfspaces in $\mathbb{R}^d$ has $O(n^{\lfloor d/2 \rfloor})$ vertices.

**Proof:** It is not hard to show that among all polytopes, simplicial polytopes maximize the number of faces for a given set of vertices and simple polytopes maximize the number of vertices for a given set of faces. We will prove just the polar form of the theorem, and the other will follow by polar equivalence.
Consider a polytope defined by the intersection of $n$ halfspaces in general position. Let us suppose by convention that the $x_d$ axis is the vertical axis. Given a face, its highest vertex and lowest vertices are defined as those having the maximum and minimum $x_d$ coordinates, respectively. (There are no ties if we assume general position.)

The proof is based on a charging argument. We will place a charge at each vertex. We will then move the charge for each vertex to a specially chosen incident face, in such a way that no face receives more than two charges. Finally, we will show that the number of faces that receive charges is at most $O(n^{\lfloor d/2 \rfloor})$.

First, we claim that every vertex $v$ is either the highest or lowest vertex for a $j$-face, where $j \geq \lceil d/2 \rceil$. To see this, recall that for a simple polytope, the neighborhood immediately surrounding any vertex is isomorphic to a simplex. Thus, $v$ is incident to exactly $d$ edges (1-faces). (See Fig. 34.7 for an example in dimension 5.) Consider a horizontal (that is, orthogonal to $x_d$) hyperplane passing through $v$. Since there are $d$ edges in all, at least $\lceil d/2 \rceil$ of these edges must lie on the same side of this hyperplane. (By general position we may assume that no edge lies exactly on the hyperplane.)

As we observed earlier in the lecture, the local neighborhood about each vertex of a simple polytope is isomorphic to a simplex, which implies that there is a face of dimension at least $\lceil d/2 \rceil$ that spans these edges and is incident to $v$. Therefore, $v$ is the lowest or highest vertex for this face. We charge this face for the charge on vertex $v$. Thus, we may charge every vertex of the polytope to face of dimension at least $\lceil d/2 \rceil$, and every such face will be charged at most twice (once by its lowest and once by its highest vertex).

![Figure 34.7](image)

Figure 34.7: Proof of the Upper Bound Theorem in dimension 5. In this case the three edges above $v$ span a 3-face whose lowest vertex is $v$.

All that remains is to count the number of faces that have been charged and multiply by 2. Recalling our earlier lemma on the naive bound on the number of $j$-faces of a simple polytope with $n$ facets is $n \choose d-j$. (Each $j$-face is arises from the intersection of $d-j$ hyperplanes and this is number of $(d-j)$-element subsets of hyperplanes.) Summing this up over all the faces of dimension $\lceil d/2 \rceil$ and higher we find that the number of vertices is at most

$$2 \sum_{j=\lfloor d/2 \rfloor}^{d} {n \choose d-j}.$$ 

By changing the summation index to $k = d-j$ and making the observation that $n \choose k$ is $O(n^k)$, we have that the number of vertices is at most

$$2 \sum_{k=0}^{\lfloor d/2 \rfloor} {n \choose k} = \sum_{k=0}^{\lfloor d/2 \rfloor} O(n^k).$$
This is a geometric series, and so is dominated asymptotically by its largest term. Therefore it follows that the number of charges, that is, the number of vertices is at most

\[ O \left( n^{\lfloor d/2 \rfloor} \right), \]

and this completes the proof.

Is this bound tight? Yes it is. There is a family of polytopes, called *cyclic polytopes*, which match this asymptotic bound. (See Boissonnat and Yvinec for a definition and proof.)