Computational Geometry
Lecture Notes HS 2013

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Preface

These lecture notes are designed to accompany a course on “Computational Geometry” that we teach at the Department of Computer Science, ETH Zürich, in every winter term since 2005. The course has evolved over the years and so have the notes, a first version of which was published in 2008. In the current setting, the course runs over 14 weeks, with three hours of lecture and two hours of exercises each week. In addition, there are three sets of graded homeworks which students have to hand in spread over the course. The target audience are third-year Bachelor or Master students of Mathematics or Computer Science.

The selection of topics and their treatment is somewhat subjective, beyond what we consider essentials driven by the (diverse) research interests within our working group. There is a certain slant towards algorithms rather than data structures and a fair amount of influx from combinatorial geometry. The focus is on low-dimensional Euclidean space (mostly 2D), although we sometimes discuss possible extensions and/or remarkable differences when going to higher dimensions. At the end of each chapter there is a list of questions that we expect our students to be able to answer in the oral exam.

Most parts of these notes have gone through a number of proof-readings, but experience tells that there are always a few mistakes that escape detection. So in case you notice some problem, please let us know, regardless of whether it is a minor typo or punctuation error, a glitch in formulation, or a hole in an argument. This way the issue can be fixed for the next edition and future readers profit from your findings.

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Chapter 1

Fundamentals

1.1 Models of Computation

When designing algorithms, one has to agree on a model of computation according to which these algorithms can be executed. There are various such models, but when it comes to geometry some are more convenient to work with than others. Even using very elementary geometric operations—such as taking the center of a circle defined by three points or computing the length of a given circular arc—the realms of rational and even algebraic numbers are quickly left behind. Representing the resulting real numbers/coordinates would be a rather painful task in, for instance, a Turing machine type model of computation.

Therefore, other models of computation are more prominent in the area of geometric algorithms and data structures. In this course we will be mostly concerned with two models: the Real RAM and the algebraic computation/decision tree model. The former is rather convenient when designing algorithms, because it sort of abstracts from the aforementioned representation issues by simply assuming that it can be done. The latter model typically appears in the context of lower bounds, that is, proofs that certain problems cannot be solved more efficiently than some function depending on the problem size (and possibly some other parameters).

So let us see what these models are in more detail.

Real RAM Model. A memory cell stores a real number (that is what the “Real” stands for\footnote{RAM stands for random access machine, meaning that every memory cell can be accessed in constant time. Not like, say, a list where one always has to start from the first element.}). Any single arithmetic operation (addition, subtraction, multiplication, division, and k-th root, for small constant k) or comparison can be computed in constant time\footnote{In addition, sometimes also logarithms, other analytic functions, indirect addressing (integral), or floor and ceiling are used. As adding some of these operations makes the model more powerful, it is usually specified and emphasized explicitly when an algorithm uses them.}. This is a quite powerful (and somewhat unrealistic) model of computation, as a single real number in principle can encode an arbitrary amount of information. Therefore we
have to ensure that we do not abuse the power of this model. For instance, we may want to restrict the numbers that are manipulated by any single arithmetic operation to be bounded by some fixed polynomial in the numbers that appear in the input.

On the positive side, the real RAM model allows to abstract from the lowlands of numeric and algebraic computation and to concentrate on the algorithmic core from a combinatorial point of view.

But there are also downsides to using such a powerful model. In particular, it may be a challenge to efficiently implement a geometric algorithm designed for the real RAM on an actual computer. With bounded memory there is no way to represent general real numbers explicitly, and operations using a symbolic representation can hardly be considered constant time.

When interested in lower bounds, it is convenient to use a model of computation that encompasses and represents explicitly all possible execution paths of an algorithm. This is what the following model is about.

Algebraic Computation Trees (Ben-Or [1]). A computation is regarded as a binary tree.

- The leaves contain the (possible) results of the computation.
- Every node $v$ with one child has an operation of the form $+, -, \times, \div, \sqrt{\ldots}$ associated to it. The operands of this operation are constant input values, or among the ancestors of $v$ in the tree.
- Every node $v$ with two children has associated to it a branching of the form $> 0$, $\geq 0$, or $= 0$. The branch is with respect to the result of $v$'s parent node. If the expression yields true, the computation continues with the left child of $v$; otherwise, it continues with the right child of $v$.

The term decision tree is used if all of the final results (leaves) are either true or false. If every branch is based on a linear function in the input values, we face a linear decision tree. Analogously one can define, say, quadratic decision trees.

The complexity of a computation or decision tree is the maximum number of vertices along any root-to-leaf path. It is well known that $\Omega(n \log n)$ comparisons are required to sort $n$ numbers. But also for some problems that appear easier than sorting at first glance, the same lower bound holds. Consider, for instance, the following problem.

Element Uniqueness

Input: $\{x_1, \ldots, x_n\} \subset \mathbb{R}$, $n \in \mathbb{N}$.

Output: Is $x_i = x_j$, for some $i, j \in \{1, \ldots, n\}$ with $i \neq j$?
Ben-Or [1] has shown that any algebraic decision tree to solve Element Uniqueness for $n$ elements has complexity $\Omega(n \log n)$.

1.2 Basic Geometric Objects

We will mostly be concerned with the $d$-dimensional Euclidean space $\mathbb{R}^d$, for small $d \in \mathbb{N}$; typically, $d = 2$ or $d = 3$. The basic objects of interest in $\mathbb{R}^d$ are the following.

**Points.** A point $p$, typically described by its $d$ Cartesian coordinates $p = (x_1, \ldots, x_d)$.

<table>
<thead>
<tr>
<th>$p$</th>
<th>$r$</th>
<th>$q$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$(-4, 0)$</td>
<td>$(7, 1)$</td>
<td>$(2, -2)$</td>
</tr>
</tbody>
</table>

**Directions.** A vector $v \in S^{d-1}$ (the $(d-1)$-dimensional unit sphere), typically described by its $d$ Cartesian coordinates $v = (x_1, \ldots, x_d)$, with $||v|| = \sqrt{\sum_{i=1}^{d} x_i^2} = 1$.

**Lines.** A line is a one-dimensional affine subspace. It can be described by two distinct points $p$ and $q$ as the set of all points $r$ that satisfy $r = p + \lambda(q - p)$, for some $\lambda \in \mathbb{R}$.

While any pair of distinct points defines a unique line, a line in $\mathbb{R}^2$ contains infinitely many points and so it may happen that a collection of three or more points lie on a line. Such a collection of points is termed *collinear*.

**Rays.** If we remove a single point from a line and take the closure of one of the connected components, then we obtain a ray. It can be described by two distinct points $p$ and $q$ as the set of all points $r$ that satisfy $r = p + \lambda(q - p)$, for some $\lambda \geq 0$. The orientation of a ray is the direction $(q - p)/||q - p||$.

**Line segment.** A line segment is a compact connected subset of a line. It can be described by two points $p$ and $q$ as the set of all points $r$ that satisfy $r = p + \lambda(q - p)$, for some $\lambda \in [0, 1]$. We will denote the line segment through $p$ and $q$ by $pq$. Depending on the context we may allow or disallow degenerate line segments consisting of a single point only ($p = q$ in the above equation).

**Hyperplanes.** A hyperplane $H$ is a $(d-1)$-dimensional affine subspace. It can be described algebraically by $d + 1$ coefficients $\lambda_1, \ldots, \lambda_{d+1} \in \mathbb{R}$, where $||\lambda_1, \ldots, \lambda_{d+1}|| = 1$, as the set of all points $(x_1, \ldots, x_d)$ that satisfy the linear equation $H : \sum_{i=1}^{d} \lambda_i x_i = \lambda_{d+1}$.

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3Not *collinear*, which refers to a notion in the theory of coalgebras.
If the above equation is converted into an inequality, we obtain the algebraic description of a halfspace (in $\mathbb{R}^2$: halfplane).

**Spheres and balls.** A sphere is the set of all points that are equidistant to a fixed point. It can be described by a point $c$ (center) and a number $\rho \in \mathbb{R}$ (radius) as the set of all points $p$ that satisfy $||p - c|| = \rho$. The ball of radius $\rho$ around $p$ consists of all points $p$ that satisfy $||p - c|| \leq \rho$.

**References**

Chapter 2

Polygons

Although we can think of a line $\ell \subset \mathbb{R}^2$ as an infinite point set that consists of all points in $\mathbb{R}^2$ that are on $\ell$, there still exists a finite description for $\ell$. Such a description is, for instance, provided by the three coefficients $a, b, c \in \mathbb{R}$ of an equation of the form $ax + by = c$, with $(a, b) \neq (0, 0)$. Actually this holds true for all of the fundamental geometric objects that were mentioned in the previous section: Each of them has constant description complexity (or, informally, just size), that is, it can be described by a constant number of parameters.

In this course we will typically deal with objects that are not of constant size. Often these are formed by merely aggregating constant-size objects, for instance, points to form a finite set of points. But sometimes we also demand additional structure that goes beyond aggregation only. Probably the most fundamental geometric objects of this type are what we call polygons. You probably learned this term in school, but what is a polygon precisely? Consider the examples shown in Figure 2.1. Are all of these polygons? If not, where would you draw the line?

Figure 2.1: What is a polygon?

2.1 Classes of Polygons

Obviously, there is not the right answer to such a question and certainly there are different types of polygons. Often the term polygon is used somewhat sloppily in place

\footnote{Unless specified differently, we will always assume that the dimension is (a small) constant. In a high-dimensional space $\mathbb{R}^d$, one has to account for a description complexity of $\Theta(d)$.}
of what we call a simple polygon, defined below.

**Definition 2.1** A simple polygon is a compact region $P \subset \mathbb{R}^2$ that is bounded by a simple closed curve $\gamma : [0, 1] \to \mathbb{R}^2$ that consists of a finite number of line segments. A curve is a continuous map $\gamma : [0, 1] \to \mathbb{R}^2$. A curve $\gamma$ is closed, if $\gamma(0) = \gamma(1)$ and it is simple if it is injective on $[0, 1)$, that is, the curve does not intersect itself.

Out of the examples shown above only Polygon 2.1a is simple. For each of the remaining polygons it is impossible to combine the bounding segments into a simple closed curve.

The term compact for subsets of $\mathbb{R}^d$ means bounded and closed. A subset of $P \subset \mathbb{R}^d$ is bounded, if it is contained in the ball of radius $r$ around the origin, for some finite $r > 0$. Being closed means that the boundary is considered to be part of the polygon.

In order to formally define these terms, let us briefly review a few basic notions from topology.

The standard topology of $\mathbb{R}^d$ is defined in terms of the Euclidean metric. A point $p \in \mathbb{R}^d$ is interior to a set $P \subset \mathbb{R}^d$, if there exists an $\varepsilon$-ball $B_\varepsilon(p) = \{ x \in \mathbb{R}^d : ||x - p|| < \varepsilon \}$ around $p$, for some $\varepsilon > 0$, that is completely contained in $P$. A set is open, if all of its points are interior; and it is closed, if its complement is open.

**Exercise 2.2** Determine for each of the following sets whether they are open or closed in $\mathbb{R}^2$. a) $B_1(0)$ b) $\{(1,0)\}$ c) $\mathbb{R}^2$ d) $\mathbb{R}^2 \setminus \mathbb{Z}^2$ e) $\mathbb{R}^2 \setminus \mathbb{Q}^2$ f) $\{(x, y) : x \in \mathbb{R}, y \geq 0 \}$

**Exercise 2.3** Show that the union of countably many open sets in $\mathbb{R}^d$ is open. Show that the union of a finite number of closed sets in $\mathbb{R}^d$ is closed. (These are two of the axioms that define a topology. So the statements are needed to assert that the metric topology is a topology, indeed.) What follows for intersections of open and closed sets? Finally, show that the union of countably many closed sets in $\mathbb{R}^d$ is not necessarily closed.

The boundary $\partial P$ of a set $P \subset \mathbb{R}^d$ consists of all points that are neither interior to $P$ nor to its complement $\mathbb{R}^d \setminus P$. By definition, for every $p \in \partial P$ every ball $B_\varepsilon(p)$ contains both points from $P$ and from $\mathbb{R}^d \setminus P$. Sometimes one wants to consider a set $P \subset \mathbb{R}^d$ open although it is not. In that case one can resort to the interior $P^\circ$ of $P$ that is formed by the subset of points interior to $P$. Similarly, the closure $\overline{P}$ of $P$ is defined by $\overline{P} = P \cup \partial P$.

Lower-dimensional objects, such as line segments in $\mathbb{R}^2$ or triangles in $\mathbb{R}^3$, do not possess any interior point (because the $\varepsilon$-balls needed around any such point are full-dimensional). Whenever we want to talk about the interior of a lower-dimensional object, we use the qualifier relative and consider it relative to the smallest affine subspace that contains the object.

For instance, the smallest affine subspace that contains a line segment is a line and so the relative interior of a line segment in $\mathbb{R}^2$ consists of all points except the endpoints, just like for an interval in $\mathbb{R}^1$. Similarly, for a triangle in $\mathbb{R}^3$ the smallest affine subspace that contains it is a plane. Hence its relative interior is just the interior of the triangle, considered as a two-dimensional object.
Exercise 2.4 Show that for any \( P \subset \mathbb{R}^d \) the interior \( P^\circ \) is open. (Why is there something to show to begin with?) Show that for any \( P \subset \mathbb{R}^d \) the closure \( \overline{P} \) is closed.

When describing a simple polygon \( P \) it is sufficient to describe only its boundary \( \partial P \). As \( \partial P \) by definition is a simple closed curve \( \gamma \) that consists of finitely many line segments, we can efficiently describe it as a sequence \( p_1, \ldots, p_n \) of points, such that \( \gamma \) is formed by the line segments \( p_1p_2, p_2p_3, \ldots, p_{n-1}p_n, p_np_1 \). These points are referred to as the vertices of the polygon, and the segments connecting them are referred as the edges of the polygon. The set of vertices of a polygon \( P \) is denoted by \( V(P) \), and the set of edges of \( P \) is denoted by \( E(P) \).

Knowing the boundary, it is easy to tell apart the (bounded) interior from the (unbounded) exterior. This is asserted even for much more general curves by the well-known Jordan-Curve Theorem.

Theorem 2.5 (Jordan 1887) Any simple closed curve \( \gamma : [0,1] \rightarrow \mathbb{R}^2 \) divides the plane into exactly two connected components whose common boundary is formed by \( \gamma \).

In full generality, the proof of the deceptively obvious claim is surprisingly difficult. We will not prove it here, the interested reader can find a proof, for instance, in the book of Mohar and Thomassen [11]. There exist different generalizations of the theorem and there also has been some debate about to which degree the original proof of Jordan is actually correct. For simple polygons the situation is easier, though. The essential idea can be worked out algorithmically, which we leave as an exercise.

Exercise 2.6 Describe an algorithm to decide whether a point lies inside or outside of a simple polygon. More precisely, given a simple polygon \( P \subset \mathbb{R}^2 \) as a list of its vertices \((v_1, v_2, \ldots, v_n)\) in counterclockwise order and a query point \( q \in \mathbb{R}^2 \), decide whether \( q \) is inside \( P \), on the boundary of \( P \), or outside. The runtime of your algorithm should be \( O(n) \).

There are good reasons to ask for the boundary of a polygon to form a simple curve: For instance, in the example depicted in Figure 2.1b there are several regions for which it is completely unclear whether they should belong to the interior or to the exterior of the polygon. A similar problem arises for the interior regions in Figure 2.1f. But there are more general classes of polygons that some of the remaining examples fall into. We will discuss only one such class here. It comprises polygons like the one from Figure 2.1d.

Definition 2.7 A region \( P \subset \mathbb{R}^2 \) is a simple polygon with holes if it can be described as \( P = F \setminus \bigcup_{H \in \mathcal{H}} H^\circ \), where \( \mathcal{H} \) is a finite collection of pairwise disjoint simple polygons (called holes) and \( F \) is a simple polygon for which \( F^\circ \supset \bigcup_{H \in \mathcal{H}} H \).

The way this definition heavily depends on the notion of simple polygons makes it straightforward to derive a similar trichotomy as the Jordan Curve Theorem provides for simple polygons, that is, every point in the plane is either inside, or on the boundary, or outside of \( P \) (exactly one of these three).
2.2 Polygon Triangulation

From a topological point of view, a simple polygon is nothing but a disk and so it is a very elementary object. But geometrically a simple polygon can be—as if mocking the label we attached to it—a pretty complicated shape, see Figure 2.2 for an example. While there is an easy and compact one-dimensional representation in terms of the boundary, as a sequence of vertices/points, it is often desirable to work with a more structured representation of the whole two-dimensional shape.

![Figure 2.2: A simple (?) polygon.](image)

For instance, it is not straightforward to compute the area of a general simple polygon. In order to do so, one usually describes the polygon in terms of simpler geometric objects, for which computing the area is easy. Good candidates for such shapes are triangles, rectangles, and trapezoids. Indeed, it is not hard to show that every simple polygon admits a “nice” partition into triangles, which we call a triangulation.

**Definition 2.8** A triangulation of a simple polygon $P$ is a collection $\mathcal{T}$ of triangles, such that

1. $P = \bigcup_{T \in \mathcal{T}} T$;
2. $V(P) = \bigcup_{T \in \mathcal{T}} V(T)$; and
3. for every distinct pair $T, U \in \mathcal{T}$, the intersection $T \cap U$ is either a common vertex, or a common edge, or empty.

**Exercise 2.9** Show that each condition in Definition 2.8 is necessary in the following sense: Give an example of a non-triangulation that would form a triangulation if the condition was omitted. Is the definition equivalent if (3) is replaced by $T^c \cap U^c = \emptyset$, for every distinct pair $T, U \in \mathcal{T}$?

If we are given a triangulation of a simple polygon $P$ it is easy to compute the area of $P$ by simply summing up the area of all triangles from $\mathcal{T}$. Triangulations are an incredibly useful tool in planar geometry, and one reason for their importance is that every simple polygon admits one.

**Theorem 2.10** Every simple polygon has a triangulation.
Proof. Let $P$ be a simple polygon on $n$ vertices. We prove the statement by induction on $n$. For $n = 3$ we face a triangle $P$ that is a triangulation by itself. For $n > 3$ consider the lexicographically smallest vertex $v$ of $P$, that is, among all vertices of $P$ with a smallest $x$-coordinate the one with smallest $y$-coordinate. Denote the neighbors of $v$ (next vertices) along $\partial P$ by $u$ and $w$. Consider the line segment $uw$. We distinguish two cases.

Case 1: except for its endpoints $u$ and $w$, the segment $uw$ lies completely in $P^\circ$. Then $uw$ splits $P$ into two smaller polygons, the triangle $uvw$ and a simple polygon $P'$ on $n - 1$ vertices (Figure 2.3a). By the inductive hypothesis, $P'$ has a triangulation that together with $T$ yields a triangulation of $P$.

Case 2: the relative interior of $uw$ does not lie completely in $P^\circ$ (Figure 2.3b). By choice of $v$, the polygon $P$ is contained in the closed halfplane to the right of the vertical line through $v$. Therefore, as the segments $uw$ and $vw$ are part of a simple closed curve defining $\partial P$, every point sufficiently close to $v$ and between the rays $vu$ and $vw$ must be in $P^\circ$. On the other hand, since $uw \nsubseteq P^\circ$, there is some point from $\partial P$ in the interior of the triangle $T = uvw$ (by the choice of $v$ the points $u, v, w$ are not collinear and so $T$ is a triangle, indeed) or on the line segment $uw$. In particular, as $\partial P$ is composed of line segments, there is a vertex of $P$ in $T^\circ$ or on $uw$ (otherwise, a line segment would have to intersect the line segment $uw$ twice, which is impossible). Let $p$ denote a leftmost such vertex. Then the open line segment $vp$ is contained in $T^\circ$ and, thus, it splits $P$ into two polygons $P_1$ and $P_2$ on less than $n$ vertices each (in one of them, $u$ does not appear as a vertex, whereas $w$ does not appear as a vertex in the other). By the inductive hypothesis, both $P_1$ and $P_2$ have triangulations and their union yields a triangulation of $P$. 

The configuration from Case 1 above is called an ear: Three consecutive vertices $u, v, w$ of a simple polygon $P$ such that the relative interior of $uw$ lies in $P^\circ$. In fact, we could have skipped the analysis for Case 2 by referring to the following theorem.

Theorem 2.11 (Meisters [9, 10]) Every simple polygon that is not a triangle has two non-overlapping ears, that is, two ears $A$ and $B$ such that $A^\circ \cap B^\circ = \emptyset$.  

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But knowing Theorem 2.10 we can obtain Theorem 2.11 as a direct consequence of the following.

**Theorem 2.12** Every triangulation of a simple polygon on \( n \geq 4 \) vertices contains at least two (triangles that are) ears.

**Exercise 2.13** Prove Theorem 2.12.

**Exercise 2.14** Let \( P \) be a simple polygon with vertices \( v_1, v_2, \ldots, v_n \) (in counterclockwise order), where \( v_i \) has coordinates \((x_i, y_i)\). Show that the area of \( P \) is

\[
\frac{1}{2} \sum_{i=1}^{n} x_{i+1}y_i - x_iy_{i+1},
\]

where \((x_{n+1}, y_{n+1}) = (x_1, y_1)\).

The number of edges and triangles in a triangulation of a simple polygon are completely determined by the number of vertices, as the following simple lemma shows.

**Lemma 2.15** Every triangulation of a simple polygon on \( n \geq 3 \) vertices consists of \( n - 2 \) triangles and \( 2n - 3 \) edges.

**Proof.** Proof by induction on \( n \). The statement is true for \( n = 3 \). For \( n > 3 \) consider a simple polygon \( P \) on \( n \) vertices and an arbitrary triangulation \( T \) of \( P \). Any edge \( uv \) in \( T \) that is not an edge of \( P \) (and there must be such an edge because \( P \) is not a triangle) partitions \( P \) into two polygons \( P_1 \) and \( P_2 \) with \( n_1 \) and \( n_2 \) vertices, respectively. Since \( n_1, n_2 < n \) we conclude by the inductive hypothesis that \( T \) partitions \( P_1 \) into \( n_1 - 2 \) triangles and \( P_2 \) into \( n_2 - 2 \) triangles, using \( 2n_1 - 3 \) and \( 2n_2 - 3 \) edges, respectively.

All vertices of \( P \) appear in exactly one of \( P_1 \) or \( P_2 \), except for \( u \) and \( v \), which appear in both. Therefore \( n_1 + n_2 = n + 2 \) and so the number of triangles in \( T \) is \((n_1 - 2) + (n_2 - 2) = (n_1 + n_2) - 4 = n + 2 - 4 = n - 2 \). Similarly, all edges of \( T \) appear in exactly one of \( P_1 \) or \( P_2 \), except for the edge \( uv \), which appears in both. Therefore the number of edges in \( T \) is \((2n_1 - 3) + (2n_2 - 3) - 1 = 2(n_1 + n_2) - 7 = 2(n + 2) - 7 = 2n - 3 \). \( \square \)

The universal presence of triangulations is something particular about the plane: The natural generalization of Theorem 2.10 to dimension three and higher does not hold. What is this generalization, anyway?

**Tetrahedralizations in \( \mathbb{R}^3 \).** A simple polygon is a planar object that is a topological disk that is locally bounded by patches of lines. The corresponding term in \( \mathbb{R}^3 \) is a polyhedron, and although we will not formally define it here yet, a literal translation of the previous sentence yields an object that topologically is a ball and is locally bounded by patches of planes. A triangle in \( \mathbb{R}^2 \) corresponds to a tetrahedron in \( \mathbb{R}^3 \) and a tetrahedralization is a nice partition into tetrahedra, where "nice" means that the union of the tetrahedra covers the object, the vertices of the tetrahedra are vertices of the polyhedron, and any
two distinct tetrahedra intersect in either a common triangular face, or a common edge, or a common vertex, or not at all.\footnote{These “nice” subdivisions can be defined in an abstract combinatorial setting, where they are called simplicial complexes.}

Unfortunately, there are polyhedra in $\mathbb{R}^3$ that do not admit a tetrahedralization. The following construction is due to Schönhardt \cite{schonhardt1928}. It is based on a triangular prism, that is, two congruent triangles placed in parallel planes where the corresponding sides of both triangles are connected by a rectangle (Figure 2.4a). Then one triangle is twisted/rotated slightly within its plane. As a consequence, the rectangular faces are not plane anymore, but they obtain an inward dent along their diagonal in direction of the rotation (Figure 2.4b). The other (former) diagonals of the rectangular faces—labeled $ab'$, $bc'$, and $ca'$ in Figure 2.4b—are now epigonals, that is, they lie in the exterior of the polyhedron. Since these epigonals are the only edges between vertices that are not part of the polyhedron, there is no way to add edges to form a tetrahedron for a subdivision. Clearly the polyhedron is not a tetrahedron by itself, and so we conclude that it does not admit a subdivision into tetrahedra without adding new vertices. If adding new vertices—so-called Steiner vertices—is allowed, then there is no problem to construct a tetrahedralization, and this holds true in general.

**Figure 2.4:** The Schönhardt polyhedron cannot be subdivided into tetrahedra without adding new vertices.

**Exercise 2.16** Convert Theorem 2.10 into an $O(n^2)$ time algorithm to construct a triangulation for a given simple polygon on $n$ vertices.

The runtime achieved by the straightforward application of Theorem 2.10 is not optimal. We will revisit this question at several times during this course and discuss improved algorithms for the problem of triangulating a simple polygon.
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The best (in terms of worst-case runtime) algorithm known due to Chazelle [4] computes a triangulation in linear time. But this algorithm is very complicated and we will not discuss it here. There is also a somewhat simpler randomized algorithm to compute a triangulation in expected linear time [2], which we will not discuss in detail, either. Instead you will later see a much simpler algorithm with a pretty-close-to linear runtime bound. The question of whether there exists a simple (which is not really a well-defined term, of course, except that Chazelle’s Algorithm does not qualify) deterministic linear time algorithm to triangulate a simple polygon remains open [7].

Polygons with holes. It is interesting to note that the complexity of the problem changes to $\Theta(n \log n)$, if the polygon may contain holes [3]. This means that there is an algorithm to construct a triangulation for a given simple polygon with holes on a total of $n$ vertices (counting both the vertices on the outer boundary and those of holes) in $O(n \log n)$ time. But there is also a lower bound of $\Omega(n \log n)$ operations that holds in all models of computation in which there exists the corresponding lower bound for comparison-based sorting. This difference in complexity is a very common pattern: There are many problems that are (sometimes much) harder for simple polygons with holes than for simple polygons. So maybe the term “simple” has some justification, after all...

General triangle covers. What if we drop the “niceness” conditions required for triangulations and just want to describe a given simple polygon as a union of triangles? It turns out this is a rather drastic change and, for instance, it is unlikely that we can efficiently find an optimal/minimal description of this type: Christ has shown [5] that it is NP-hard to decide whether for a simple polygon $P$ on $n$ vertices and a positive integer $k$, there exists a set of at most $k$ triangles whose union is $P$. In fact, the problem is not even known to be in NP, because it is not clear whether the coordinates of solutions can always be encoded compactly.

2.3 The Art Gallery Problem

In 1973 Victor Klee posed the following question: “How many guards are necessary, and how many are sufficient to patrol the paintings and works of art in an art gallery with $n$ walls?” From a geometric point of view, we may think of an “art gallery with $n$ walls” as a simple polygon bounded by $n$ edges, that is, a simple polygon $P$ with $n$ vertices. And a guard can be modeled as a point where we imagine the guard to stand and observe everything that is in sight. In sight, finally, refers to the walls of the gallery (edges of the polygon) that are opaque and, thus, prevent a guard to see what is behind. In other words, a guard (point) $g$ can watch over every point $p \in P$, for which the line segment $gp$ lies completely in $P^o$, see Figure 2.5.

It is not hard to see that $\lfloor n/3 \rfloor$ guards are necessary in general.
2.3. The Art Gallery Problem

Figure 2.5: The region that a guard g can observe.

Exercise 2.17 Describe a family \((P_n)_{n \geq 3}\) of simple polygons such that \(P_n\) has \(n\) vertices and at least \(\lfloor n/3 \rfloor\) guards are needed to guard it.

What is more surprising: \(\lfloor n/3 \rfloor\) guards are always sufficient as well. Chvátal \[6\] was the first to prove that, but then Fisk \[8\] gave a much simpler proof using—you may have guessed it—triangulations. Fisk’s proof was considered so beautiful that it was included into “Proofs from THE BOOK” \[1\], a collection inspired by Paul Erdős’ belief in “a place where God keeps aesthetically perfect proofs”. The proof is based on the following lemma.

Lemma 2.18 Every triangulation of a simple polygon is 3-colorable. That is, each vertex can be assigned one of three colors in such a way that adjacent vertices receive different colors.

Proof. Induction on \(n\). For \(n = 3\) the statement is obvious. For \(n > 3\), by Theorem 2.12 the triangulation contains an ear \(uvw\). Cutting off the ear creates a triangulation of a polygon on \(n - 1\) vertices, which by the inductive hypothesis admits a 3-coloring. Now whichever two colors the vertices \(u\) and \(w\) receive in this coloring, there remains a third color to be used for \(v\).

Theorem 2.19 (Fisk \[8\]) Every simple polygon on \(n\) vertices can be guarded using at most \(\lfloor n/3 \rfloor\) guards.

Proof. Consider a triangulation of the polygon and a 3-coloring of the vertices as ensured by Lemma 2.18. Take the smallest color class, which clearly consists of at most \(\lfloor n/3 \rfloor\) vertices, and put a guard at each vertex. As every point of the polygon is contained in at least one triangle and every triangle has exactly one vertex in the guarding set, the whole polygon is guarded.

Questions

1. What is a simple polygon/a simple polygon with holes Explain the definitions and provide some examples of members and non-members of the respective classes.
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Figure 2.6: A triangulation of a simple polygon on 17 vertices and a 3-coloring of it. The vertices shown solid orange form the smallest color class and guard the polygon using $\lfloor 17/3 \rfloor = 5$ guards.

For a given polygon you should be able to tell which of these classes it belongs to or does not belong to and argue why this is the case.

2. **What is a closed/open/bounded set in $\mathbb{R}^d$? What is the interior/closure of a point set?** Explain the definitions and provide some illustrative examples. For a given set you should be able to argue which of the properties mentioned it possesses.

3. **What is a triangulation of a simple polygon? Does it always exist?** Explain the definition and provide some illustrative examples. Present the proof of Theorem 2.10 in detail.

4. **How about higher dimensional generalizations? Can every polyhedron in $\mathbb{R}^3$ be nicely subdivided into tetrahedra?** Explain Schönhardt’s construction.

5. **How many points are needed to guard a simple polygon?** Present the proofs of Theorem 2.12, Lemma 2.18 and Theorem 2.19 in detail.

References


2.3. The Art Gallery Problem


Chapter 3

Convex Hull

There exists an incredible variety of point sets and polygons. Among them, some have certain properties that make them “nicer” than others in some respect. For instance, look at the two polygons shown below.

![Convex Polygon](image1.png) ![Non-Convex Polygon](image2.png)

(a) A convex polygon. (b) A non-convex polygon.

Figure 3.1: Examples of polygons: Which do you like better?

As it is hard to argue about aesthetics, let us take a more algorithmic stance. When designing algorithms, the polygon shown on the left appears much easier to deal with than the visually and geometrically more complex polygon shown on the right. One particular property that makes the left polygon nice is that one can walk between any two vertices along a straight line without ever leaving the polygon. In fact, this statement holds true not only for vertices but for any two points within the polygon. A polygon or, more generally, a set with this property is called convex.

**Definition 3.1** A set $P \subseteq \mathbb{R}^d$ is convex if $\overline{pq} \subseteq P$, for any $p, q \in P$.

An alternative, equivalent way to phrase convexity would be to demand that for every line $\ell \subseteq \mathbb{R}^d$ the intersection $\ell \cap P$ be connected. The polygon shown in Figure 3.1b is not convex because there are some pairs of points for which the connecting line segment is not completely contained within the polygon. An immediate consequence of the definition is the following
Observation 3.2. For any family \((P_i)_{i \in I}\) of convex sets, the intersection \(\bigcap_{i \in I} P_i\) is convex.

Indeed there are many problems that are comparatively easy to solve for convex sets but very hard in general. We will encounter some particular instances of this phenomenon later in the course. However, not all polygons are convex and a discrete set of points is never convex, unless it consists of at most one point only. In such a case it is useful to make a given set \(P\) convex, that is, approximate \(P\) with or, rather, encompass \(P\) within a convex set \(H \supseteq P\). Ideally, \(H\) differs from \(P\) as little as possible, that is, we want \(H\) to be a smallest convex set enclosing \(P\).

At this point let us step back for a second and ask ourselves whether this wish makes sense at all: Does such a set \(H\) always exist? Fortunately, we are on the safe side because the whole space \(\mathbb{R}^d\) is certainly convex. It is less obvious, but we will see below that \(H\) is actually unique. Therefore it is legitimate to refer to \(H\) as the smallest convex set enclosing \(P\) or—shortly—the convex hull of \(P\).

3.1 Convexity

In this section we will derive an algebraic characterization of convexity. Such a characterization allows to investigate convexity using the machinery from linear algebra.

Consider \(P \subseteq \mathbb{R}^d\). From linear algebra courses you should know that the linear hull

\[
\text{lin}(P) := \{ q \mid q = \sum \lambda_i p_i \land \forall i : p_i \in P, \lambda_i \in \mathbb{R} \}
\]

is the set of all linear combinations of \(P\) (smallest linear subspace containing \(P\)). For instance, if \(P = \{p\} \subseteq \mathbb{R}^2 \setminus \{0\}\) then \(\text{lin}(P)\) is the line through \(p\) and the origin.

Similarly, the affine hull

\[
\text{aff}(P) := \{ q \mid q = \sum \lambda_i p_i \land \sum \lambda_i = 1 \land \forall i : p_i \in P, \lambda_i \in \mathbb{R} \}
\]

is the set of all affine combinations of \(P\) (smallest affine subspace containing \(P\)). For instance, if \(P = \{p, q\} \subseteq \mathbb{R}^2\) and \(p \neq q\) then \(\text{aff}(P)\) is the line through \(p\) and \(q\).

It turns out that convexity can be described in a very similar way algebraically, which leads to the notion of convex combinations.

Proposition 3.3. A set \(P \subseteq \mathbb{R}^d\) is convex if and only if \(\sum_{i=1}^n \lambda_i p_i \in P\), for all \(n \in \mathbb{N}\), \(p_1, \ldots, p_n \in P\), and \(\lambda_1, \ldots, \lambda_n \geq 0\) with \(\sum_{i=1}^n \lambda_i = 1\).

Proof. “\(\Rightarrow\)”: obvious with \(n = 2\).

“\(\Leftarrow\)”: Induction on \(n\). For \(n = 1\) the statement is trivial. For \(n \geq 2\), let \(p_1 \in P\) and \(\lambda_1 \geq 0\), for \(1 \leq i \leq n\), and assume \(\sum_{i=1}^n \lambda_i = 1\). We may suppose that \(\lambda_i > 0\), for all \(i\). (Simply omit those points whose coefficient is zero.) We need to show that \(\sum_{i=1}^n \lambda_i p_i \in P\).
3.1. Convexity

Define \( \lambda = \sum_{i=1}^{n-1} \lambda_i \) and for \( 1 \leq i \leq n-1 \) set \( \mu_i = \lambda_i/\lambda \). Observe that \( \mu_i \geq 0 \) and \( \sum_{i=1}^{n-1} \mu_i = 1 \). By the inductive hypothesis, \( q := \sum_{i=1}^{n-1} \mu_i p_i \in P \), and thus by convexity of \( P \) also \( \lambda q + (1 - \lambda) p_n \in P \). We conclude by noting that \( \lambda q + (1 - \lambda) p_n = \lambda \sum_{i=1}^{n-1} \mu_i p_i + \lambda_n p_n = \sum_{i=1}^{n} \lambda_i p_i \).

**Definition 3.4** The convex hull \( \text{conv}(P) \) of a set \( P \subseteq \mathbb{R}^d \) is the intersection of all convex supersets of \( P \).

At first glance this definition is a bit scary: There may be a whole lot of supersets for any given \( P \) and it not clear that taking the intersection of all of them yields something sensible to work with. However, by Observation 3.2 we know that the resulting set is convex, at least. The missing bit is provided by the following proposition, which characterizes the convex hull in terms of exactly those convex combinations that appeared in Proposition 3.3 already.

**Proposition 3.5** For any \( P \subseteq \mathbb{R}^d \) we have

\[
\text{conv}(P) = \left\{ \sum_{i=1}^{n} \lambda_i p_i \mid n \in \mathbb{N} \land \sum_{i=1}^{n} \lambda_i = 1 \land \forall i \in \{1, \ldots, n\} : \lambda_i \geq 0 \land p_i \in P \right\}.
\]

The elements of the set on the right hand side are referred to as convex combinations of \( P \).

**Proof.** “\( \supseteq \)”: Consider a convex set \( C \supseteq P \). By Proposition 3.3 (only-if direction) the right hand side is contained in \( C \). As \( C \) was arbitrary, the claim follows.

“\( \subseteq \)”: Denote the set on the right hand side by \( R \). Clearly \( R \supseteq P \). We show that \( R \) forms a convex set. Let \( p = \sum_{i=1}^{n} \lambda_i p_i \) and \( q = \sum_{i=1}^{n} \mu_i p_i \) be two convex combinations. (We may suppose that both \( p \) and \( q \) are expressed over the same \( p_i \) by possibly adding some terms with a coefficient of zero.)

Then for \( \lambda \in [0,1] \) we have \( \lambda p + (1 - \lambda) q = \sum_{i=1}^{n} (\lambda \lambda_i + (1 - \lambda) \mu_i) p_i \in R \), as \( \lambda \lambda_i + (1 - \lambda) \mu_i \geq 0 \), for all \( 1 \leq i \leq n \), and \( \sum_{i=1}^{n} (\lambda \lambda_i + (1 - \lambda) \mu_i) = \lambda + (1 - \lambda) = 1 \). \( \square \)

In linear algebra the notion of a basis in a vector space plays a fundamental role. In a similar way we want to describe convex sets using as few entities as possible, which leads to the notion of extremal points, as defined below.

**Definition 3.6** The convex hull of a finite point set \( P \subseteq \mathbb{R}^d \) forms a convex polytope. Each \( p \in P \) for which \( p \notin \text{conv}(P \setminus \{p\}) \) is called a vertex of \( \text{conv}(P) \). A vertex of \( \text{conv}(P) \) is also called an extremal point of \( P \). A convex polytope in \( \mathbb{R}^2 \) is called a convex polygon.

Essentially, the following proposition shows that the term vertex above is well defined.

**Proposition 3.7** A convex polytope in \( \mathbb{R}^d \) is the convex hull of its vertices.
Proof. Let \( P = \{p_1, \ldots, p_n\}, \ n \in \mathbb{N}\), such that without loss of generality \( p_1, \ldots, p_k\) are the vertices of \( \mathcal{P} := \text{conv}(P)\). We prove by induction on \( n \) that \( \text{conv}(p_1, \ldots, p_n) \subseteq \text{conv}(p_1, \ldots, p_k)\). For \( n = k \) the statement is trivial.

For \( n > k\), \( p_n\) is not a vertex of \( \mathcal{P} \) and hence \( p_n\) can be expressed as a convex combination \( p_n = \sum_{i=1}^{n-1} \lambda_i p_i \). Thus for any \( x \in \mathcal{P} \) we can write \( x = \sum_{i=1}^{n-1} \mu_i p_i = \sum_{i=1}^{n-1} \mu_i p_i + \mu_n \sum_{i=1}^{n-1} \lambda_i p_i - \sum_{i=1}^{n-1} (\mu_i + \mu_n \lambda_i) p_i \). As \( \sum_{i=1}^{n-1} (\mu_i + \mu_n \lambda_i) = 1 \), we conclude inductively that \( x \in \text{conv}(p_1, \ldots, p_{n-1}) \subseteq \text{conv}(p_1, \ldots, p_k)\). \( \square \)

### 3.2 Classical Theorems for Convex Sets

Next we will discuss a few fundamental theorems about convex sets in \( \mathbb{R}^d \). The proofs typically use the algebraic characterization of convexity and then employ some techniques from linear algebra.

**Theorem 3.8 (Radon [8])** Any set \( P \subset \mathbb{R}^d \) of \( d + 2 \) points can be partitioned into two disjoint subsets \( P_1 \) and \( P_2 \) such that \( \text{conv}(P_1) \cap \text{conv}(P_2) \neq \emptyset \).

**Proof.** Let \( P = \{p_1, \ldots, p_{d+2}\} \). No more than \( d + 1 \) points can be affinely independent in \( \mathbb{R}^d \). Hence suppose without loss of generality that \( p_{d+2} \) can be expressed as an affine combination of \( p_1, \ldots, p_{d+1} \), that is, there exist \( \lambda_1, \ldots, \lambda_{d+1} \in \mathbb{R} \) with \( \sum_{i=1}^{d+1} \lambda_i = 1 \) and \( \sum_{i=1}^{d+1} \lambda_i p_i = p_{d+2} \). Let \( P_1 \) be the set of all points \( p_i \) for which \( \lambda_i \) is positive and let \( P_2 = P \setminus P_1 \). Then setting \( \lambda_{d+2} = -1 \) we can write \( \sum_{p_i \in P_1} \lambda_i p_i = \sum_{p_i \in P_2} \lambda_i p_i \), where all coefficients on both sides are non-negative. Renormalizing by \( \mu_i = \lambda_i / \mu \) and \( \nu_i = \lambda_i / \nu \), where \( \mu = \sum_{p_i \in P_1} \lambda_i \) and \( \nu = - \sum_{p_i \in P_2} \lambda_i \), yields convex combinations \( \sum_{p_i \in P_1} \mu_i p_i = \sum_{p_i \in P_2} \nu_i p_i \) that describe a common point of \( \text{conv}(P_1) \) and \( \text{conv}(P_2) \). \( \square \)

**Theorem 3.9 (Helly)** Consider a collection \( \mathcal{C} = \{C_1, \ldots, C_n\} \) of \( n \geq d + 1 \) convex subsets of \( \mathbb{R}^d \), such that any \( d + 1 \) pairwise distinct sets from \( \mathcal{C} \) have non-empty intersection. Then also the intersection \( \bigcap_{i=1}^{n} C_i \) of all sets from \( \mathcal{C} \) is non-empty.

**Proof.** Induction on \( n \). The base case \( n = d + 1 \) holds by assumption. Hence suppose that \( n \geq d + 2 \). Consider the sets \( D_i = \bigcap_{j \neq i} C_j \), for \( i \in \{1, \ldots, n\} \). As \( D_i \) is an intersection of \( n - 1 \) sets from \( \mathcal{C} \), by the inductive hypothesis we know that \( D_i \neq \emptyset \). Therefore we can find some point \( p_i \in D_i \), for each \( i \in \{1, \ldots, n\} \). Now by Theorem 3.8 the set \( P = \{p_1, \ldots, p_n\} \) can be partitioned into two disjoint subsets \( P_1 \) and \( P_2 \) such that \( \text{conv}(P_1) \cap \text{conv}(P_2) \neq \emptyset \). We claim that any point \( p \in \text{conv}(P_1) \cap \text{conv}(P_2) \) also lies in \( \bigcap_{i=1}^{n} C_i \), which completes the proof.

Consider some \( C_i \), for \( i \in \{1, \ldots, n\} \). By construction \( D_j \subseteq C_i \), for \( j \neq i \). Thus \( p_i \) is the only point from \( P \) that may not be in \( C_i \). As \( p_i \) is part of only one of \( P_1 \) or \( P_2 \), say, of \( P_1 \), we have \( P_2 \subseteq C_i \). The convexity of \( C_i \) implies \( \text{conv}(P_2) \subseteq C_i \) and, therefore, \( p \in C_i \). \( \square \)
Theorem 3.10 (Carathéodory [3]) For any $P \subset \mathbb{R}^d$ and $q \in \text{conv}(P)$ there exist $k \leq d + 1$ points $p_1, \ldots, p_k \in P$ such that $q \in \text{conv}(p_1, \ldots, p_k)$.

Exercise 3.11 Prove Theorem 3.10.

Theorem 3.12 (Separation Theorem) Any two compact convex sets $C, D \subset \mathbb{R}^d$ with $C \cap D = \emptyset$ can be separated strictly by a hyperplane, that is, there exists a hyperplane $h$ such that $C$ and $D$ lie in the opposite open halfspaces bounded by $h$.

Proof. Consider the distance function $\delta : C \times D \to \mathbb{R}$ with $(c, d) \mapsto ||c - d||$. Since $C \times D$ is compact and $\delta$ is continuous and strictly bounded from below by 0, the function $\delta$ attains its minimum at some point $(c_0, d_0) \in C \times D$ with $\delta(c_0, d_0) > 0$. Let $h$ be the hyperplane perpendicular to the line segment $c_0d_0$ and passing through the midpoint of $c_0$ and $d_0$.

If there was a point, say, $c'$ in $C \cap h$, then by convexity of $C$ the whole line segment $c_0c'$ lies in $C$ and some point along this segment is closer to $d_0$ than is $c_0$, in contradiction to the choice of $c_0$. The figure shown to the right depicts the situation in $\mathbb{R}^2$. If, say, $C$ has points on both sides of $h$, then by convexity of $C$ it has also a point on $h$, but we just saw that there is no such point. Therefore, $C$ and $D$ must lie in different open halfspaces bounded by $h$. \qed

The statement above is wrong for arbitrary (not necessarily compact) convex sets. However, if the separation is not required to be strict (the hyperplane may intersect the sets), then such a separation always exists, with the proof being a bit more involved (cf. [7], but also check the errata on Matoušek’s webpage).

Exercise 3.13 Show that the Separation Theorem does not hold in general, if not both of the sets are convex.

Exercise 3.14 Prove or disprove:

(a) The convex hull of a compact subset of $\mathbb{R}^d$ is compact.

(b) The convex hull of a closed subset of $\mathbb{R}^d$ is closed.

Altogether we obtain various equivalent definitions for the convex hull, summarized in the following theorem.

Theorem 3.15 For a compact set $P \subset \mathbb{R}^d$ we can characterize $\text{conv}(P)$ equivalently as one of

(a) the smallest (w. r. t. set inclusion) convex subset of $\mathbb{R}^d$ that contains $P$;

(b) the smallest (w. r. t. measure) convex subset of $\mathbb{R}^d$ that contains $P$;

(c) the smallest (w. r. t. volume) convex subset of $\mathbb{R}^d$ that contains $P$;
(b) the set of all convex combinations of points from \( P \);
(c) the set of all convex combinations formed by \( d + 1 \) or fewer points from \( P \);
(d) the intersection of all convex supersets of \( P \);
(e) the intersection of all closed halfspaces containing \( P \).

Exercise 3.16 Prove Theorem 3.15.

3.3 Planar Convex Hull

Although we know by now what is the convex hull of point set, it is not yet clear how to construct it algorithmically. As a first step, we have to find a suitable representation for convex hulls. In this section we focus on the problem in \( \mathbb{R}^2 \), where the convex hull of a finite point set forms a convex polygon. A convex polygon is easy to represent, for instance, as a sequence of its vertices in counterclockwise orientation. In higher dimensions finding a suitable representation for convex polytopes is a much more delicate task.

Problem 3.17 (Convex hull)

Input: \( P = \{ p_1, \ldots, p_n \} \subset \mathbb{R}^2, \ n \in \mathbb{N} \).

Output: Sequence \( (q_1, \ldots, q_h) \), \( 1 \leq h \leq n \), of the vertices of \( \text{conv}(P) \) (ordered counterclockwise).

![Figure 3.2: Convex Hull of a set of points in \( \mathbb{R}^2 \).](image)

Another possible algorithmic formulation of the problem is to ignore the structure of the convex hull and just consider it as a point set.
Problem 3.18 (Extremal points)

Input: \( P = \{p_1, \ldots, p_n\} \subset \mathbb{R}^2, n \in \mathbb{N} \).

Output: Set \( Q \subseteq P \) of the vertices of \( \text{conv}(P) \).

Degeneracies. A couple of further clarifications regarding the above problem definitions are in order.

First of all, for efficiency reasons an input is usually specified as a sequence of points. Do we insist that this sequence forms a set or are duplications of points allowed?

What if three points are collinear? Are all of them considered extremal? According to our definition from above, they are not and that is what we will stick to. But note that there may be cases where one wants to include all such points, nevertheless.

By the Separation Theorem, every extremal point \( p \) can be separated from the convex hull of the remaining points by a halfplane. If we take such a halfplane and translate its defining line such that it passes through \( p \), then all points from \( P \) other than \( p \) should lie in the resulting open halfplane. In \( \mathbb{R}^2 \) it turns out convenient to work with the following “directed” reformulation.

Proposition 3.19 A point \( p \in P = \{p_1, \ldots, p_n\} \subset \mathbb{R}^2 \) is extremal for \( P \) \iff there is a directed line \( g \) through \( p \) such that \( P \setminus \{p\} \) is to the left of \( g \).

The interior angle at a vertex \( v \) of a polygon \( P \) is the angle between the two edges of \( P \) incident to \( v \) whose corresponding angular domain lies in \( P^\circ \). If this angle is smaller than \( \pi \), the vertex is called convex; if the angle is larger than \( \pi \), the vertex is called reflex. For instance, the vertex labeled \( c \) in the polygon depicted to the right is a convex vertex, whereas the vertex labeled \( r \) is a reflex vertex.

Exercise 3.20

A set \( S \subset \mathbb{R}^2 \) is star-shaped if there exists a point \( c \in S \), such that for every point \( p \in S \) the line segment \( cp \) is contained in \( S \). A simple polygon with exactly three convex vertices is called a pseudotriangle (see the example shown on the right).

In the following we consider subsets of \( \mathbb{R}^2 \). Prove or disprove:

a) Every convex vertex of a simple polygon lies on its convex hull.

b) Every star-shaped set is convex.

c) Every convex set is star-shaped.

d) The intersection of two convex sets is convex.
e) The union of two convex sets is convex.

f) The intersection of two star-shaped sets is star-shaped.

g) The intersection of a convex set with a star-shaped set is star-shaped.

h) Every triangle is a pseudotriangle.

i) Every pseudotriangle is star-shaped.

3.4 Trivial algorithms

One can compute the extremal points using Carathéodory’s Theorem as follows: Test for every point \( p \in P \) whether there are \( q, r, s \in P \setminus \{p\} \) such that \( p \) is inside the triangle with vertices \( q, r, \) and \( s \). Runtime \( O(n^4) \).

Another option, inspired by the Separation Theorem: test for every pair \( (p, q) \in P^2 \) whether all points from \( P \setminus \{p, q\} \) are to the left of the directed line through \( p \) and \( q \) (or on the line segment \( pq \)). Runtime \( O(n^3) \).

Exercise 3.21 Let \( P = (p_0, \ldots, p_{n-1}) \) be a sequence of \( n \) points in \( \mathbb{R}^2 \). Someone claims that you can check by means of the following algorithm whether or not \( P \) describes the boundary of a convex polygon in counterclockwise order:

```c
bool is_convex(p0, \ldots, pn-1) {
    for i = 0, \ldots, n - 1:
        if (p_i, p_{(i+1) \mod n}, p_{(i+2) \mod n}) form a right turn:
            return false;
    return true;
}
```

Disprove the claim and describe a correct algorithm to solve the problem.

Exercise 3.22 Let \( P \subset \mathbb{R}^2 \) be a convex polygon, given as an array \( p[0] \ldots p[n-1] \) of its \( n \) vertices in counterclockwise order.

a) Describe an \( O(\log(n)) \) time algorithm to determine whether a point \( q \) lies inside, outside or on the boundary of \( P \).

b) Describe an \( O(\log(n)) \) time algorithm to find a (right) tangent to \( P \) from a query point \( q \) located outside \( P \). That is, find a vertex \( p[i] \), such that \( P \) is contained in the closed halfplane to the left of the oriented line \( qp[i] \).
3.5 Jarvis' Wrap

We are now ready to describe a first simple algorithm to construct the convex hull. It works as follows:

Find a point \( p_1 \) that is a vertex of \( \text{conv}(P) \) (e.g., the one with smallest \( x \)-coordinate). “Wrap” \( P \) starting from \( p_1 \), i.e., always find the next vertex of \( \text{conv}(P) \) as the one that is rightmost with respect to the direction given by the previous two vertices.

Besides comparing \( x \)-coordinates, the only geometric primitive needed is an orientation test: Denote by \( \text{rightturn}(p, q, r) \), for three points \( p, q, r \in \mathbb{R}^2 \), the predicate that is true if and only if \( r \) is (strictly) to the right of the oriented line \( pq \).

![Diagram of Jarvis' Wrap](image)

**Code for Jarvis' Wrap.**

\( p[0..N) \) contains a sequence of \( N \) points.

\( p_{\text{start}} \) point with smallest \( x \)-coordinate.

\( q_{\text{next}} \) some other point in \( p[0..N) \).

```cpp
int h = 0;
Point_2 q_now = p_start;
do {
    q[h] = q_now;
    h = h + 1;
    for (int i = 0; i < N; i = i + 1)
        if (rightturn_2(q_now, q_next, p[i]))
            q_next = p[i];
    q_now = q_next;
} while (q_now != p_start);
```

\( q[0,h) \) describes a convex polygon bounding the convex hull of \( p[0..N) \).
Analysis. For every output point the above algorithm spends \( n \) rightturn tests, which is \( \Rightarrow O(nh) \) in total.

**Theorem 3.23** [6] Jarvis’ Wrap computes the convex hull of \( n \) points in \( \mathbb{R}^2 \) using \( O(nh) \) rightturn tests, where \( h \) is the number of hull vertices.

In the worst case we have \( h = n \), that is, \( O(n^2) \) rightturn tests. Jarvis’ Wrap has a remarkable property that is called output sensitivity: the runtime depends not only on the size of the input but also on the size of the output. For a huge point set it constructs the convex hull in optimal linear time, if the convex hull consists of a constant number of vertices only. Unfortunately the worst case performance of Jarvis’ Wrap is suboptimal, as we will see soon.

**Degeneracies.** The algorithm may have to cope with various degeneracies.

- Several points have smallest \( x \)-coordinate \( \Rightarrow \) lexicographic order:
  \[
  (p_x, p_y) < (q_x, q_y) \iff p_x < q_x \lor p_x = q_x \land p_y < q_y .
  \]

- Three or more points collinear \( \Rightarrow \) choose the point that is farthest among those that are rightmost.

**Predicates.** Besides the lexicographic comparison mentioned above, the Jarvis’ Wrap (and most other 2D convex hull algorithms for that matter) need one more geometric predicate: the rightturn or—more generally—orientation test. The computation amounts to evaluating a polynomial of degree two, see the exercise below. We therefore say that the orientation test has algebraic degree two. In contrast, the lexicographic comparison has degree one only. The algebraic degree not only has a direct impact on the efficiency of a geometric algorithm (lower degree \( \leftrightarrow \) less multiplications), but also an indirect one because high degree predicates may create large intermediate results, which may lead to overflows and are much more costly to compute with exactly.

**Exercise 3.24** Prove that for three points \( (p_x, p_y), (q_x, q_y), (r_x, r_y) \in \mathbb{R}^2 \), the sign of the determinant

\[
\begin{vmatrix}
1 & p_x & p_y \\
1 & q_x & q_y \\
1 & r_x & r_y
\end{vmatrix}
\]

determines if \( r \) lies to the right, to the left or on the directed line through \( p \) and \( q \).
3.6 Graham Scan (Successive Local Repair)

There exist many algorithms that exhibit a better worst-case runtime than Jarvis’ Wrap. Here we discuss only one of them: a particularly elegant and easy-to-implement variant of the so-called Graham Scan [5]. This algorithm is referred to as Successive Local Repair because it starts with some polygon enclosing all points and then step-by-step repairs the deficiencies of this polygon, by removing non-convex vertices. It goes as follows:

Sort points lexicographically and remove duplicates: \((p_1, \ldots, p_n)\).

As long as there is a (consecutive) triple \((p, q, r)\) such that \(r\) is to the right of or on the directed line \(\overrightarrow{pq}\), remove \(q\) from the sequence.

**Code for Graham Scan.**

\(p[0..N)\) lexicographically sorted sequence of pairwise distinct points, \(N \geq 2\).

\[
\begin{align*}
q[0] &= p[0] ; \\
\text{int } h &= 0 ; \\
&// \text{ Lower convex hull (left to right):} \\
&\text{for } (\text{int } i = 1 ; i < N ; i = i + 1 ) \{ \\
&\quad \text{while } (h > 0 \&\& !\text{leftturn}_2(q[h-1], q[h], p[i])) \\
&\quad \quad h = h - 1 ; \\
&\quad \quad h = h + 1 ; \\
&\quad q[h] = p[i] ; \\
&\} \\
&// \text{ Upper convex hull (right to left):} \\
&\text{for } (\text{int } i = N - 2 ; i >= 0 ; i = i - 1 ) \{ \\
&\quad \text{while } (!\text{leftturn}_2(q[h-1], q[h], p[i])) \\
&\quad \quad h = h - 1 ; \\
&\quad \quad h = h + 1 ; \\
&\quad q[h] = p[i] ; \\
&\}
\end{align*}
\]

\(q[0,h)\) describes a convex polygon bounding the convex hull of \(p[0..N)\).
Analysis.

**Theorem 3.25** The convex hull of a set $P \subset \mathbb{R}^2$ of $n$ points can be computed using $O(n \log n)$ geometric operations.

**Proof.**

1. Sorting and removal of duplicate points: $O(n \log n)$.

2. At the beginning we have a sequence of $2n - 1$ points; at the end the sequence consists of $h$ points. Observe that for every positive orientation test, one point is discarded from the sequence for good. Therefore, we have exactly $2n - h - 1$ such shortcuts/positive orientation tests. In addition there are at most $2n - 2$ negative tests (#iterations of the outer for loops). Altogether we have at most $4n - h - 3$ orientation tests.

In total the algorithm uses $O(n \log n)$ geometric operations. Note that the number of orientation tests is linear only, but $O(n \log n)$ lexicographic comparisons are needed. □

### 3.7 Lower Bound

It is not hard to see that the runtime of Graham Scan is asymptotically optimal in the worst-case.

**Theorem 3.26** $\Omega(n \log n)$ geometric operations are needed to construct the convex hull of $n$ points in $\mathbb{R}^2$ (in the algebraic computation tree model).

**Proof.** Reduction from sorting (for which it is known that $\Omega(n \log n)$ comparisons are needed in the algebraic computation tree model). Given $n$ real numbers $x_1, \ldots, x_n$, construct a set $P = \{p_i \mid 1 \leq i \leq n\}$ of $n$ points in $\mathbb{R}^2$ by setting $p_i = (x_i, x_i^2)$. This construction can be regarded as embedding the numbers into $\mathbb{R}^2$ along the $x$-axis and then projecting the resulting points vertically onto the unit parabola. The order in which the points appear along the lower convex hull of $P$ corresponds to the sorted order of the $x_i$. Therefore, if we could construct the convex hull in $o(n \log n)$ time, we could also sort in $o(n \log n)$ time. □

Clearly this reduction does not work for the Extremal Points problem. But using a reduction from Element Uniqueness (see Section 1.1) instead, one can show that $\Omega(n \log n)$ is also a lower bound for the number of operations needed to compute the set of extremal points only. This was first shown by Avis [1] for linear computation trees, then by Yao [9] for quadratic computation trees, and finally by Ben-Or [2] for general algebraic computation trees.
3.8 Chan’s Algorithm

Given matching upper and lower bounds we may be tempted to consider the algorithmic complexity of the planar convex hull problem settled. However, this is not really the case: Recall that the lower bound is a worst case bound. For instance, the Jarvis’ Wrap runs in $O(nh)$ time an thus beats the $\Omega(n \log n)$ bound in case that $h = o(\log n)$. The question remains whether one can achieve both output dependence and optimal worst case performance at the same time. Indeed, Chan [4] presented an algorithm to achieve this runtime by cleverly combining the “best of” Jarvis’ Wrap and Graham Scan. Let us look at this algorithm in detail. The algorithm consists of two steps that are executed one after another.

**Divide.** Input: a set $P \subset \mathbb{R}^2$ of $n$ points and a number $H \in \{1, \ldots, n\}$.

1. Divide $P$ into $k = \lceil n/H \rceil$ sets $P_1, \ldots, P_k$ with $|P_i| \leq H$.
2. Construct $\text{conv}(P_i)$ for all $i$, $1 \leq i \leq k$.

**Analysis.** Step 1 takes $O(n)$ time. Step 2 can be handled using Graham Scan in $O(H \log H)$ time for any single $P_i$, that is, $O(n \log H)$ time in total.

**Conquer.** Output: the vertices of $\text{conv}(P)$ in counterclockwise order, if $\text{conv}(P)$ has less than $H$ vertices; otherwise, the message that $\text{conv}(P)$ has at least $H$ vertices.

1. Find the lexicographically smallest point in $\text{conv}(P_i)$ for all $i$, $1 \leq i \leq k$.
2. Starting from the lexicographically smallest point of $P$ find the first $H$ points of $\text{conv}(P)$ oriented counterclockwise (simultaneous Jarvis’ Wrap on the sequences $\text{conv}(P_i)$).

Determine in every wrap step the point $q_i$ of tangency from the current point of $\text{conv}(P)$ to $\text{conv}(P_i)$, for all $1 \leq i \leq k$. We have seen in Exercise 3.22 how to compute $q_i$ in $O(\log \text{conv}(P_i)) = O(\log H)$ time. Among the $k$ candidates $q_1, \ldots, q_k$ we find the next vertex of $\text{conv}(P)$ in $O(k)$ time.

**Analysis.** Step 1 takes $O(n)$ time. Step 2 consists of at most $H$ wrap steps. Each wrap step needs $O(k \log H + k) = O(k \log H)$ time, which amounts to $O(Hk \log H) = O(n \log H)$ time for Step 2 in total.

**Remark.** Using a more clever search strategy instead of many tangency searches one can handle the conquer phase in $O(n)$ time, see Exercise 3.27 below. However, this is irrelevant as far as the asymptotic runtime is concerned, given that already the divide step takes $O(n \log H)$ time.
Exercise 3.27 Consider \( k \) convex polygons \( P_1, \ldots, P_k \), for some constant \( k \in \mathbb{N} \), where each polygon is given as a list of its vertices in counterclockwise orientation. Show how to construct the convex hull of \( P_1 \cup \ldots \cup P_k \) in \( O(n) \) time, where \( n = \sum_{i=1}^{k} n_i \) and \( n_i \) is the number of vertices of \( P_i \), for \( 1 \leq i \leq k \).

Searching for \( h \). While the runtime bound for \( H = h \) is exactly what we were heading for, it looks like in order to actually run the algorithm we would have to know \( h \), which—in general—we do not. Fortunately we can circumvent this problem rather easily, by applying what is called a doubly exponential search. It works as follows.

Call the algorithm from above iteratively with parameter \( H = \min\{2^{2^t}, n\} \), for \( t = 0, \ldots \), until the conquer step finds all extremal points of \( P \) (i.e., the wrap returns to its starting point).

**Analysis:** Let \( 2^{2^t} \) be the last parameter for which the algorithm is called. Since the previous call with \( H = 2^{2^{t-1}} \) did not find all extremal points, we know that \( 2^{2^{t-1}} < h \), that is, \( 2^{t-1} < \log h \), where \( h \) is the number of extremal points of \( P \). The total runtime is therefore at most

\[
\sum_{i=0}^{s} cn \log 2^i = cn \sum_{i=0}^{s} 2^i = cn(2^{s+1} - 1) < 4cn \log h = O(n \log h),
\]

for some constant \( c \in \mathbb{R} \). In summary, we obtain the following theorem.

**Theorem 3.28** The convex hull of a set \( P \subset \mathbb{R}^2 \) of \( n \) points can be computed using \( O(n \log h) \) geometric operations, where \( h \) is the number of convex hull vertices.

**Questions**

6. How is convexity defined? What is the convex hull of a set in \( \mathbb{R}^d \)? Give at least three possible definitions.

7. What does it mean to compute the convex hull of a set of points in \( \mathbb{R}^2 \)? Discuss input and expected output and possible degeneracies.

8. How can the convex hull of a set of \( n \) points in \( \mathbb{R}^2 \) be computed efficiently? Describe and analyze (incl. proofs) Jarvis’ Wrap, Successive Local Repair, and Chan’s Algorithm.

9. Is there a linear time algorithm to compute the convex hull of \( n \) points in \( \mathbb{R}^2 \)? Prove the lower bound and define/explain the model in which it holds.

10. Which geometric primitive operations are used to compute the convex hull of \( n \) points in \( \mathbb{R}^2 \)? Explain the two predicates and how to compute them.
Remarks. The sections on Jarvis’ Wrap and Graham Scan are based on material that Emo Welzl prepared for a course on “Geometric Computing” in 2000.

References


Chapter 4

Plane Graphs and the DCEL

So far we have been talking about geometric structures such as triangulations of polygons without paying much attention to how to represent them. This will change now. As many of these structures can be regarded as plane graphs, we start by reviewing a bit of terminology from (planar) graph theory. We assume that this is—at least somewhat—familiar ground for you. If you would like to see more details and examples, please refer to any standard textbook on graph theory, such as Bondy and Murty [2], Diestel [3], or West [10].

A (simple undirected) graph $G = (V, E)$ where $V$ is a finite set of vertices and $E$ is the set of edges, $E \subseteq (V^2) := \{(u, v) \mid u, v \in V, u \neq v\}$. For brevity, one often writes $uv$ rather than $\{u, v\}$ to denote an edge. The two vertices defining an edge are adjacent to each other and they are incident to the edge.

For a vertex $v \in V$, denote by $N_G(v)$ the neighborhood of $v$ in $G$, that is, the set of vertices from $G$ that are adjacent to $v$. Similarly, for a set $W \subset V$ of vertices define $N_G(W) := \bigcup_{w \in W} N_G(w)$. The degree $\deg_G(v)$ of a vertex $v \in V$ is the size of its neighborhood, that is, the number of edges from $E$ incident to $v$. The subscript is often omitted when it is clear to which graph it refers to. As every edge is incident to exactly two vertices, we have the following simple but useful relationship, often called handshaking-lemma: $\sum_{v \in V} \deg(v) = 2|E|$. A walk in a graph $G$ is a sequence $W = (v_1, \ldots, v_k), k \in \mathbb{N}$, of vertices such that $v_i$ and $v_{i+1}$ are adjacent in $G$, for all $1 \leq i < k$. A path is a walk whose vertices are pairwise distinct. A cycle is a walk whose vertices are pairwise distinct, except for $v_1 = v_k$. A graph is connected, if there is a path between any pair of vertices. If a graph is not connected, it is disconnected. A disconnected graph can be decomposed into maximal connected subgraphs, its (connected) components.

A tree is a connected graph that does not have any cycle. It is not hard to show that trees on $n$ vertices are exactly the graphs on $n$ vertices that have $n - 1$ edges.

In geometry we are typically concerned with graphs that are embedded on some surface, here $\mathbb{R}^2$. An embedding or drawing is just a “nice” mapping of a graph $G$ into the plane. More formally, each vertex $v$ is mapped to a distinct point $\phi(v) \in \mathbb{R}^2$ (that is, $\phi : V \to \mathbb{R}^2$ is injective) and each edge $uv$ is mapped to an arc—a simple Jordan
curve—\( \phi(uv) \subset \mathbb{R}^2 \) from \( \phi(u) \) to \( \phi(v) \) such that no vertex is mapped to the relative interior of \( \phi(uv) \).

A graph is \textit{planar} if it admits a \textit{crossing-free} drawing, that is, a drawing in which no two arcs intersect except at common endpoints. For example, \( K_4 \) (the complete graph on four vertices) is planar, as demonstrated by the drawing shown in Figure 4.1a.

![Crossing-free drawing.](image1)

![Straight-line drawing.](image2)

\textbf{Figure 4.1: Embeddings of }K_4\textit{ into the plane.}

If a graph is planar, then by Fáry-Wagner’s Theorem \cite{4,8} there also exists a \textit{straight-line} drawing, that is, a drawing in which all arcs are line segments. In order to obtain such a drawing for \( K_4 \), we have to put one vertex inside the convex hull of the other three, see Figure 4.1b.

Sometimes we refer to graphs not as abstract graphs but in a concrete embedding. If this embedding is a crossing-free embedding into the plane, the embedded graph is referred to as a \textit{plane graph}. Note the distinction between “planar” and “plane”: The former refers to an abstract graph and expresses the possibility of a crossing-free drawing, whereas the latter refers to a geometric object that is a concrete crossing-free drawing of a graph in the plane.

A \textit{plane straight-line graph} (PSLG) is a crossing-free straight-line drawing of a graph in the plane. Both graphs in Figure 4.1 are plane, but only the one shown on the right is also a plane straight-line graph.

\section*{4.1 The Euler Formula}

If we remove all vertices (points) and edges (arcs) of a plane graph \( G \) from \( \mathbb{R}^2 \), then what remains is a finite collection of open sets. These sets are referred to as the \textit{faces} of \( G \). For instance, both plane graphs in Figure 4.1 have 4 vertices, 6 edges and 4 faces (one of which is unbounded). In general, if \(|V|\) is the number of vertices of a connected plane graph, \(|E|\) its number of edges and \(|F|\) the number of faces, then the Euler Formula states that

\[ |V| - |E| + |F| = 2. \]

In the example, we get \( 4 - 6 + 4 = 2 \).
4.2 The Doubly-Connected Edge List

If you do not insist on being too formal, the proof is simple and works by induction on the number of edges. If we fix the number of vertices, the base case occurs for \(|V| - 1\) edges where the plane graph is a tree. Then we have \(|F| = 1\) and the formula holds. A graph with more edges always contains a cycle and therefore at least one bounded face. Choose one edge from a bounded face and remove it. The resulting graph is still connected and has one edge less but also one face less since the edge removal merges the bounded face with another one. Consequently, since the Euler Formula holds for the smaller graph by induction, it also holds for the larger graph.

The Euler Formula can be used to prove the following important fact about planar graphs.

**Lemma 4.1** Any planar graph on \(n \geq 3\) vertices has at most \(3n - 6\) edges and at most \(2n - 4\) faces.

This lemma shows that the overall complexity—the sum of the number of vertices, edges, and faces—of a planar graph is linear in \(n\). A planar graph with a maximal number \((3n - 6)\) of edges is called **maximal planar**.

**Exercise 4.2** Prove Lemma 4.1 using the Euler formula.

**Exercise 4.3** Prove that every planar graph has a vertex of degree at most 5.

**Exercise 4.4** Show that \(K_5\) (the complete graph on five vertices) is not planar.

### 4.2 The Doubly-Connected Edge List

Many algorithms in computational geometry work with plane graphs. The **doubly-connected edge list** (DCEL) is a data structure to represent a plane graph in such a way that it is easy to traverse and to manipulate. In order to avoid unnecessary complications, let us discuss only connected graphs here that contain at least two vertices. It is not hard to extend the data structure to cover all plane graphs. For simplicity we also assume that we deal with a straight-line embedding and so the geometry of edges is defined by the mapping of their endpoints already. For more general embeddings, the geometric description of edges has to be stored in addition.

The main building block of a DCEL is a list of **halfedges**. Every actual edge is represented by two halfedges going in opposite direction, and these are called **twins**, see Figure 4.2. Along the boundary of each face, halfedges are oriented counterclockwise.

A DCEL stores a list of halfedges, a list of vertices, and a list of faces. These lists are unordered but interconnected by various pointers. A vertex \(v\) stores a pointer \(\text{halfedge}(v)\) to an arbitrary halfedge originating from \(v\). Every vertex also knows its coordinates, that is, the point \(\text{point}(v)\) it is mapped to in the represented embedding. A face \(f\) stores a pointer \(\text{halfedge}(f)\) to an arbitrary halfedge within the face. A halfedge \(h\) stores **five** pointers:
A constant amount of information is stored for every vertex, (half-)edge, and face of the graph. Therefore the whole DCEL needs storage proportional to $|V| + |E| + |F|$, which is $O(n)$ for a plane graph with $n$ vertices by Lemma 4.1. This information is sufficient for most tasks. For example, traversing all edges around a face $f$ can be done as follows:

$$s \leftarrow \text{halfedge}(f)$$
$$h \leftarrow s$$
$$\text{do}$$
$$\quad \text{something with } h$$
$$\quad h \leftarrow \text{next}(h)$$
$$\text{while } h \neq s$$

**Exercise 4.5** Give pseudocode to traverse all edges incident to a given vertex $v$ of a DCEL.

**Exercise 4.6** Why is the previous halfedge $\text{prev}(\cdot)$ stored explicitly and the source vertex of a halfedge is not?

### 4.2.1 Manipulating a DCEL

In many geometric algorithms, plane graphs appear not just as static objects but rather they evolve over the course of the algorithm. Therefore the data structure used to represent the graph must allow for efficient update operations to change it.
4.2. The Doubly-Connected Edge List

First of all, we need to be able to generate new vertices, edges, and faces, to be added to the corresponding list within the DCEL and—symmetrically—the ability to delete an existing entity. Then it should be easy to add a new vertex \( v \) to the graph within some face \( f \). As we maintain a connected graph, we better link the new vertex to somewhere, say, to an existing vertex \( u \). For such a connection to be possible, we require that the open line segment \( uv \) lies completely in \( f \).

Of course, two halfedges are to be added connecting \( u \) and \( v \). But where exactly? Given that from a vertex and from a face only some arbitrary halfedge is directly accessible, it turns out convenient to use a halfedge in the interface. Let \( h \) denote the halfedge incident to \( f \) for which \( \text{target}(h) = u \). Our operation becomes then (see also Figure 4.3)

\[
\text{add-vertex-at}(v, h)
\]

Precondition: the open line segment \( \text{point}(v)\text{point}(u) \), where \( u := \text{target}(h) \), lies completely in \( f := \text{face}(h) \).

Postcondition: a new vertex \( v \) has been inserted into \( f \), connected by an edge to \( u \).

![Figure 4.3: Add a new vertex connected to an existing vertex \( u \).](image)

and it can be realized by manipulating a constant number of pointers as follows.

\[
\text{add-vertex-at}(v, h) \{
    h_1 \leftarrow \text{a new halfedge}
    h_2 \leftarrow \text{a new halfedge}
    \text{halfedge}(v) \leftarrow h_2
    \text{twin}(h_1) \leftarrow h_2
    \text{twin}(h_2) \leftarrow h_1
    \text{target}(h_1) \leftarrow v
    \text{target}(h_2) \leftarrow u
    \text{face}(h_1) \leftarrow f
    \text{face}(h_2) \leftarrow f
    \text{next}(h_1) \leftarrow h_2
    \text{next}(h_2) \leftarrow \text{next}(h)
    \text{prev}(h_1) \leftarrow h
\}
\]
Similarly, it should be possible to add an edge between two existing vertices \( u \) and \( v \), provided the open line segment \( uv \) lies completely within a face \( f \) of the graph, see Figure 4.4. Since such an edge insertion splits \( f \) into two faces, the operation is called \textit{split-face}. Again we use the halfedge \( h \) that is incident to \( f \) and for which \( \text{target}(h) = u \).

Our operation becomes then

\[
\text{split-face}(h, v)
\]

Precondition: \( v \) is incident to \( f := \text{face}(h) \) but not adjacent to \( u := \text{target}(h) \).

The open line segment \( \text{point}(v)\text{point}(u) \) lies completely in \( f \).
Postcondition: \( f \) has been split by a new edge \( uv \).

The implementation is slightly more complicated compared to \textit{add-vertex-at} above, because the face \( f \) is destroyed and so we have to update the face information of all incident halfedges. In particular, this is not a constant time operation, but its time complexity is proportional to the size of \( f \).

\[
\text{split-face}(h, v) \{
\begin{align*}
f_1 &\leftarrow \text{a new face} \\
f_2 &\leftarrow \text{a new face} \\
h_1 &\leftarrow \text{a new halfedge} \\
h_2 &\leftarrow \text{a new halfedge} \\
\text{halfedge}(f_1) &\leftarrow h_1 \\
\text{halfedge}(f_2) &\leftarrow h_2 \\
\text{twin}(h_1) &\leftarrow h_2 \\
\text{twin}(h_2) &\leftarrow h_1 \\
\text{target}(h_1) &\leftarrow v
\end{align*}
\}
\]
4.2. The Doubly-Connected Edge List

\[
\begin{align*}
\text{target}(h_2) & \leftarrow u \\
\text{next}(h_2) & \leftarrow \text{next}(h) \\
\text{prev}(\text{next}(h_2)) & \leftarrow h_2 \\
\text{prev}(h_1) & \leftarrow h \\
\text{next}(h_1) & \leftarrow h_1 \\
i & \leftarrow h_2 \\
\text{loop} & \\
\text{face}(i) & \leftarrow f_2 \\
\text{if} \quad \text{target}(i) = v & \text{break the loop} \\
i & \leftarrow \text{next}(i) \\
\text{endloop} & \\
\text{next}(h_1) & \leftarrow \text{next}(i) \\
\text{prev}(\text{next}(h_1)) & \leftarrow h_1 \\
\text{next}(i) & \leftarrow h_2 \\
\text{prev}(h_2) & \leftarrow i \\
i & \leftarrow h_1 \\
\text{do} & \\
\text{face}(i) & \leftarrow f_1 \\
i & \leftarrow \text{next}(i) \\
\text{until} \quad \text{target}(i) = u & \\
\text{delete the face } f & \\
\end{align*}
\]

In a similar fashion one can realize the inverse operation \text{join-face}(h) that removes the edge (represented by the halfedge) \( h \), thereby joining the faces \text{face}(h) and \text{face}(\text{twin}(h)).

It is easy to see that every connected plane graph on at least two vertices can be constructed using the operations \text{add-vertex-at} and \text{split-face}, starting from an embedding of \( K_2 \) (two vertices connected by an edge).

**Exercise 4.7** Give pseudocode for the operation \text{join-face}(h). Also specify preconditions, if needed.

**Exercise 4.8** Give pseudocode for the operation \text{split-edge}(h), that splits the edge (represented by the halfedge) \( h \) into two by a new vertex \( w \), see Figure 4.5.

### 4.2.2 Graphs with Unbounded Edges

In some cases it is convenient to consider plane graphs, in which some edges are not mapped to a line segment but to an unbounded curve, such as a ray. This setting is not really much different from the one we studied before, except that one vertex is placed “at infinity”. One way to think of it is in terms of stereographic projection: Imagine \( \mathbb{R}^2 \) being the \( x/y \)-plane in \( \mathbb{R}^3 \) and place a unit sphere \( S \) such that its southpole touches the origin. We obtain a bijective continuous mapping between \( \mathbb{R}^2 \) and \( S \setminus \{n\} \), where \( n \)
Figure 4.5: Split an edge by a new vertex.

is the northpole of $S$, as follows: A point $p \in \mathbb{R}^2$ is mapped to the point $p'$ that is the intersection of the line through $p$ and $n$ with $S$, see Figure 4.6. The further away a point

Figure 4.6: Stereographic projection.

in $\mathbb{R}^2$ is from the origin, the closer its image on $S$ gets to $n$. But there is no way to reach $n$ except in the limit. Therefore, we can imagine drawing the graph on $S$ instead of in $\mathbb{R}^2$ and putting the “infinite vertex” at $n$.

All this is just for the sake of a proper geometric interpretation. As far as a DCEL representation of such a graph is concerned, there is no need to consider spheres or, in fact, anything beyond what we have discussed before. The only difference to the case with all finite edges is that there is this special infinite vertex, which does not have any point/coordinates associated to it. But other than that, the infinite vertex is treated in exactly the same way as the finite vertices: it has in– and outgoing halfedges along which the unbounded faces can be traversed (Figure 4.7).

4.2.3 Remarks

It is actually not so easy to point exactly to where the DCEL data structure originates from. Often Muller and Preparata [7] are credited, but while they use the term DCEL,
the data structure they describe is different from what we discussed above and from what people usually consider a DCEL nowadays. Overall, there is a large number of variants of this data structure, which appear under the names winged edge data structure [1], halfedge data structure [9], or quad-edge data structure [5]. Kettner [6] provides a comparison of all these and some additional references.

Questions

11. What are planar/plane graphs and straight-line embeddings? Give the definitions and explain the difference between planar and plane.

12. How many edges can a planar graph have? What is the average vertex degree in a planar graph? Explain Euler’s formula and derive your answers from it (see Exercise 4.2 and 4.3).

13. How can plane graphs be represented on a computer? Explain the DCEL data structure and how to work with it.

References

Chapter 4. Plane Graphs and the DCEL


Chapter 5

Line Sweep

In this chapter we will discuss a simple and widely applicable paradigm to design geometric algorithms: the so-called Line-Sweep (or Plane-Sweep) technique. It can be used to solve a variety of different problems, some examples are listed below. The first part may come as a reminder to many of you, because you should have heard something about line-sweep in one of the basic CS courses already. However, we will soon proceed and encounter a couple of additional twists that were most likely not covered there.

Consider the following geometric problems.

**Problem 5.1 (Simple Polygon Test)** Given a sequence \( P = (p_1, \ldots, p_n) \) of points in \( \mathbb{R}^2 \), does \( P \) describe the boundary of a simple polygon?

**Problem 5.2 (Polygon Intersection)** Given two simple polygons \( P \) and \( Q \) in \( \mathbb{R}^2 \) as a (counterclockwise) sequence of their vertices, is \( P \cap Q = \emptyset \)?

**Problem 5.3 (Segment Intersection Test)** Given a set \( S \) of \( n \) closed line segments in \( \mathbb{R}^2 \), do any two of them intersect?

**Remark:** In principle it is clear what is meant by “two segments intersect”. But there are a few special cases that one may have to consider carefully. For instance, does it count if an endpoint lies on another segment? What if two segments share an endpoint? What about overlapping segments and segments of length zero? In general, let us count all these as intersections. However, sometimes we may want to exclude some of these cases. For instance, in a simple polygon test, we do not want to consider the shared endpoint between two consecutive edges of the boundary as an intersection.

**Problem 5.4 (Segment Intersections)** Given a set \( S \) of \( n \) closed line segments in \( \mathbb{R}^2 \), compute all pairs of segments that intersect.
Problem 5.5 (Segment Arrangement) Given a set \( S \) of \( n \) closed line segments in \( \mathbb{R}^2 \), construct the arrangement induced by \( S \), that is, the subdivision of \( \mathbb{R}^2 \) induced by \( S \).

Problem 5.6 (Map Overlay) Given two sets \( S \) and \( T \) of \( n \) and \( m \), respectively, pairwise interior disjoint line segments in \( \mathbb{R}^2 \), construct the arrangement induced by \( S \cup T \).

In the following we will use Problem 5.4 as our flagship example.

**Trivial Algorithm.** Test all the \( \binom{n}{2} \) pairs of segments from \( S \) in \( O(n^2) \) time and \( O(n) \) space. For Problem 5.4 this is worst-case optimal because there may be \( \Omega(n^2) \) intersecting pairs.

But in case that the number of intersecting pairs is, say, linear in \( n \) there is still hope to obtain a subquadratic algorithm. Given that there is a lower bound of \( \Omega(n \log n) \) for **Element Uniqueness** (Given \( x_1, \ldots, x_n \in \mathbb{R} \), is there an \( i \neq j \) such that \( x_i = x_j \)?) in the algebraic computation tree model, all we can hope for is an output-sensitive runtime of the form \( O(n \log n + k) \), where \( k \) denotes the number of intersecting pairs (output size).

### 5.1 Interval Intersections

As a warmup let us consider the corresponding problem in \( \mathbb{R}^1 \).

**Problem 5.7** Given a set \( I \) of \( n \) intervals \( [\ell_i, r_i] \subset \mathbb{R}, 1 \leq i \leq n \). Compute all pairs of intervals from \( I \) that intersect.

**Theorem 5.8** Problem 5.7 can be solved in \( O(n \log n + k) \) time and \( O(n) \) space, where \( k \) is the number of intersecting pairs from \( \binom{n}{2} \).

**Proof.** First observe that two real intervals intersect if and only if one contains the right endpoint of the other.

Sort the set \( \{(\ell_i, 0) \mid 1 \leq i \leq n\} \cup \{(r_i, 1) \mid 1 \leq i \leq n\} \) in increasing lexicographic order and denote the resulting sequence by \( P \). Store along with each point from \( P \) its origin \( i \). Walk through \( P \) from start to end while maintaining a list \( L \) of intervals that contain the current point \( p \in P \).

Whenever \( p = (\ell_i, 0), 1 \leq i \leq n \), insert \( i \) into \( L \). Whenever \( p = (r_i, 1), 1 \leq i \leq n \), remove \( i \) from \( L \) and then report for all \( j \in L \) the pair \( \{i, j\} \) as intersecting. \( \square \)

### 5.2 Segment Intersections

How can we transfer the (optimal) algorithm for the corresponding problem in \( \mathbb{R}^1 \) to the plane? In \( \mathbb{R}^1 \) we moved a point from left to right and at any point resolved the situation locally around this point. More precisely, at any point during the algorithm, we knew all
5.2. Segment Intersections

intersections that are to the left of the current (moving) point. A point can be regarded a hyperplane in $\mathbb{R}^1$, and the corresponding object in $\mathbb{R}^2$ is a line.

**General idea.** Move a line $\ell$ (so-called *sweep line*) from left to right over the plane, such that at any point during this process all intersections to the left of $\ell$ have been reported.

**Sweep line status.** The list of intervals containing the current point corresponds to a list $L$ of segments (sorted by $y$-coordinate) that intersect the current sweep line $\ell$. This list $L$ is called *sweep line status* (SLS). Considering the situation locally around $L$, it is obvious that only segments that are adjacent in $L$ can intersect each other. This observation allows to reduce the overall number of intersection tests, as we will see. In order to allow for efficient insertion and removal of segments, the SLS is usually implemented as a balanced binary search tree.

**Event points.** The order of segments in SLS can change at certain points only: whenever the sweep line moves over a segment endpoint or a point of intersection of two segments from $S$. Such a point is referred to as an *event point* (EP) of the sweep. Therefore we can reduce the conceptually continuous process of moving the sweep line over the plane to a discrete process that moves the line from EP to EP. This discretization allows for an efficient computation.

At any EP several events can happen simultaneously: several segments can start and/or end and at the same point a couple of other segments can intersect. In fact the sweep line does not even make a difference between any two event points that have the same $x$-coordinate. To properly resolve the order of processing, EPs are considered in lexicographic order and wherever several events happen at a single point, these are considered simultaneously as a single EP. In this light, the sweep line is actually not a line but an infinitesimal step function (see Figure 5.1).

**Event point schedule.** In contrast to the one-dimensional situation, in the plane not all EP are known in advance because the points of intersection are discovered during the algorithm only. In order to be able to determine the next EP at any time, we use a priority queue data structure, the so-called *event point schedule* (EPS).

Along with every EP $p$ store a list $\text{end}(p)$ of all segments that end at $p$, a list $\text{begin}(p)$ of all segments that begin at $p$, and a list $\text{int}(p)$ of all segments in SLS that intersect at $p$ a segment that is adjacent to it in SLS.

Along with every segment we store pointers to all its appearances in an $\text{int}(-)$ list of some EP. As a segment appears in such a list only if it intersects one of its neighbors there, every segment needs to store at most two such pointers.
Invariants. The following conditions can be shown to hold before and after any event point is handled. (We will not formally prove this here.) In particular, the last condition at the end of the sweep implies the correctness of the algorithm.

1. \( L \) is the sequence of segments from \( S \) that intersect \( \ell \), ordered by \( y \)-coordinate of their point of intersection.

2. \( E \) contains all endpoints of segments from \( S \) and all points where two segments that are adjacent in \( L \) intersect to the right of \( \ell \).

3. All pairs from \( (S^2) \) that intersect to the left of \( \ell \) have been reported.

Event point handling. An EP \( p \) is processed as follows.

1. If \( \text{end}(p) \cup \text{int}(p) = \emptyset \), localize \( p \) in \( L \).

2. Report all pairs of segments from \( \text{end}(p) \cup \text{begin}(p) \cup \text{int}(p) \) as intersecting.

3. Remove all segments in \( \text{end}(p) \) from \( L \).

4. Reverse the subsequence in \( L \) that is formed by the segments from \( \text{int}(p) \).

5. Insert segments from \( \text{begin}(p) \) into \( L \), sorted by slope.

6. Test the topmost and bottommost segment in \( L \) from \( \text{begin}(p) \cup \text{int}(p) \) for intersection with its successor and predecessor, respectively, and update EP if necessary.
Updating EPS. Insert an EP \( p \) corresponding to an intersection of two segments \( s \) and \( t \). Without loss of generality let \( s \) be above \( t \) at the current position of \( \ell \).

1. If \( p \) does not yet appear in \( E \), insert it.

2. If \( s \) is contained in an \( \text{int}(\cdot) \) list of some other EP \( q \), where it intersects a segment \( t' \) from above: Remove both \( s \) and \( t' \) from the \( \text{int}(\cdot) \) list of \( q \) and possibly remove \( q \) from \( E \) (if \( \text{end}(q) \cup \text{begin}(q) \cup \text{int}(q) = \emptyset \)). Proceed analogously in case that \( t \) is contained in an \( \text{int}(\cdot) \) list of some other EP, where it intersects a segment from below.

3. Insert \( s \) and \( t \) into \( \text{int}(p) \).

Sweep.

1. Insert all segment endpoints into \( \text{begin}(\cdot) \)/\( \text{end}(\cdot) \) list of a corresponding EP in \( E \).

2. As long as \( E \neq \emptyset \), handle the first EP and then remove it from \( E \).

Runtime analysis. Initialization: \( O(n \log n) \). Processing of an EP \( p \):

\[
O(\#\text{intersecting pairs} + |\text{end}(p)| \log n + |\text{int}(p)| + |\text{begin}(p)| \log n + \log n).
\]

In total: \( O(k + n \log n + k \log n) = O((n + k) \log n) \), where \( k \) is the number of intersecting pairs in \( S \).

Space analysis. Clearly \( |S| \leq n \). At begin we have \( |E| \leq 2n \) and \( |S| = 0 \). Furthermore the number of additional EPs corresponding to points of intersection is always bounded by \( 2|S| \). Thus the space needed is \( O(n) \).

**Theorem 5.9** Problem 5.4 and Problem 5.5 can be solved in \( O((n + k) \log n) \) time and \( O(n) \) space.

**Theorem 5.10** Problem 5.1, Problem 5.2 and Problem 5.3 can be solved in \( O(n \log n) \) time and \( O(n) \) space.

**Exercise 5.11** Flesh out the details of the sweep line algorithm for Problem 5.2 that is referred to in Theorem 5.10. What if you have to construct the intersection rather than just to decide whether or not it is empty?

**Exercise 5.12** You are given \( n \) axis-parallel rectangles in \( \mathbb{R}^2 \) with their bottom sides lying on the x-axis. Construct their union in \( O(n \log n) \) time.
Chapter 5. Line Sweep

5.3 Improvements

The basic ingredients of the line sweep algorithm go back to work by Bentley and Ottmann [2] from 1979. The particular formulation discussed here, which takes all possible degeneracies into account, is due to Mehlhorn and Näher [9].

Theorem 5.10 is obviously optimal for Problem 5.3, because this is just the 2-dimensional generalization of Element Uniqueness (see Section 1.1). One might suspect there is also a similar lower bound of $\Omega(n \log n)$ for Problem 5.1 and Problem 5.2. However, this is not the case: Both can be solved in $O(n)$ time, albeit using a very complicated algorithm of Chazelle [4] (the famous triangulation algorithm).

Similarly, it is not clear why $O(n \log n + k)$ time should not be possible in Theorem 5.9. Indeed this was a prominent open problem in the 1980’s that has been solved in several steps by Chazelle and Edelsbrunner in 1988. The journal version of their paper [5] consists of 54 pages and the space usage is suboptimal $O(n + k)$.

Clarkson and Shor [6] and independently Mulmuley [10, 11] described randomized algorithms with expected runtime $O(n \log n + k)$ using $O(n)$ and $O(n + k)$, respectively, space.

An optimal deterministic algorithm, with runtime $O(n \log n + k)$ and using $O(n)$ space, is known since 1995 only due to Balaban [1].

5.4 Algebraic degree of geometric primitives

We already have encountered different notions of complexity during this course: We can analyze the time complexity and the space complexity of algorithms and both usually depend on the size of the input. In addition we have seen examples of output-sensitive algorithms, whose complexity also depends on the size of the output. In this section, we will discuss a different kind of complexity that is the algebraic complexity of the underlying geometric primitives. In this way, we try to shed a bit of light into the bottom layer of geometric algorithms that is often swept under the rug because it consists of constant time operations only. Nevertheless, it would be a mistake to disregard this layer completely, because in most—if not every—real world application it plays a crucial role, by affecting efficiency as well as correctness. Regarding efficiency, the value of the constants involved can make a big difference. The possible effects on correctness will hopefully become clear in the course of this section.

In all geometric algorithms there are some fundamental geometric predicates and/or constructions on the bottom level. Both are operations on geometric objects, the difference is only in the result: The result of a construction is a geometric object (for instance, the point common to two non-parallel lines), whereas the result of a predicate is Boolean ($\text{true}$ or $\text{false}$).

Geometric predicates and constructions in turn are based on fundamental arithmetic operations. For instance, we formulated planar convex hull algorithms in terms of an orientation predicate—given three points $p, q, r \in \mathbb{R}^2$, is $r$ strictly to the right of the ori-
5.4. Algebraic degree of geometric primitives

The rightturn/orientation predicate for three points in $\mathbb{R}^2$ has algebraic degree two.

The degree of a predicate depends on the algebraic expression used to evaluate it. Any such expression is intimately tied to the representation of the geometric objects used. Where not stated otherwise, we assume that geometric objects are represented as described in Section 1.2. So in the proposition above we assume that points are represented using Cartesian coordinates. The situation might be different, if, say, a polar representation is used instead.

But even once a representation is agreed upon, there is no obvious correspondence to an algebraic expression that describes a given predicate. In fact, it is not even clear why such an expression should exist in general. But if it does—as, for instance, in case of the orientation predicate—we prefer to work with a polynomial of smallest possible degree. Therefore we define the (algebraic) degree of a geometric predicate as the minimum degree of a polynomial that defines it (predicate true $\iff$ polynomial positive). So there still is something to be shown in order to complete Proposition 5.13.

Exercise 5.14 Show that there is no polynomial of degree at most one that describes the orientation test in $\mathbb{R}^2$.

Hint: Let $p = (0,0)$, $q = (x,0)$, and $r = (0,y)$ and show that there is no linear function in $x$ and $y$ that distinguishes the region where $pqr$ form a rightturn from its complement.

The degree of a predicate corresponds to the size of numbers that arise during its evaluation. If all input coordinates are $k$-bit integers then the numbers that occur during an evaluation of a degree $d$ predicate on these coordinates are of size about $1^{\text{d}k}$. If the number type used for the computation can represent all such numbers, the predicate can be evaluated exactly and thus always correctly. For instance, when using a standard IEEE double precision floating point implementation which has a mantissa length of 53 bit then the above orientation predicate can be evaluated exactly if the input coordinates are integers between 0 and $2^{25}$, say.

Let us now get back to the line segment intersection problem. It needs a few new geometric primitives: most prominently, constructing the intersection point of two line segments.

---

1It is only about $dk$ because not only multiplications play a role but also additions. As a rule of thumb, a multiplication may double the bitsize, while an addition may increase it by one.
Two segments. Given two line segments \( s = \lambda a + (1 - \lambda)b, \lambda \in [0,1] \) and \( t = \mu c + (1 - \mu)d, \mu \in [0,1] \), it is a simple exercise in linear algebra to compute \( s \cap t \). Note that \( s \cap t \) is either empty or a single point or a line segment.

For \( a = (a_x, a_y) \), \( b = (b_x, b_y) \), \( c = (c_x, c_y) \), and \( d = (d_x, d_y) \) we obtain two linear equations in two variables \( \lambda \) and \( \mu \).

\[
\begin{align*}
\lambda a_x + (1 - \lambda) b_x &= \mu c_x + (1 - \mu) d_x \\
\lambda a_y + (1 - \lambda) b_y &= \mu c_y + (1 - \mu) d_y
\end{align*}
\]

Rearranging terms yields

\[
\begin{align*}
\lambda(a_x - b_x) + \mu(d_x - c_x) &= d_x - b_x \\
\lambda(a_y - b_y) + \mu(d_y - c_y) &= d_y - b_y
\end{align*}
\]

Assuming that the lines underlying \( s \) and \( t \) have distinct slopes (that is, they are neither identical nor parallel) we have

\[
D = \begin{vmatrix} a_x - b_x & d_x - c_x \\ a_y - b_y & d_y - c_y \end{vmatrix} = \begin{vmatrix} a_x & a_y & 1 & 0 \\ b_x & b_y & 1 & 0 \\ c_x & c_y & 0 & 1 \\ d_x & d_y & 0 & 1 \end{vmatrix} \neq 0
\]

and using Cramer’s rule

\[
\lambda = \frac{1}{D} \begin{vmatrix} d_x - b_x & d_x - c_x \\ d_y - b_y & d_y - c_y \end{vmatrix} \quad \text{and} \quad \mu = \frac{1}{D} \begin{vmatrix} a_x - b_x & d_x - b_x \\ a_y - b_y & d_y - b_y \end{vmatrix}.
\]

To test if \( s \) and \( t \) intersect, we can—after having sorted out the degenerate case in which both segments have the same slope—compute \( \lambda \) and \( \mu \) and then check whether \( \lambda, \mu \in [0,1] \).

Observe that both \( \lambda \) and \( D \) result from multiplying two differences of input coordinates. Computing the \( x \)-coordinate of the point of intersection via \( b_x + \lambda(a_x - b_x) \) uses another multiplication. Overall this computation yields a fraction whose numerator is a polynomial of degree three in the input coordinates and whose denominator is a polynomial of degree two in the input coordinates.

In order to maintain the sorted order of event points in the EPS, we need to compare event points lexicographically. In case that both are intersection points, this amounts to comparing two fractions of the type discussed above. In order to formulate such a comparison as a polynomial, we have to cross-multiply the denominators, and so obtain a polynomial of degree \( 3 + 2 = 5 \). It can be shown (but we will not do it here) that this bound is tight and so we conclude that

**Proposition 5.15** The algebraic degree of the predicate that compares two intersection points of line segments lexicographically is five.
Therefore the coordinate range in which this predicate can be evaluated exactly using IEEE double precision numbers shrinks down to integers between 0 and about $2^{10} = 1'024$.

**Exercise 5.16** What is the algebraic degree of the predicate checking whether two line segments intersect? (Above we were interested in the actual intersection point, but now we consider the predicate that merely answers the question whether two segments intersect or not by yes or no).

**Exercise 5.17** The goal of this exercise is to analyze the algebraic degree of the InCircle predicate: Given three points $p, q, r \in \mathbb{R}^2$ that define a circle $C$ and a fourth point $s$, is $s$ located inside $C$ or not? To this end we employ the so-called parabolic lifting map, which will also play a prominent role in the next chapter of the course.

   a) The parabolic lifting map $\ell : \mathbb{R}^2 \to \mathbb{R}^3$ is defined for a point $p = (x, y) \in \mathbb{R}^2$ by $\ell(p) = (x, y, x^2 + y^2)$. For a circle $C \subseteq \mathbb{R}^2$ of positive radius, show that the “lifted circle” $\ell(C) = \{\ell(p) | p \in C\}$ is contained in a unique plane $h_C \subseteq \mathbb{R}^3$. Moreover, a point $p \in \mathbb{R}^2$ is strictly inside (outside, respectively) of $C$ if and only if the lifted point $\ell(p)$ is strictly below (above, respectively) $h_C$.

   b) What is the degree of the polynomial you need to evaluate in order to decide the InCircle predicate?

**5.5 Red-Blue Intersections**

Although the Bentley-Ottmann sweep appears to be rather simple, its implementation is not straightforward. For once, the original formulation did not take care of possible degeneracies—as we did in the preceding section. But also the algebraic degree of the predicates used is comparatively high. In particular, comparing two points of intersection lexicographically is a predicate of degree five, that is, in order to compute it, we need to evaluate a polynomial of degree five. When evaluating such a predicate with plain floating point arithmetic, one easily gets incorrect results due to limited precision roundoff errors. Such failures frequently render the whole computation useless. The line sweep algorithm is problematic in this respect, as a failure to detect one single point of intersection often implies that no other intersection of the involved segments to the right is found.

In general, predicates of degree four are needed to construct the arrangement of line segments because one needs to determine the orientation of a triangle formed by three segments. This is basically an orientation test where two points are segment endpoints and the third point is an intersection point of segments. Given that the coordinates of the latter are fractions, whose numerator is a degree three polynomial and the common denominator is a degree two polynomial, we obtain a degree four polynomial overall.

Motivated by the Map overlay application we consider here a restricted case. In the red-blue intersection problem, the input consists of two sets $R$ (red) and $B$ (blue) of
segments such that the segments in each set are *interior-disjoint*, that is, for any pair of distinct segments their relative interior is disjoint. In this case there are no triangles of intersecting segments, and it turns out that predicates of degree two suffice to construct the arrangement. This is optimal because already the intersection test for two segments is a predicate of degree two.

**Predicates of degree two.** Restricting to degree two predicates has certain consequences. While it is possible to determine the position of a segment endpoint relative to another segment using an orientation test, one cannot, for example, compare a segment endpoint with a point of intersection lexicographically. Even computing the coordinates for a point of intersection is not possible. Therefore the output of intersection points is done implicitly, as “intersection of $s$ and $t$”.

Graphically speaking we can deform any segment—keeping its endpoints fixed—as long as it remains monotone and it does not reach or cross any segment endpoint (it did not touch before). With help of degree two predicates there is no way to tell the difference.

**Witnesses.** Using such transformations the processing of intersection points is deferred as long as possible (lazy computation). The last possible point $w(s, t)$ where an intersection between two segments $s$ and $t$ has to be processed we call the *witness* of the intersection. The witness $w(s, t)$ is the lexicographically smallest segment endpoint that is located within the closed wedge formed by the two intersecting segments $s$ and $t$ to the right of the point of intersection (Figure 5.2). Note that each such wedge contains at least two segment endpoints, namely the right endpoints of the two intersecting segments. Therefore for any pair of intersecting segments its witness is well-defined.

As a consequence, only the segment endpoints are EPs and the EPS can be determined by lexicographic sorting during initialization. On the other hand, the SLS structure gets more complicated because its order of segments does not necessarily reflect the order in which the segments intersect the sweep line.

**Invariants.** The invariants of the algorithm have to take the relaxed notion of order into account. Denote the sweep line by $\ell$. 
1. \( L \) is the sequence of segments from \( S = R \cup B \) intersecting \( \ell \); \( s \) appears before \( t \) in \( L \implies s \) intersects \( \ell \) above \( t \) or \( s \) intersects \( t \) and the witness of this intersection is to the right of \( \ell \).

2. All intersections of segments from \( S \) whose witness is to the left of \( \ell \) have been reported.

**SLS Data Structure.** The SLS structure consist of three levels. We use the fact that segments of the same color do not interact, except by possibly sharing endpoints.

1. Collect adjacent segments of the same color in bundles, stored as balanced search trees. For each bundle store pointers to the topmost and bottommost segment. (As the segments within one bundle are interior-disjoint, their order remains static and thus correct under possible deformations due to lazy computation.)

2. All bundles are stored in a doubly linked list, sorted by \( y \)-coordinate.

3. All red bundles are stored in a balanced search tree (bundle tree).

![Graphical representation of the SLS data structure.](image)

The search tree structure should support insert, delete, split and merge in (amortized) logarithmic time each. For instance, splay trees \([12]\) meet these requirements. (If you have never heard about splay trees so far, you do not need to know for our purposes here. Just treat them as a black box. However, you should take a note and read about splay trees still. They are a fascinating data structure.)

**EP handling.** An EP \( p \) is processed as follows.

1. We first want to classify all bundles with respect to their position relative to \( p \): as either lying \textit{above} or \textit{below} \( p \) or as \textit{ending} at \( p \). There are \( \leq 2 \) bundles that have no clear such characterization.
(a) Localize \( p \) in bundle tree → Find \( \leq 2 \) bundles without clear characterization. For the localization we use the pointers to the topmost and bottommost segment within a bundle.

(b) Localize \( p \) in \( \leq 2 \) red bundles found and split them at \( p \). (If \( p \) is above the topmost or below the bottommost segment of the bundle, there is no split.) All red bundles are now either above, ending, or below with respect to \( p \).

(c) Localize \( p \) within the blue bundles by linear search.

(d) Localize \( p \) in the \( \leq 2 \) blue bundles found and split them at \( p \). (If \( p \) is above the topmost or below the bottommost segment of the bundle, there is no split.) All bundles are now either above, ending, or below with respect to \( p \).

2. Run through the list of bundles around \( p \). More precisely, start from one of the bundles containing \( p \) found in Step 1 and walk up in the doubly linked list of bundles until two successive bundles (hence of opposite color) are both above \( p \). Similarly, walk down in the doubly linked list of bundles, until two successive bundles are both below \( p \). Handle all adjacent pairs of bundles \((A, B)\) that are in wrong order and report all pairs of segments as intersecting. (Exchange \( A \) and \( B \) in the bundle list and merge them with their new neighbors.)

3. Report all two-colored pairs from \( \text{begin}(p) \times \text{end}(p) \) as intersecting.

4. Remove ending bundles and insert starting segments, sorted by slope and bundled by color and possibly merge with the closest bundle above or below.

Remark: As both the red and the blue segments are interior-disjoint, at every EP there can be at most one segment that passes through the EP. Should this happen, for the purpose of EP processing split this segment into two, one ending and one starting at this EP. But make sure to not report an intersection between the two parts!

Analysis. Sorting the EPS: \( O(n \log n) \) time. Every EP generates a constant number of tree searches and splits of \( O(\log n) \) each. Every exchange in Step 2 generates at least one intersection. New bundles are created only by inserting a new segment or by one of the constant number of splits at an EP. Therefore \( O(n) \) bundles are created in total. The total number of merge operations is \( O(n) \), as every merge kills one bundle and \( O(n) \) bundles are created overall. The linear search in steps 1 and 2 can be charged either to the ending bundle or—for continuing bundles—to the subsequent merge operation. In summary, we have a runtime of \( O(n \log n + k) \) and space usage is linear obviously.

**Theorem 5.18** For two sets \( R \) and \( B \), each consisting of interior-disjoint line segments in \( \mathbb{R}^2 \), one can find all intersecting pairs of segments in \( O(n \log n + k) \) time and linear space, using predicates of maximum degree two. Here \( n = |R| + |B| \) and \( k \) is the number of intersecting pairs.
Remarks. The first optimal algorithm for the red-blue intersection problem was published in 1988 by Harry Mairson and Jorge Stolfi [7]. In 1994 Timothy Chan [3] described a trapezoidal-sweep algorithm that uses predicates of degree three only. The approach discussed above is due to Andrea Mantler and Jack Snoeyink [8] from 2000.

Exercise 5.19 Let $S$ be a set of $n$ segments each of which is either horizontal or vertical. Describe an $O(n \log n)$ time and $O(n)$ space algorithm that counts the number of pairs in $\binom{n}{2}$ that intersect.

Questions

14. How can one test whether a polygon on $n$ vertices is simple? Describe an $O(n \log n)$ time algorithm.

15. How can one test whether two simple polygons on altogether $n$ vertices intersect? Describe an $O(n \log n)$ time algorithm.

16. How does the line sweep algorithm work that finds all $k$ intersecting pairs among $n$ line segments in $\mathbb{R}^2$? Describe the algorithm, using $O((n + k) \log n)$ time and $O(n)$ space. In particular, explain the data structures used, how event points are handled, and how to cope with degeneracies.

17. Given two line segments $s$ and $t$ in $\mathbb{R}^2$ whose endpoints have integer coordinates in $[0, 2^b)$; suppose $s$ and $t$ intersect in a single point $q$, what can you say about the coordinates of $q$? Give a good (tight up to small additive number of bits) upper bound on the size of these coordinates. (No need to prove tightness, that is, give an example which achieves the bound.)

18. What is the degree of a predicate and why is it an important parameter? What is the degree of the orientation test and the incircle test in $\mathbb{R}^2$? Explain the term and give two reasons for its relevance. Provide tight upper bounds for the two predicates mentioned. (No need to prove tightness.)

19. What is the map overlay problem and how can it be solved optimally using degree two predicates only? Give the problem definition and explain the term “interior-disjoint”. Explain the bundle-sweep algorithm, in particular, the data structures used, how an event point is processed, and the concept of witnesses.

References


Chapter 6

Delaunay Triangulations

In Chapter 2 we have discussed triangulations of simple polygons. A triangulation nicely partitions a polygon into triangles, which allows, for instance, to easily compute the area or a guarding of the polygon. Another typical application scenario is to use a triangulation $T$ for interpolation: Suppose a function $f$ is defined on the vertices of the polygon $P$, and we want to extend it “reasonably” and continuously to $P^\circ$. Then for a point $p \in P^\circ$ find a triangle $t$ of $T$ that contains $p$. As $p$ can be written as a convex combination $\sum_{i=1}^{3} \lambda_i v_i$ of the vertices $v_1, v_2, v_3$ of $t$, we just use the same coefficients to obtain an interpolation $f(p) := \sum_{i=1}^{3} \lambda_i f(v_i)$ of the function values.

If triangulations are a useful tool when working with polygons, they might also turn out useful to deal with other geometric objects, for instance, point sets. But what could be a triangulation of a point set? Polygons have a clearly defined interior, which naturally lends itself to be covered by smaller polygons such as triangles. A point set does not have an interior, except... Here the notion of convex hull comes handy, because it allows us to treat a point set as a convex polygon. Actually, not really a convex polygon, because points in the interior of the convex hull should not be ignored completely. But one way to think of a point set is as a convex polygon—its convex hull—possibly with some holes—which are points—in its interior. A triangulation should then partition the convex hull while respecting the points in the interior, as shown in the example in Figure 6.1b.

![Figure 6.1: Examples of (non-)triangulations.](image)

(a) Simple polygon triangulation.  (b) Point set triangulation.  (c) Not a triangulation.

In contrast, the example depicted in Figure 6.1c nicely subdivides the convex hull
but should not be regarded a triangulation: Two points in the interior are not respected
but simply swallowed by a large triangle.
This interpretation directly leads to the following adaption of Definition 2.8.

Definition 6.1 A triangulation of a finite point set \( P \subset \mathbb{R}^2 \) is a collection \( T \) of triangles, such that

1. \( \text{conv}(P) = \bigcup_{T \in T} T \);
2. \( P = \bigcup_{T \in T} V(T) \); and
3. for every distinct pair \( T, U \in T \), the intersection \( T \cap U \) is either a common
   vertex, or a common edge, or empty.

Just as for polygons, triangulations are universally available for point sets, meaning
that (almost) every point set admits at least one.

Proposition 6.2 Every set \( P \subseteq \mathbb{R}^2 \) of \( n \geq 3 \) points has a triangulation, unless all points
in \( P \) are collinear.

Proof. In order to construct a triangulation for \( P \), consider the lexicographically sorted
sequence \( p_1, \ldots, p_n \) of points in \( P \). Let \( m \) be minimal such that \( p_1, \ldots, p_m \) are not
collinear. We triangulate \( p_1, \ldots, p_m \) by connecting \( p_m \) to all of \( p_1, \ldots, p_{m-1} \) (which are
on a common line), see Figure 6.2a

![Figure 6.2: Constructing the scan triangulation of \( P \).](image)

(a) Getting started.
(b) Adding a point.

Then we add \( p_{m+1}, \ldots, p_n \). When adding \( p_i \), for \( i > m \), we connect \( p_i \) with all
vertices of \( C_{i-1} := \text{conv}(\{p_1, \ldots, p_{i-1}\}) \) that it “sees”, that is, every vertex \( v \) of \( C_{i-1} \) for
which \( p_i \cap V \cap C_{i-1} = \{v\} \). In particular, among these vertices are the two points of tangency
from \( p_i \) to \( C_{i-1} \), which shows that we always add triangles (Figure 6.2b) whose union
after each step covers \( C_1 \). □

The triangulation that is constructed in Proposition 6.2 is called a scan triangulation.
Such a triangulation (Figure 6.3a (left) shows a larger example) is usually “ugly”, though,
since it tends to have many long and skinny triangles. This is not just an aesthetic deficit.
Having long and skinny triangles means that the vertices of a triangle tend to be spread
out far from each other. You can probably imagine that such a behavior is undesirable,
for instance, in the context of interpolation. In contrast, the Delaunay triangulation of the same point set (Figure 6.3b) looks much nicer, and we will discuss in the next section how to get this triangulation.

Exercise 6.3 Describe an $O(n \log n)$ time algorithm to construct a scan triangulation for a set of $n$ points in $\mathbb{R}^2$.

On another note, if you look closely into the SLR-algorithm to compute planar convex hull that was discussed in Chapter 3, then you will realize that we also could have used this algorithm in the proof of Proposition 6.2. Whenever a point is discarded during SLR, a triangle is added to the polygon that eventually becomes the convex hull.

In view of the preceding chapter, we may regard a triangulation as a plane graph: the vertices are the points in $P$ and there is an edge between two points $p \neq q$, if and only if there is a triangle with vertices $p$ and $q$. Therefore we can use Euler’s formula to determine the number of edges in a triangulation.

Lemma 6.4 Any triangulation of a set $P \subset \mathbb{R}^2$ of $n$ points has exactly $3n-h-3$ edges, where $h$ is the number of points from $P$ on $\partial \text{conv}(P)$.

Proof. Consider a triangulation $T$ of $P$ and denote by $E$ the set of edges and by $F$ the set of faces of $T$. We count the number of edge-face incidences in two ways. Denote $J = \{(e, f) \in E \times F : e \subset \partial f\}$.

On the one hand, every edge is incident to exactly two faces and therefore $|J| = 2|E|$. On the other hand, every bounded face of $T$ is a triangle and the unbounded face has $h$ edges on its boundary. Therefore, $|J| = 3(|F| - 1) + h$. 67
Together we obtain $3|F| = 2|E| - h + 3$. Using Euler's formula $(3n - 3|E| + 3|F| = 6)$ we conclude that $3n - |E| - h + 3 = 6$ and so $|E| = 3n - h - 3$. \hfill \Box

In graph theory, the term “triangulation” is sometimes used as a synonym for “maximal planar”. But geometric triangulations are different, they are maximal planar in the sense that no straight-line edge can be added without sacrificing planarity.

**Corollary 6.5** A triangulation of a set $P \subset \mathbb{R}^2$ of $n$ points is maximal planar, if and only if $\text{conv}(P)$ is a triangle.

**Proof.** Combine Lemma 4.1 and Lemma 6.4. \hfill \Box

**Exercise 6.6** Find for every $n \geq 3$ a simple polygon $P$ with $n$ vertices such that $P$ has exactly one triangulation. $P$ should be in general position, meaning that no three vertices are collinear.

**Exercise 6.7** Show that every set of $n \geq 5$ points in general position (no three points are collinear) has at least two different triangulations.

**Hint:** Show first that every set of five points in general position contains a convex 4-hole, that is, a subset of four points that span a convex quadrilateral that does not contain the fifth point.

### 6.1 The Empty Circle Property

We will now move on to study the ominous and supposedly nice Delaunay triangulations mentioned above. They are defined in terms of an empty circumcircle property for triangles. The *circumcircle* of a triangle is the unique circle passing through the three vertices of the triangle, see Figure 6.4.

![Circumcircle of a triangle](image)

**Figure 6.4:** Circumcircle of a triangle.

**Definition 6.8** A triangulation of a finite point set $P \subset \mathbb{R}^2$ is called a Delaunay triangulation, if the circumcircle of every triangle is empty, that is, there is no point from $P$ in its interior.
Consider the example depicted in Figure 6.5. It shows a Delaunay triangulation of a set of six points: The circumcircles of all five triangles are empty (we also say that the triangles satisfy the empty circle property). The dashed circle is not empty, but that is fine, since it is not a circumcircle of any triangle.

It is instructive to look at the case of four points in convex position. Obviously, there are two possible triangulations, but in general, only one of them will be Delaunay, see Figure 6.6a and 6.6b. If the four points are on a common circle, though, this circle is empty; at the same time it is the circumcircle of all possible triangles; therefore, both triangulations of the point set are Delaunay, see Figure 6.6c.

Proposition 6.9 Given a set $P \subset \mathbb{R}^2$ of four points that are in convex position but not cocircular. Then $P$ has exactly one Delaunay triangulation.

Proof. Consider a convex polygon $P = pqrs$. There are two triangulation of $P$: a triangulation $T_1$ using the edge $pr$ and a triangulation $T_2$ using the edge $qs$.

Consider the family $C_1$ of circles through $pr$, which contains the circumcircles $C_1 = pqr$ and $C'_1 = rps$ of the triangles in $T_1$. By assumption $s$ is not on $C_1$. If $s$ is outside of
Chapter 6. Delaunay Triangulations

C₁, then q is outside of C₁': Consider the process of continuously moving from C₁ to C₁' in C₁ (Figure 6.7a), the point q is “left behind” immediately when going beyond C₁ and only the final circle C₁' “grabs” the point s.

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

Figure 6.7: Circumcircles and containment for triangulations of four points.

Similarly, consider the family C₂ of circles through pq, which contains the circumcircles C₁ = pqr and C₂ = spq, the latter belonging to a triangle in T₂. As s is outside of C₁, it follows that r is inside C₂: Consider the process of continuously moving from C₁ to C₂ in C₂ (Figure 6.7b), the point r is on C₁ and remains within the circle all the way up to C₂. This shows that T₁ is Delaunay, whereas T₂ is not.

The case that s is located inside C₁ is symmetric: just cyclically shift the roles of pqrs to qrsp. □

6.2 The Lawson Flip algorithm

It is not clear yet that every point set actually has a Delaunay triangulation (given that not all points are on a common line). In this and the next two sections, we will prove that this is the case. The proof is algorithmic. Here is the Lawson flip algorithm for a set P of n points.

1. Compute some triangulation of P (for example, the scan triangulation).

2. While there exists a subtriangulation of four points in convex position that is not Delaunay (like in Figure 6.6b), replace this subtriangulation by the other triangulation of the four points (Figure 6.6a).

We call the replacement operation in the second step a (Lawson) flip.

**Theorem 6.10** Let P ⊆ R² be a set of n points, equipped with some triangulation T. The Lawson flip algorithm terminates after at most \( \binom{n}{2} = O(n²) \) flips, and the resulting triangulation D is a Delaunay triangulation of P.
We will prove Theorem 6.10 in two steps: First we show that the program described above always terminates and, therefore, is an algorithm, indeed (Section 6.3). Then we show that the algorithm does what it claims to do, namely the result is a Delaunay triangulation (Section 6.4).

6.3 Termination of the Lawson Flip Algorithm: The Lifting Map

In order to prove Theorem 6.10, we invoke the (parabolic) lifting map. This is the following: given a point \( p = (x, y) \in \mathbb{R}^2 \), its lifting \( \ell(p) \) is the point

\[
\ell(p) = (x, y, x^2 + y^2) \in \mathbb{R}^3.
\]

Geometrically, \( \ell \) "lifts" the point vertically up until it lies on the unit paraboloid

\[
\{(x, y, z) \mid z = x^2 + y^2\} \subseteq \mathbb{R}^3,
\]

see Figure 6.8a.

Recall the following important property of the lifting map that we proved in Exercise 5.17. It is illustrated in Figure 6.8b.

Lemma 6.11 Let \( C \subseteq \mathbb{R}^2 \) be a circle of positive radius. The "lifted circle" \( \ell(C) = \{\ell(p) \mid p \in C\} \) is contained in a unique plane \( h_C \subseteq \mathbb{R}^3 \). Moreover, a point \( p \in \mathbb{R}^2 \) is strictly inside (outside, respectively) of \( C \) if and only if the lifted point \( \ell(p) \) is strictly below (above, respectively) \( h_C \).

Using the lifting map, we can now prove Theorem 6.10. Let us fix the point set \( P \) for this and the next section. First, we need to argue that the algorithm indeed terminates (if you think about it a little, this is not obvious). So let us interpret a flip operation in
the lifted picture. The flip involves four points in convex position in $\mathbb{R}^2$, and their lifted images form a tetrahedron in $\mathbb{R}^3$ (think about why this tetrahedron cannot be “flat”).

The tetrahedron is made up of four triangles; when you look at it from the top, you see two of the triangles, and when you look from the bottom, you see the other two. In fact, what you see from the top and the bottom are the lifted images of the two possible triangulations of the four-point set in $\mathbb{R}^2$ that is involved in the flip.

Here is the crucial fact that follows from Lemma 6.11: The two top triangles come from the non-Delaunay triangulation before the flip, see Figure 6.9a. The reason is that both top triangles have the respective fourth point below them, meaning that in $\mathbb{R}^2$, the circumcircles of these triangles contain the respective fourth point—the empty circle property is violated. In contrast, the bottom two triangles come from the Delaunay triangulation of the four points: they both have the respective fourth point above them, meaning that in $\mathbb{R}^2$, the circumcircles of the triangles do not contain the respective fourth point, see Figure 6.9b.

![Figure 6.9: Lawson flip: the height of the surface of lifted triangles decreases.](image)

In the lifted picture, a Lawson flip can therefore be interpreted as an operation that replaces the top two triangles of a tetrahedron by the bottom two ones. If we consider the lifted image of the current triangulation, we therefore have a surface in $\mathbb{R}^3$ whose pointwise height can only decrease through Lawson flips. In particular, once an edge has been flipped, this edge will be strictly above the resulting surface and can therefore never be flipped a second time. Since $n$ points can span at most $\binom{n}{2}$ edges, the bound on the number of flips follows.

### 6.4 Correctness of the Lawson Flip Algorithm

It remains to show that the triangulation of $P$ that we get upon termination of the Lawson flip algorithm is indeed a Delaunay triangulation. Here is a first observation telling us that the triangulation is “locally Delaunay”.

1. Before the flip: the top two triangles of the tetrahedron and the corresponding non-Delaunay triangulation in the plane.
2. After the flip: the bottom two triangles of the tetrahedron and the corresponding Delaunay triangulation in the plane.

Figure 6.9: Lawson flip: the height of the surface of lifted triangles decreases.
Observation 6.12 Let $\Delta, \Delta'$ be two adjacent triangles in the triangulation $\mathcal{D}$ that results from the Lawson flip algorithm. Then the circumcircle of $\Delta$ does not have any vertex of $\Delta'$ in its interior, and vice versa.

If the two triangles together form a convex quadrilateral, this follows from the fact that the Lawson flip algorithm did not flip the common edge of $\Delta$ and $\Delta'$. If the four vertices are not in convex position, this is basic geometry: given a triangle $\Delta$, its circumcircle $C$ can only contain points of $C \setminus \Delta$ that form a convex quadrilateral with the vertices of $\Delta$.

Now we show that the triangulation is also "globally Delaunay".

Proposition 6.13 The triangulation $\mathcal{D}$ that results from the Lawson flip algorithm is a Delaunay triangulation.

Proof. Suppose for contradiction that there is some triangle $\Delta \in \mathcal{D}$ and some point $p \in P$ strictly inside the circumcircle $C$ of $\Delta$. Among all such pairs $(\Delta, p)$, we choose one for which we the distance of $p$ to $\Delta$ is minimal. Note that this distance is positive since $\mathcal{D}$ is a triangulation of $P$. The situation is as depicted in Figure 6.10a.

Now consider the edge $e$ of $\Delta$ that is facing $p$. There must be another triangle $\Delta' \in \mathcal{D}$ that is incident to the edge $e$. By the local Delaunay property of $\mathcal{D}$, the third vertex $q$ of $\Delta'$ is on or outside of $C$, see Figure 6.10b. But then the circumcircle $C'$ of $\Delta'$ contains the whole portion of $C$ on $p$'s side of $e$, hence it also contains $p$; moreover, $p$ is closer to $\Delta'$ than to $\Delta$ (Figure 6.10c). But this is a contradiction to our choice of $\Delta$ and $p$. Hence there was no $(\Delta, p)$, and $\mathcal{D}$ is a Delaunay triangulation.

Exercise 6.14 The Euclidean minimum spanning tree (EMST) of a finite point set $P \subset \mathbb{R}^2$ is a spanning tree for which the sum of the edge lengths is minimum (among all spanning trees of $P$). Show:

Figure 6.10: Correctness of the Lawson flip algorithm.
a) Every EMST of $P$ is a plane graph.

b) Every EMST of $P$ contains a closest pair, i.e., an edge between two points $p, q \in P$ that have minimum distance to each other among all point pairs in $(P^2)$.

c) Every Delaunay Triangulation of $P$ contains an EMST of $P$.

6.5 The Delaunay Graph

Despite the fact that a point set may have more than one Delaunay triangulation, there are certain edges that are present in every Delaunay triangulation, for instance, the edges of the convex hull.

**Definition 6.15** The Delaunay graph of $P \subseteq \mathbb{R}^2$ consists of all line segments $pq$, for $p, q \in P$, that are contained in every Delaunay triangulation of $P$.

The following characterizes the edges of the Delaunay graph.

**Lemma 6.16** The segment $pq$, for $p, q \in P$, is in the Delaunay graph of $P$ if and only if there exists a circle through $p$ and $q$ that has $p$ and $q$ on its boundary and all other points of $P$ are strictly outside.

**Proof.** “⇒”: Let $pq$ be an edge in the Delaunay graph of $P$, and let $D$ be a Delaunay triangulation of $P$. Then there exists a triangle $\Delta = pqr$ in $D$, whose circumcircle $C$ does not contain any point from $P$ in its interior.

If there is a point $s$ on $\partial C$ such that $rs$ intersects $pq$, then let $\Delta' = pqt$ denote the other ($\neq \Delta$) triangle in $D$ that is incident to $pq$ (Figure 6.11a). Flipping the edge $pq$ to $rt$ yields another Delaunay triangulation of $P$ that does not contain the edge $pq$, in contradiction to $pq$ being an edge in the Delaunay graph of $P$. Therefore, there is no such point $s$.

![Diagram](a) Another point $s \in \partial C$.  

![Diagram](b) Moving $C$ away from $s$.

**Figure 6.11:** Characterization of edges in the Delaunay graph (I).
Otherwise we can slightly change the circle \( C \) by moving away from \( r \) while keeping \( p \) and \( q \) on the circle. As \( P \) is a finite point set, we can do such a modification without catching another point from \( P \) with the circle. In this way we obtain a circle \( C' \) through \( p \) and \( q \) such that all other points from \( P \) are strictly outside \( C' \) (Figure 6.12b).

\[
\text{⇐: Let } D \text{ be a Delaunay triangulation of } P. \text{ If } pq \text{ is not an edge of } D, \text{ there must be another edge of } D \text{ that crosses } pq \text{ (otherwise, we could add } pq \text{ to } D \text{ and still have a plane graph, a contradiction to } D \text{ being a triangulation of } P). \text{ Let } rs \text{ denote the first edge of } D \text{ that intersects the directed line segment } pq. \]

Consider the triangle \( \Delta \) of \( D \) that is incident to \( rs \) on the side that faces \( p \) (given that \( rs \) intersects \( pq \) this is a well defined direction). By the choice of \( rs \) neither of the other two edges of \( \Delta \) intersects \( pq \), and \( p \notin \Delta^o \) because \( \Delta \) is part of a triangulation of \( P \). The only remaining option is that \( p \) is a vertex of \( \Delta = prs \). As \( \Delta \) is part of a Delaunay triangulation, its circumcircle \( C_{\Delta} \) is empty (i.e., \( C_{\Delta}^o \cap P = \emptyset \)).

Consider now a circle \( C \) through \( p \) and \( q \), which exists by assumption. Fixing \( p \) and \( q \), expand \( C \) towards \( r \) to eventually obtain the circle \( C' \) through \( p \), \( q \), and \( r \) (Figure 6.12a). Recall that \( r \) and \( s \) are on different sides of the line through \( p \) and \( q \). Therefore, \( s \) lies strictly outside of \( C' \). Next fix \( p \) and \( r \) and expand \( C' \) towards \( s \) to eventually obtain the circle \( C_{\Delta} \) through \( p \), \( r \), and \( s \) (Figure 6.12b). Recall that \( s \) and \( q \) are on the same side of the line through \( p \) and \( r \). Therefore, \( q \in C_{\Delta} \), which is in contradiction to \( C_{\Delta} \) being empty. It follows that there is no Delaunay triangulation of \( P \) that does not contain the edge \( pq \).

\[\begin{align*}
\text{(a) Expanding } C \text{ towards } r. \\
\text{(b) Expanding } C' \text{ towards } s.
\end{align*}\]

**Figure 6.12:** Characterization of edges in the Delaunay graph (II).

The Delaunay graph is useful to prove uniqueness of the Delaunay triangulation in case of general position.

**Corollary 6.17** Let \( P \subset \mathbb{R}^2 \) be a finite set of points in general position, that is, no four points of \( P \) are cocircular. Then \( P \) has a unique Delaunay triangulation. \( \square \)
6.6 Every Delaunay Triangulation Maximizes the Smallest Angle

Why are we actually interested in Delaunay triangulations? After all, having empty circumcircles is not a goal in itself. But it turns out that Delaunay triangulations satisfy a number of interesting properties. Here we show just one of them.

Recall that when we compared a scan triangulation with a Delaunay triangulation of the same point set in Figure 6.3, we claimed that the scan triangulation is “ugly” because it contains many long and skinny triangles. The triangles of the Delaunay triangulation, at least in this example, look much nicer, that is, much closer to an equilateral triangle. One way to quantify this “niceness” is to look at the angles that appear in a triangulation: If all angles are large, then all triangles are reasonably close to an equilateral triangle. Indeed, we will show that Delaunay triangulations maximize the smallest angle among all triangulations of a given point set. Note that this does not imply that there are no long and skinny triangles in a Delaunay triangulation. But if there is a long and skinny triangle in a Delaunay triangulation, then there is an at least as long and skinny triangle in every triangulation of the point set.

Given a triangulation \( T \) of \( P \), consider the sorted sequence \( A(T) = (\alpha_1, \alpha_2, \ldots, \alpha_m) \) of interior angles, where \( m \) is the number of triangles (we have already remarked earlier that \( m \) is a function of \( P \) only and does not depend on \( T \)). Being sorted means that \( \alpha_1 \leq \alpha_2 \leq \cdots \leq \alpha_m \). Let \( T, T' \) be two triangulations of \( P \). We say that \( A(T) < A(T') \) if there exists some \( i \) for which \( \alpha_i < \alpha'_i \) and \( \alpha_j = \alpha'_j \), for all \( j < i \). (This is nothing but the lexicographic order on these sequences.)

**Theorem 6.18** Let \( P \subseteq \mathbb{R}^2 \) be a finite set of points in general position (not all collinear and no four cocircular). Let \( D^* \) be the unique Delaunay triangulation of \( P \), and let \( T \) be any triangulation of \( P \). Then \( A(T) \leq A(D^*) \).

In particular, \( D^* \) maximizes the smallest angle among all triangulations of \( P \).

![Diagram](a) Four cocircular points and the induced eight angles. (b) The situation before a flip.

**Figure 6.13:** Angle-optimality of Delaunay triangulations.
6.6. Every Delaunay Triangulation Maximizes the Smallest Angle

**Proof.** We know that $T$ can be transformed into $D^*$ through the Lawson flip algorithm, and we are done if we can show that each such flip lexicographically increases the sorted angle sequence. A flip replaces six interior angles by six other interior angles, and we will actually show that the smallest of the six angles strictly increases under the flip. This implies that the whole angle sequence increases lexicographically.

Let us first look at the situation of four cocircular points, see Figure 6.13a. In this situation, the *inscribed angle theorem* (a generalization of Thales’ Theorem, stated below as Theorem 6.19) tells us that the eight depicted angles come in four equal pairs. For instance, the angles labeled $\alpha_1$ at $s$ and $r$ are angles on the same side of the chord $pq$ of the circle.

In Figure 6.13b, we have the situation in which we perform a Lawson flip (replacing the solid with the dashed diagonal). By the symbol $\alpha$ ($\bar{\alpha}$, respectively) we denote an angle strictly smaller (larger, respectively) than $\alpha$. Here are the six angles before the flip:

$$\alpha_1 + \alpha_2, \quad \alpha_3, \quad \alpha_4, \quad \alpha_1, \quad \alpha_2, \quad \bar{\alpha}_3 + \bar{\alpha}_4.$$ 

After the flip, we have

$$\alpha_1, \quad \alpha_2, \quad \bar{\alpha}_3, \quad \bar{\alpha}_4, \quad \alpha_1 + \alpha_4, \quad \alpha_2 + \alpha_3.$$ 

Now, for every angle after the flip there is at least one smaller angle before the flip:

$$\alpha_1 > \alpha_1,$$
$$\alpha_2 > \alpha_2,$$
$$\bar{\alpha}_3 > \alpha_3,$$
$$\bar{\alpha}_4 > \alpha_4,$$
$$\alpha_1 + \alpha_4 > \alpha_4,$$
$$\alpha_2 + \alpha_3 > \alpha_3.$$ 

It follows that the smallest angle strictly increases. ☐

**Theorem 6.19 (Inscribed Angle Theorem)** Let $C$ be a circle with center $c$ and positive radius and $p, q \in C$. Then the angle $\angle prq \mod \pi = \frac{1}{2} \angle pcq$ is the same, for all $r \in C$.

**Proof.** Without loss of generality we may assume that $c$ is located to the left of or on the oriented line $pq$. 

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Consider first the case that the triangle \( \triangle = \text{pqr} \) contains \( c \). Then \( \Delta \) can be partitioned into three triangles: \( \text{pcr}, \text{qcr}, \text{and cpq} \). All three triangles are isosceles, because two sides of each form the radius of C. Denote \( \alpha = \angle \text{prc}, \beta = \angle \text{crq}, \gamma = \angle \text{cpq}, \) and \( \delta = \angle \text{pcq} \) (see the figure shown to the right). The angles we are interested in are \( \theta = \angle \text{prq} = \alpha + \beta \) and \( \delta \), for which we have to show that \( \delta = 2\theta \).

Indeed, the angle sum in \( \Delta \) is \( \pi = 2(\alpha + \beta + \gamma) \) and the angle sum in the triangle \( \text{cpq} \) is \( \pi = \delta + 2\gamma \). Combining both yields \( \delta = 2(\alpha + \beta) = 2\theta \).

Next suppose that \( \text{pqcr} \) are in convex position and \( r \) is to the left of or on the oriented line \( \text{pq} \). Without loss of generality let \( r \) be to the left of or on the oriented line \( \text{qc} \). (The case that \( r \) lies to the right of or on the oriented line \( \text{pc} \) is symmetric.) Define \( \alpha, \beta, \gamma, \delta \) as above and observe that \( \theta = \alpha - \beta \). Again have to show that \( \delta = 2\theta \).

The angle sum in the triangle \( \text{cpq} \) is \( \pi = \delta + 2\gamma \) and the angle sum in the triangle \( \text{rqp} \) is \( \pi = (\alpha - \beta) + \alpha + \gamma + (\gamma - \beta) = 2(\alpha + \gamma - \beta) \). Combining both yields \( \delta = \pi - 2\gamma = 2(\alpha - \beta) = 2\theta \).
It remains to consider the case that \( r \) is to the right of the oriented line \( pq \).

Consider the point \( r' \) that is antipodal to \( r \) on \( C \), and the quadrilateral \( Q = prqr' \). We are interested in the angle \( \phi \) of \( Q \) at \( r \). By Thales' Theorem the inner angles of \( Q \) at \( p \) and \( q \) are both \( \pi/2 \). Hence the angle sum of \( Q \) is \( 2\pi = \theta + \phi + 2\pi/2 \) and so \( \phi = \pi - \theta \).

What happens in the case where the Delaunay triangulation is not unique? The following still holds.

**Theorem 6.20** Let \( P \subseteq \mathbb{R}^2 \) be a finite set of points, not all on a line. Every Delaunay triangulation \( D \) of \( P \) maximizes the smallest angle among all triangulations \( T \) of \( P \).

**Proof.** Let \( D \) be some Delaunay triangulation of \( P \). We infinitesimally perturb the points in \( P \) such that no four are on a common circle anymore. Then the Delaunay triangulation becomes unique (Corollary 6.17). Starting from \( D \), we keep applying Lawson flips until we reach the unique Delaunay triangulation \( D^* \) of the perturbed point set. Now we examine this sequence of flips on the original *unperturbed* point set. All these flips must involve four cocircular points (only in the cocircular case, an infinitesimal perturbation can change “good” edges into “bad” edges that still need to be flipped). But as Figure 6.13 (a) easily implies, such a “degenerate” flip does not change the smallest of the six involved angles. It follows that \( D \) and \( D^* \) have the same smallest angle, and since \( D^* \) maximizes the smallest angle among all triangulations \( T \) (Theorem 6.18), so does \( D \). \( \square \)

### 6.7 Constrained Triangulations

Sometimes one would like to have a Delaunay triangulation, but certain edges are already prescribed, for example, a Delaunay triangulation of a simple polygon. Of course, one cannot expect to be able to get a proper Delaunay triangulation where all triangles satisfy the empty circle property. But it is possible to obtain some triangulation that comes as close as possible to a proper Delaunay triangulation, given that we are forced to include the edges in \( E \). Such a triangulation is called a *constrained Delaunay triangulation*, a formal definition of which follows.

Let \( P \subseteq \mathbb{R}^2 \) be a finite point set and \( G = (P, E) \) a geometric graph with vertex set \( P \) (we consider the edges \( e \in E \) as line segments). A triangulation \( T \) of \( P \) *respects* \( G \) if it contains all segments \( e \in E \). A triangulation \( T \) of \( P \) that respects \( G \) is said to be a *constrained Delaunay triangulation* of \( P \) with respect to \( G \) if the following holds for every triangle \( \Delta \) of \( T \):
The circumcircle of $\Delta$ contains only points $q \in P$ in its interior that are not visible from the interior of $\Delta$. A point $q \in P$ is visible from the interior of $\Delta$ if there exists a point $p$ in the interior of $\Delta$ such that the line segment $pq$ does not intersect any segment $e \in E$. We can thus imagine the line segments of $E$ as “blocking the view”.

For illustration, consider the simple polygon and its constrained Delaunay triangulation shown in Figure 6.15. The circumcircle of the shaded triangle $\Delta$ contains a whole other triangle in its interior. But these points cannot be seen from $\Delta^\circ$, because all possible connecting line segments intersect the blocking polygon edge $e$ of $\Delta$.

**Figure 6.15:** Constrained Delaunay triangulation of a simple polygon.

**Theorem 6.21** For every finite point set $P$ and every plane graph $G = (P, E)$, there exists a constrained Delaunay triangulation of $P$ with respect to $G$.

**Exercise 6.22** Prove Theorem 6.21. Also describe a polynomial algorithm to construct such a triangulation.

**Questions**

20. **What is a triangulation?** Provide the definition and prove a basic property: every triangulation with the same set of vertices and the same outer face has the same number of triangles.

21. **What is a triangulation of a point set?** Give a precise definition.
22. Does every point set (not all points on a common line) have a triangulation? You may, for example, argue with the scan triangulation.

23. What is a Delaunay triangulation of a set of points? Give a precise definition.

24. What is the Delaunay graph of a point set? Give a precise definition and a characterization.

25. How can you prove that every set of points (not all on a common line) has a Delaunay triangulation? You can for example sketch the Lawson flip algorithm and the Lifting Map, and use these to show the existence.

26. When is the Delaunay triangulation of a point set unique? Show that general position is a sufficient condition. Is it also necessary?

27. What can you say about the “quality” of a Delaunay triangulation? Prove that every Delaunay triangulation maximizes the smallest interior angle in the triangulation, among the set of all triangulations of the same point set.
Chapter 7

Delaunay Triangulation: Incremental Construction

In the last lecture, we have learned about the Lawson flip algorithm that computes a Delaunay triangulation of a given $n$-point set $P \subseteq \mathbb{R}^2$ with $O(n^2)$ Lawson flips. One can actually implement this algorithm to run in $O(n^2)$ time, and there are point sets where it may take $\Omega(n^2)$ flips.

In this lecture, we will discuss a different algorithm. The final goal is to show that this algorithm can be implemented to run in $O(n \log n)$ time; this lecture, however, is concerned only with the correctness of the algorithm. Throughout the lecture we assume that $P$ is in general position (no 3 points on a line, no 4 points on a common circle), so that the Delaunay triangulation is unique (Corollary 6.17). There are techniques to deal with non-general position, but we don’t discuss them here.

7.1 Incremental construction

The idea is to build the Delaunay triangulation of $P$ by inserting one point after another. We always maintain the Delaunay triangulation of the point set $R$ inserted so far, and when the next point $s$ comes along, we simply update the triangulation to the Delaunay triangulation of $R \cup \{s\}$. Let $DT(R)$ denote the Delaunay triangulation of $R \subseteq P$.

To avoid special cases, we enhance the point set $P$ with three artificial points “far out”. The convex hull of the resulting point set is a triangle; later, we can simply remove the extra points and their incident edges to obtain $DT(P)$. The incremental algorithm starts off with the Delaunay triangulation of the three artificial points which consists of one big triangle enclosing all other points. (In our figures, we suppress the far-away points, since they are merely a technicality.)

Now assume that we have already built $DT(R)$, and we next insert $s \in P \setminus R$. Here is the outline of the update step.

1. Find the triangle $\Delta = \Delta(p, q, r)$ of $DT(R)$ that contains $s$, and replace it with the three triangles resulting from connecting $s$ with all three vertices $p, q, r$; see Figure
We now have a triangulation $\mathcal{T}$ of $R \cup \{s\}$.

2. Perform Lawson flips on $\mathcal{T}$ until $\mathcal{DT}(R \cup \{s\})$ is obtained; see Figure 7.2

How to organize the Lawson flips. The Lawson flips can be organized quite systematically, since we always know the candidates for “bad” edges that may still have to be flipped. Initially (after step 1), only the three edges of $\Delta$ can be bad, since these are the only edges for which an incident triangle has changed (by inserting $s$ in Step 1). Each of the three new edges is good, since the 4 vertices of its two incident triangles are not in convex position.

Now we have the following invariant (part (a) certainly holds in the first flip):
(a) In every flip, the convex quadrilateral $Q$ in which the flip happens has exactly two edges incident to $s$, and the flip generates a new edge incident to $s$.

(b) Only the two edges of $Q$ that are \textit{not} incident to $s$ can become bad through the flip.

We will prove part (b) in the next lemma. The invariant then follows since (b) entails (a) in the next flip. This means that we can maintain a queue of potentially bad edges that we process in turn. A good edge will simply be removed from the queue, and a bad edge will be flipped and replaced according to (b) with two new edges in the queue. In this way, we never flip edges incident to $s$; the next lemma proves that this is correct and at the same time establishes part (b) of the invariant.

**Lemma 7.1** Every edge incident to $s$ that is created during the update is an edge of the Delaunay graph of $R \cup \{s\}$ and thus an edge that will be in $DT(R \cup \{s\})$. It easily follows that edges incident to $s$ will never become bad during the update step\(^1\).

**Proof.** Let us consider one of the first three new edges, $sp$, say. Since the triangle $\Delta$ has a circumcircle $C$ strictly containing only $s$ ($\Delta$ is in $DT(R)$), we can shrink that circumcircle to a circle $C'$ through $s$ and $p$ with no interior points, see Figure 7.3 (a). This proves that $sp$ is in the Delaunay graph. If $st$ is an edge created by a flip, a similar argument works. The flip destroys exactly one triangle $\Delta$ of $DT(R)$. Its circumcircle $C$ contains $s$ only, and shrinking it yields an empty circle $C'$ through $s$ and $t$. Thus, $st$ is in the Delaunay graph also in this case. \(\square\)

![Figure 7.3](attachment:image.png)

**Figure 7.3:** Newly created edges incident to $s$ are in the Delaunay graph

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\(^1\)If such an edge was bad, it could be flipped, but then it would be “gone forever” according to the lifting map interpretation from the previous lecture.
7.2 The History Graph

What can we say about the performance of the incremental construction? Not much yet. First of all, we did not specify how we find the triangle \( \Delta \) of \( DT(R) \) that contains the point \( s \) to be inserted. Doing this in the obvious way (checking all triangles) is not good, since already the find steps would then amount to \( O(n^2) \) work throughout the whole algorithm. Here is a smarter method, based on the history graph.

**Definition 7.2** Given \( R \subseteq P \) (regarded as a sequence that reflects the insertion order), the history graph of \( R \) is a directed acyclic graph whose vertices are all triangles that have ever been created during the incremental construction of \( DT(R) \). There is a directed edge from \( \Delta \) to \( \Delta' \) whenever \( \Delta \) has been destroyed during an insertion step, \( \Delta' \) has been created during the same insertion step, and \( \Delta \) overlaps with \( \Delta' \) in its interior.

It follows that the history graph contains triangles of outdegrees 3, 2 and 0. The ones of outdegree 0 are clearly the triangles of \( DT(R) \).

The triangles of outdegree 3 are the ones that have been destroyed during Step 1 of an insertion. For each such triangle \( \Delta \), its three outneighbors are the three new triangles that have replaced it, see Figure 7.4.

The triangles of outdegree 2 are the ones that have been destroyed during Step 2 of an insertion. For each such triangle \( \Delta \), its two outneighbors are the two new triangles created during the flip that has destroyed \( \Delta \), see Figure 7.5.

The history graph can be built during the incremental construction at asymptotically no extra cost; but it may need extra space since it keeps all triangles ever created. Given the history graph, we can search for the triangle \( \Delta \) of \( DT(R) \) that contains \( s \), as follows. We start from the big triangle spanned by the three far-away points; this one certainly
Contains $s$. Then we follow a directed path in the history graph. If the current triangle still has outneighbors, we find the unique outneighbor containing $s$ and continue the search with this neighbor. If the current triangle has no outneighbors anymore, it is in $\Delta T(R)$ and contains $s$—we are done.

**Types of triangles in the history graph.** After each insertion of a point $s$, several triangles are created and added to the history graph. It is important to note that these triangles come in two types: Some of them are valid Delaunay triangles of $R \cup \{s\}$, and they survive to the next stage of the incremental construction. Other triangles are immediately destroyed by subsequent Lawson flips, because they are not Delaunay triangles of $R \cup \{s\}$. These "ephemeral" triangles will give us some headache (though not much) in the algorithm’s analysis in the next chapter.

Note that, whenever a Lawson flip is performed, of the two triangles destroyed one of them is always a “valid” triangle from a previous iteration, and the other one is an “ephemeral” triangle that was created at this iteration. The ephemeral triangle is always the one that has $s$, the newly inserted point, as a vertex.

### 7.3 The structural change

Concerning the actual update (Steps 1 and 2), we can make the following

**Observation 7.3** *Given $\Delta T(R)$ and the triangle $\Delta$ of $\Delta T(R)$ that contains $s$, we can build $\Delta T(R \cup \{s\})$ in time proportional to the degree of $s$ in $\Delta T(R \cup \{s\})$, which is the number of triangles of $\Delta T(R \cup \{s\})$ containing $s$.***

Indeed, since every flip generates exactly one new triangle incident to $s$, the number of flips is the degree of $s$ minus three. Step 1 of the update takes constant time, and
since also every flip can be implemented in constant time, the observation follows.

In the next lecture, we will show that a clever insertion order guarantees that the
search paths traversed in the history graph are short, and that the structural change (the
number of new triangles) is small. This will then give us the $O(n \log n)$ algorithm.

**Exercise 7.4** For a sequence of $n$ pairwise distinct numbers $y_1, \ldots, y_n$ consider the se-
quence of pairs $(\min\{y_1, \ldots, y_i\}, \max\{y_1, \ldots, y_i\})_{i=0,1,\ldots,n}$ ($\min\emptyset := +\infty$, $\max\emptyset := -\infty$).
How often do these pairs change in expectation if the sequence is permuted ran-
domly, each permutation appearing with the same probability? Determine the ex-
pected value.

**Questions**

28. Describe the algorithm for the incremental construction of $\mathcal{DT}(P)$: how do we
find the triangle containing the point $s$ to be inserted into $\mathcal{DT}(R)$? How do we
transform $\mathcal{DT}(R)$ into $\mathcal{DT}(R \cup \{s\})$? How many steps does the latter transformation
take, in terms of $\mathcal{DT}(R \cup \{s\})$?

29. What are the two types of triangles that the history graph contains?
Chapter 8

The Configuration Space Framework

In Section 7.1, we have discussed the incremental construction of the Delaunay triangulation of a finite point set. In this lecture, we want to analyze the runtime of this algorithm if the insertion order is chosen uniformly at random among all insertion orders. We will do the analysis not directly for the problem of constructing the Delaunay triangulation but in a somewhat more abstract framework, with the goal of reusing the analysis for other problems.

Throughout this lecture, we again assume general position: no three points on a line, no four on a circle.

8.1 The Delaunay triangulation — an abstract view

The incremental construction constructs and destroys triangles. In this section, we want to take a closer look at these triangles, and we want to understand exactly when a triangle is “there”.

Lemma 8.1  Given three points \( p, q, r \in R \), the triangle \( \Delta(p, q, r) \) with vertices \( p, q, r \) is a triangle of \( \mathcal{D}(R) \) if and only if the circumcircle of \( \Delta(p, q, r) \) is empty of points from \( R \).

Proof.  The “only if” direction follows from the definition of a Delaunay triangulation (Definition 6.8). The “if” direction is a consequence of general position and Lemma 6.16: if the circumcircle \( C \) of \( \Delta(p, q, r) \) is empty of points from \( R \), then all the three edges \( \overline{pq}, \overline{qr}, \overline{pr} \) are easily seen to be in the Delaunay graph of \( R \). \( C \) being empty also implies that the triangle \( \Delta(p, q, r) \) is empty, and hence it forms a triangle of \( \mathcal{D}(R) \).

Next we develop a somewhat more abstract view of \( \mathcal{D}(R) \).

Definition 8.2

(i)  For all \( p, q, r \in P \), the triangle \( \Delta = \Delta(p, q, r) \) is called a configuration. The points \( p, q \) and \( r \) are called the defining elements of \( \Delta \).
(ii) A configuration $\Delta$ is in conflict with a point $s \in P$ if $s$ is strictly inside the circumcircle of $\Delta$. In this case, the pair $(\Delta, s)$ is called a conflict.

(iii) A configuration $\Delta$ is called active w.r.t. $R \subseteq P$ if (a) the defining elements of $\Delta$ are in $R$, and (b) if $\Delta$ is not in conflict with any element of $R$.

According to this definition and Lemma 8.1, $\mathcal{DT}(R)$ consists of exactly the configurations that are active w.r.t. $R$. Moreover, if we consider $\mathcal{DT}(R)$ and $\mathcal{DT}(R \cup \{s\})$ as sets of configurations, we can exactly say how these two sets differ.

There are the configurations in $\mathcal{DT}(R)$ that are not in conflict with $s$. These configurations are still in $\mathcal{DT}(R \cup \{s\})$. The configurations of $\mathcal{DT}(R)$ that are in conflict with $s$ will be removed when going from $R$ to $R \cup \{s\}$. Finally, $\mathcal{DT}(R \cup \{s\})$ contains some new configurations, all of which must have $s$ in their defining set. According to Lemma 8.1, it cannot happen that we get a new configuration without $s$ in its defining set, as such a configuration would have been present in $\mathcal{DT}(R)$ already.

8.2 Configuration Spaces

Here is the abstract framework that generalizes the previous configuration view of the Delaunay triangulation.

**Definition 8.3** Let $X$ (the ground set) and $\Pi$ (the set of configurations) be finite sets. Furthermore, let

$$D : \Pi \rightarrow 2^X$$

be a function that assigns to every configuration $\Delta$ a set of defining elements $D(\Delta)$. We assume that only a constant number of configurations have the same defining elements. Let

$$K : \Pi \rightarrow 2^X$$

be a function that assigns to every configuration $\Delta$ a set of elements in conflict with $\Delta$ (the “killer” elements). We stipulate that $D(\Delta) \cap K(\Delta) = \emptyset$ for all $\Delta \in \Pi$.

Then the quadruple $S = (X, \Pi, D, K)$ is called a configuration space. The number

$$d = d(S) := \max_{\Delta \in \Pi} |D(\Delta)|$$

is called the dimension of $S$.

Given $R \subseteq X$, a configuration $\Delta$ is called active w.r.t. $R$ if

$$D(\Delta) \subseteq R \quad \text{and} \quad K(\Delta) \cap R = \emptyset,$$

i.e. if all defining elements are in $R$ but no element of $R$ is in conflict with $\Delta$. The set of active configurations w.r.t. $R$ is denoted by $\mathcal{T}_S(R)$, where we drop the subscript if the configuration space is clear from the context.
In case of the Delaunay triangulation, we set $X = P$ (the input point set). $\Pi$ consists of all triangles $\Delta = \Delta(p, q, r)$ spanned by three points $p, q, r \in X \cup \{a, b, c\}$, where $a, b, c$ are the three artificial far-away points. We set $D(\Delta) := \{p, q, r\} \cap X$. The set $K(\Delta)$ consists of all points strictly inside the circumcircle of $\Delta$. The resulting configuration space has dimension 3, and the technical condition that only a constant number of configurations share the defining set is satisfied as well. In fact, every set of three points defines a unique configuration (triangle) in this case. A set of two points or one point defines three triangles (we have to add one or two artificial points which can be done in three ways). The empty set defines one triangle, the initial triangle consisting of just the three artificial points.

Furthermore, in the setting of the Delaunay triangulation, a configuration is active w.r.t. $R$ if it is in $\mathcal{D}(R \cup \{a, b, c\})$, i.e. we have $\mathcal{I}(R) = \mathcal{D}(R \cup \{a, b, c\})$.

### 8.3 Expected structural change

Let us fix a configuration space $S = (X, \Pi, D, K)$ for the remainder of this lecture. We can also interpret the incremental construction in $S$. Given $R \subseteq X$ and $s \in X \setminus R$, we want to update $\mathcal{I}(R)$ to $\mathcal{I}(R \cup \{s\})$. What is the number of new configurations that arise during this step? For the case of Delaunay triangulations, this is the relevant question when we want to bound the number of Lawson flips during one update step, since this number is exactly the number of new configurations minus three.

Here is the general picture.

**Definition 8.4** For $Q \subseteq X$ and $s \in Q$, $\deg(s, Q)$ is defined as the number of configurations of $\mathcal{I}(Q)$ that have $s$ in their defining set.

With this, we can say that the number of new configurations in going from $\mathcal{I}(R)$ to $\mathcal{I}(R \cup \{s\})$ is precisely $\deg(s, R \cup \{s\})$, since the new configurations are by definition exactly the ones that have $s$ in their defining set.

Now the random insertion order comes in for the first time: what is

$$E(\deg(s, R \cup \{s\})),$$

averaged over all insertion orders? In such a random insertion order, $R$ is a random $r$-element subset of $X$ (when we are about to insert the $(r+1)$-st element), and $s$ is a random element of $X \setminus R$. Let $\mathcal{T}_r$ be the “random variable” for the set of active configurations after $r$ insertion steps.

It seems hard to average over all $R$, but there is a trick: we make a movie of the randomized incremental construction, and then we watch the movie backwards. What we see is elements of $X$ being deleted one after another, again in random order. This is due to the fact that the reverse of a random order is also random. At the point where the $(r+1)$-st element is being deleted, it is going to be a random element $s$ of the currently
present \((r+1)\)-element subset \(Q\). For fixed \(Q\), the expected degree of \(s\) is simply the average degree of an element in \(Q\) which is

\[
\frac{1}{r+1} \sum_{s \in Q} \deg(s, Q) \leq \frac{d}{r+1}|\mathcal{T}(Q)|,
\]

since the sum counts every configuration of \(\mathcal{T}(Q)\) at most \(d\) times. Since \(Q\) is a random \((r+1)\)-element subset, we get

\[
E(\deg(s, R \cup \{s\})) \leq \frac{d}{r+1} t_{r+1},
\]

where \(t_{r+1}\) is defined as the expected number of active configurations w.r.t. a random \((r+1)\)-element set.

Here is a more formal derivation that does not use the backwards movie view. It exploits the bijection

\[(R, s) \mapsto (R \cup \{s\}, s)\]

between pairs \((R, s)\) with \(|R| = r\) and \(s \notin R\) and pairs \((Q, s)\) with \(|Q| = r+1\) and \(s \in Q\). Let \(n = |X|\).

\[
E(\deg(s, R \cup \{s\})) = \frac{1}{\binom{n}{r}} \sum_{R \subseteq X, |R| = r} \frac{1}{n-r} \sum_{s \in X \setminus R} \deg(s, R \cup \{s\})
\]

\[
= \frac{1}{\binom{n}{r}} \sum_{Q \subseteq X, |Q| = r+1} \frac{1}{n-r} \sum_{s \in Q} \deg(s, Q)
\]

\[
= \frac{1}{\binom{n}{r+1}} \sum_{Q \subseteq X, |Q| = r+1} \frac{\binom{n}{r+1}}{\binom{n}{r}} \frac{1}{n-r} \sum_{s \in Q} \deg(s, Q)
\]

\[
\leq \frac{1}{\binom{n}{r+1}} \sum_{Q \subseteq X, |Q| = r+1} \frac{d}{r+1} |\mathcal{T}(Q)|
\]

\[
= \frac{d}{r+1} t_{r+1}.
\]

Thus, the expected number of new configurations in going from \(\mathcal{T}_r\) to \(\mathcal{T}_{r+1}\) is bounded by

\[
\frac{d}{r+1} t_{r+1},
\]
where \( t_{r+1} \) is the expected size of \( T_{r+1} \).

What do we get for Delaunay triangulations? We have \( d = 3 \) and \( t_{r+1} \leq 2(r+4) - 4 \) (the maximum number of triangles in a triangulation of \( r + 4 \) points). Hence,

\[
E(\deg(s, R \cup \{s\})) \leq \frac{6r + 12}{r + 1} \approx 6.
\]

This means that on average, \( \approx 3 \) Lawson flips are done to update \( DJ_r \) (the Delaunay triangulation after \( r \) insertion steps) to \( DJ_{r+1} \). Over the whole algorithm, the expected update cost is thus \( O(n) \).

### 8.4 Bounding location costs by conflict counting

Before we can even update \( DJ_r \) to \( DJ_{r+1} \) during the incremental construction of the Delaunay triangulation, we need to locate the new point \( s \) in \( DJ_r \), meaning that we need to find the triangle that contains \( s \). We have done this with the history graph: During the insertion of \( s \) we "visit" a sequence of triangles from the history graph, each of which contains \( s \) and was created at some previous iteration \( k < r \).

However, some of these visited triangles are "ephemeral" triangles (recall the discussion at the end of Section 7.2), and they present a problem to the generic analysis we want to perform. Therefore, we will do a charging scheme, so that all triangles charged are valid Delaunay triangles.

The charging scheme is as follows: If the visited triangle \( \Delta \) is a valid Delaunay triangle (from some previous iteration), then we simply charge the visit of \( \Delta \) during the insertion of \( s \) to the triangle-point pair \( (\Delta, s) \).

If, on the other hand, \( \Delta \) is an "ephemeral" triangle, then \( \Delta \) was destroyed, together with some neighbor \( \Delta' \), by a Lawson flip into another pair \( \Delta'', \Delta''' \). Note that this neighbor \( \Delta' \) was a valid triangle. Thus, in this case we charge the visit of \( \Delta \) during the insertion of \( s \) to the pair \( (\Delta', s) \). Observe that \( s \) is contained in the circumcircle of \( \Delta' \), so \( s \) is in conflict with \( \Delta' \).

This way, we have charged each visit to a triangle in the history graph to a triangle-point pair of the form \( (\Delta, s) \), such that \( \Delta \) is in conflict with \( s \). Furthermore, it is easy to see that no such pair gets charged more than once.

We define the notion of a conflict in general:

**Definition 8.5** A conflict is a configuration-element pair \( (\Delta, s) \) where \( \Delta \in T_r \) for some \( r \) and \( s \in K(\Delta) \).

Thus, the running time of the Delaunay algorithm is proportional to the number of conflicts. We now proceed to derive a bound on the expected number of conflicts in the generic configuration-space framework.
8.5 Expected number of conflicts

Since every configuration involved in a conflict has been created in some step \( r \) (we include step 0), the total number of conflicts is

\[
\sum_{r=0}^{n} \sum_{\Delta \in \mathcal{I}_r \setminus \mathcal{I}_{r-1}} |K(\Delta)|,
\]

where \( \mathcal{I}_{-1} := \emptyset \). \( \mathcal{I}_0 \) consists of constantly many configurations only (namely those where the set of defining elements is the empty set), each of which is in conflict with at most all elements; moreover, no conflict is created in step \( n \). Hence,

\[
\sum_{r=0}^{n} \sum_{\Delta \in \mathcal{I}_r \setminus \mathcal{I}_{r-1}} |K(\Delta)| = O(n) + \sum_{r=1}^{n-1} \sum_{\Delta \in \mathcal{I}_r \setminus \mathcal{I}_{r-1}} |K(\Delta)|,
\]

and we will bound the latter quantity. Let

\[
K(r) := \sum_{\Delta \in \mathcal{I}_r \setminus \mathcal{I}_{r-1}} |K(\Delta)|, \quad r = 1, \ldots, n-1.
\]

and \( k(r) := E(K(r)) \) the expected number of conflicts created in step \( r \).

Bounding \( k(r) \). We know that \( \mathcal{I}_r \) arises from a random \( r \)-element set \( R \). Fixing \( R \), the backwards movie view tells us that \( \mathcal{I}_{r-1} \) arises from \( \mathcal{I}_r \) by deleting a random element \( s \) of \( R \). Thus,

\[
k(r) \leq \frac{1}{\binom{n}{r}} \sum_{R \subseteq X, |R| = r} \frac{1}{r} \sum_{s \in R} \sum_{\Delta \in \mathcal{I}_r \setminus \mathcal{I}_{r-1}(R \setminus \{s\})} |K(\Delta)|
\]

\[
= \frac{1}{\binom{n}{r}} \sum_{R \subseteq X, |R| = r} \frac{1}{r} \sum_{s \in R} \sum_{\Delta \in \mathcal{I}_r \setminus \mathcal{I}_{r-1}} |K(\Delta)|
\]

\[
\leq \frac{1}{\binom{n}{r}} \sum_{R \subseteq X, |R| = r} \frac{d}{r} \sum_{\Delta \in \mathcal{I}_r} |K(\Delta)|,
\]

since in the sum over \( s \in R \), every configuration is counted at most \( d \) times. Since we can rewrite

\[
\sum_{\Delta \in \mathcal{I}_r} |K(\Delta)| = \sum_{y \in X \setminus R} |\{\Delta \in \mathcal{I}_r : y \in K(\Delta)\}|,
\]

we thus have

\[
k(r) \leq \frac{1}{\binom{n}{r}} \sum_{R \subseteq X, |R| = r} \frac{d}{r} \sum_{y \in X \setminus R} |\{\Delta \in \mathcal{I}_r : y \in K(\Delta)\}|.
\]

To estimate this further, here is a simple but crucial
Lemma 8.6 The configurations in $\mathcal{T}(R)$ that are not in conflict with $y$ are the configurations in $\mathcal{T}(R \cup \{y\})$ that do not have $y$ in their defining set; in formulas:

$$|\mathcal{T}(R)| - |\{\Delta \in \mathcal{T}(R) : y \in K(\Delta)\}| = |\mathcal{T}(R \cup \{y\})| - \deg(y, R \cup \{y\}).$$

The proof is a direct consequence of the definitions: every configuration in $\mathcal{T}(R)$ not in conflict with $y$ is by definition still present in $\mathcal{T}(R \cup \{y\})$ and still does not have $y$ in its defining set. And a configuration in $\mathcal{T}(R \cup \{y\})$ with $y$ not in its defining set is by definition already present in $\mathcal{T}(R)$ and already there not in conflict with $y$.

The lemma implies that

$$k(r) \leq k_1(r) - k_2(r) + k_3(r),$$

where

$$k_1(r) = \frac{1}{\binom{n}{r}} \sum_{R \subseteq X, |R|=r} \frac{d}{r} \sum_{y \in X \setminus R} |\mathcal{T}(R)|,$$

$$k_2(r) = \frac{1}{\binom{n}{r}} \sum_{R \subseteq X, |R|=r} \frac{d}{r} \sum_{y \in X \setminus R} |\mathcal{T}(R \cup \{y\})|,$$

$$k_3(r) = \frac{1}{\binom{n}{r}} \sum_{R \subseteq X, |R|=r} \frac{d}{r} \sum_{y \in X \setminus R} \deg(y, R \cup \{y\}).$$

**Estimating $k_1(r)$.** This is really simple.

$$k_1(r) = \frac{1}{\binom{n}{r}} \sum_{R \subseteq X, |R|=r} \frac{d}{r} \sum_{y \in X \setminus R} |\mathcal{T}(R)|$$

$$= \frac{1}{\binom{n}{r}} \sum_{R \subseteq X, |R|=r} \frac{d}{r} (n - r) |\mathcal{T}(R)|$$

$$= \frac{d}{r} (n - r) t_r.$$
Estimating $k_2(r)$. For this, we need to employ our earlier $(R, y) \mapsto (R \cup \{y\}, y)$ bijection again.

\[
k_2(r) = \frac{1}{\binom{n}{r}} \sum_{R \subseteq X, |R| = r} \frac{d}{r} \sum_{y \in X \setminus R} |\mathcal{T}(R \cup \{y\})|
\]

\[
= \frac{1}{\binom{n}{r}} \sum_{Q \subseteq X, |Q| = r+1} \frac{d}{r} \sum_{y \in Q} |\mathcal{T}(Q)|
\]

\[
= \frac{1}{\binom{n}{r+1}} \sum_{Q \subseteq X, |Q| = r+1} \binom{n}{r+1} \frac{d}{r} \frac{n}{r+1} |\mathcal{T}(Q)|
\]

\[
= \frac{1}{\binom{n}{r+1}} \sum_{Q \subseteq X, |Q| = r+1} \frac{d}{r} (n-r) |\mathcal{T}(Q)|
\]

\[
= \frac{d}{r} (n-r) t_{r+1}
\]

\[
= \frac{d}{r+1} (n - (r + 1)) t_{r+1} + \frac{dn}{r(r+1)} t_{r+1}
\]

\[
= k_1(r + 1) + \frac{dn}{r(r+1)} t_{r+1}.
\]

Estimating $k_3(r)$. This is similar to $k_2(r)$ and in addition uses a fact that we have employed before: $\sum_{y \in Q} \deg(y, Q) \leq d |\mathcal{T}(Q)|$.

\[
k_3(r) = \frac{1}{\binom{n}{r}} \sum_{R \subseteq X, |R| = r} \frac{d}{r} \sum_{y \in X \setminus R} \deg(y, R \cup \{y\})
\]

\[
= \frac{1}{\binom{n}{r}} \sum_{Q \subseteq X, |Q| = r+1} \frac{d}{r} \sum_{y \in Q} \deg(y, Q)
\]

\[
\leq \frac{1}{\binom{n}{r}} \sum_{Q \subseteq X, |Q| = r+1} \frac{d^2}{r} |\mathcal{T}(Q)|
\]

\[
= \frac{1}{\binom{n}{r+1}} \sum_{Q \subseteq X, |Q| = r+1} \binom{n}{r+1} \frac{d^2}{r} \frac{n}{r+1} |\mathcal{T}(Q)|
\]

\[
= \frac{1}{\binom{n}{r+1}} \sum_{Q \subseteq X, |Q| = r+1} \frac{n-r}{r+1} \frac{d^2}{r} |\mathcal{T}(Q)|
\]

\[
= \frac{d^2}{r(r+1)} (n-r) t_{r+1}
\]

\[
= \frac{d^2n}{r(r+1)} t_{r+1} - \frac{d^2}{r+1} t_{r+1}.
\]
Summing up. Let us recapitulate: the overall expected number of conflicts is \(O(n)\) plus 
\[
\sum_{r=1}^{n-1} k(r) = \sum_{r=1}^{n-1} (k_1(r) - k_2(r) + k_3(r)).
\]

Using our previous estimates, \(k_1(2), \ldots, k_1(n-1)\) are canceled by the first terms of \(k_2(1), \ldots, k_2(n-2)\). The second term of \(k_2(r)\) can be combined with the first term of \(k_3(r)\), so that we get
\[
\sum_{r=1}^{n-1} (k_1(r) - k_2(r) + k_3(r)) \leq k_1(1) - k_1(n) + n \sum_{r=1}^{n-1} \frac{d(d-1)}{r(r+1)} t_{r+1} - \sum_{r=1}^{n-1} \frac{d^2}{r+1} t_{r+1}
\]
\[
\leq d(n-1)t_1 + d(d-1)n \sum_{r=1}^{n-1} \frac{t_{r+1}}{r(r+1)}
\]
\[
= O\left(d^2n \sum_{r=1}^{n} \frac{t_r}{r^2}\right).
\]

The Delaunay case. We have argued that the expected number of conflicts asymptotically bounds the expected total location cost over all insertion steps. The previous equation tells us that this cost is proportional to \(O(n)\) plus
\[
O\left(9n\sum_{r=1}^{n} \frac{2(r+4) - 4}{r^2}\right) = O\left(n\sum_{r=1}^{n} \frac{1}{r}\right) = O(n \log n).
\]

Here,
\[
\sum_{r=1}^{n} \frac{1}{r} =: H_n
\]
is the \(n\)-th Harmonic Number which is known to be approximately \(\ln n\).

By going through the abstract framework of configuration spaces, we have thus analyzed the randomized incremental construction of the Delaunay triangulation of \(n\) points. According to Section 8.3, the expected update cost itself is only \(O(n)\). The steps dominating the runtime are the location steps via the history graph. According to Section 8.5, all history graph searches (whose number is proportional to the number of conflicts) can be performed in expected time \(O(n \log n)\), and this then also bounds the space requirements of the algorithm.

Exercise 8.7 Design and analyze a sorting algorithm based on randomized incremental construction in configuration spaces. The input is a set \(S\) of numbers, and the output should be the sorted sequence (in increasing order).
a) Define an appropriate configuration space for the problem! In particular, the set of active configurations w.r.t. \( S \) should represent the desired sorted sequence.

b) Provide an efficient implementation of the incremental construction algorithm. "Efficient" means that the runtime of the algorithm is asymptotically dominated by the number of conflicts.

c) What is the expected number of conflicts (and thus the asymptotic runtime of your sorting algorithm) for a set \( S \) of \( n \) numbers?

Questions

30. What is a configuration space? Give a precise definition! What is an active configuration?

31. How do we get a configuration space from the problem of computing the Delaunay triangulation of a finite point set?

32. How many new active configurations do we get on average when inserting the \( r \)-th element? Provide an answer for configuration spaces in general, and for the special case of the Delaunay triangulation.

33. What is a conflict? Provide an answer for configuration spaces in general, and for the special case of the Delaunay triangulation.

34. Explain why counting the expected number of conflicts asymptotically bounds the cost for the history searches during the randomized incremental construction of the Delaunay triangulation!
Chapter 9

Voronoi Diagrams

9.1 Post Office Problem

Suppose there are \(n\) post offices \(p_1, \ldots, p_n\) in a city. Someone who is located at a position \(q\) within the city would like to know which post office is closest to him. Modeling the city as a planar region, we think of \(p_1, \ldots, p_n\) and \(q\) as points in the plane. Denote the set of post offices by \(P = \{p_1, \ldots, p_n\}\).

While the locations of post offices are known and do not change so frequently, we do not know in advance for which—possibly many—query locations the closest post office is to be found. Therefore, our long term goal is to come up with a data structure on top of \(P\) that allows to answer any possible query efficiently. The basic idea is to apply a so-called \textit{locus approach}: we partition the query space into regions on which is the answer is the same. In our case, this amounts to partition the plane into regions such that for all points within a region the same point from \(P\) is closest (among all points from \(P\)).

As a warmup, consider the problem for two post offices \(p_i, p_j \in P\). For which query locations is the answer \(p_i\) rather than \(p_j\)? This region is bounded by the bisector of \(p_i\) and \(p_j\), that is, the set of points which have the same distance to both points.
Proposition 9.1 For any two distinct points in $\mathbb{R}^d$ the bisector is a hyperplane, that is, in $\mathbb{R}^2$ it is a line.

Proof. Let $p = (p_1, \ldots, p_d)$ and $q = (q_1, \ldots, q_d)$ be two points in $\mathbb{R}^d$. The bisector of $p$ and $q$ consists of those points $x = (x_1, \ldots, x_d)$ for which
\[ ||p - x|| = ||q - x|| \iff ||p - x||^2 = ||q - x||^2 \iff ||p||^2 - ||q||^2 = 2(p - q)^\top x. \]
As $p$ and $q$ are distinct, this is the equation of a hyperplane. □

Denote by $H(p_i, p_j)$ the closed halfspace bounded by the bisector of $p_i$ and $p_j$ that contains $p_i$. In $\mathbb{R}^2$, the region $H(p_i, p_j)$ is a halfplane; see Figure 9.2.

Exercise 9.2

a) What is the bisector of a line $\ell$ and a point $p \in \mathbb{R}^2 \setminus \ell$, that is, the set of all points $x \in \mathbb{R}^2$ with $||x - p|| = ||x - \ell||$ (=$\min_{q \in \ell} ||x - q||$)?

b) For two points $p \neq q \in \mathbb{R}^2$, what is the region that contains all points whose distance to $p$ is exactly twice their distance to $q$?

9.2 Voronoi Diagram

In the following we work with a set $P = \{p_1, \ldots, p_n\}$ of points in $\mathbb{R}^2$.

Definition 9.3 (Voronoi cell) For $p_i \in P$ denote the Voronoi cell $V_P(i)$ of $p_i$ by
\[ V_P(i) := \{ q \in \mathbb{R}^2 \mid ||q - p_i|| \leq ||q - p|| \mbox{ for all } p \in P \}. \]

Proposition 9.4
\[ V_P(i) = \bigcap_{j \neq i} H(p_i, p_j). \]

Proof. For $j \neq i$ we have $||q - p_i|| \leq ||q - p_j|| \iff q \in H(p_i, p_j)$. □
Corollary 9.5 $V_P(i)$ is non-empty and convex.

Proof. According to Proposition 9.4, the region $V_P(i)$ is the intersection of a finite number of halfplanes and hence convex. As $p_i \in V_P(i)$, we have $V_P(i) \neq \emptyset$. 

Observe that every point of the plane lies in some Voronoi cell but no point lies in the interior of two Voronoi cells. Therefore these cells form a subdivision of the plane. See Figure 9.3 for an example.

Definition 9.6 (Voronoi Diagram) The Voronoi Diagram $VD(P)$ of a set $P = \{p_1, \ldots, p_n\}$ of points in $\mathbb{R}^2$ is the subdivision of the plane induced by the Voronoi cells $V_P(i)$, for $i = 1, \ldots, n$.

Denote by $VV(P)$ the set of vertices, by $VE(P)$ the set of edges, and by $VR(P)$ the set of regions (faces) of $VD(P)$.

![Figure 9.3: Example: The Voronoi diagram of a point set.](image)

Lemma 9.7 For every vertex $v \in VV(P)$ the following statements hold.

a) $v$ is the common intersection of at least three edges from $VE(P)$;

b) $v$ is incident to at least three regions from $VR(P)$;

c) $v$ is the center of a circle $C(v)$ through at least three points from $P$ such that

d) $C(v)^{\circ} \cap P = \emptyset$.

Proof. Consider a vertex $v \in VV(P)$. As all Voronoi cells are convex, $k \geq 3$ of them must be incident to $v$. This proves Part a) and b).
Without loss of generality let these cells be $V_P(i)$, for $1 \leq i \leq k$. Denote by $e_i$, $1 \leq i \leq k$, the edge incident to $v$ that bounds $V_P(i)$ and $V_P((i \text{mod } k) + 1)$.

For any $i = 1, \ldots, k$ we have $v \in e_i \Rightarrow ||v - p_i|| = ||v - p_{(i \text{mod } k) + 1}||$. In other words, $p_1, p_2, \ldots, p_k$ are cocircular, which proves Part c).

Part d): Suppose there exists a point $p_i \in C(v)^\circ$. Then the vertex $v$ is closer to $p_i$ than it is to any of $p_1, \ldots, p_k$, in contradiction to the fact that $v$ is contained in all of $V_P(1), \ldots, V_P(k)$. □

**Corollary 9.8** If $P$ is in general position (no four points from $P$ are cocircular), then for every vertex $v \in \text{VV}(P)$ the following statements hold.

a) $v$ is the common intersection of exactly three edges from $\text{VE}(P)$;

b) $v$ is incident to exactly three regions from $\text{VR}(P)$;

c) $v$ is the center of a circle $C(v)$ through exactly three points from $P$ such that

d) $C(v)^\circ \cap P = \emptyset$. □

**Lemma 9.9** There is an unbounded Voronoi edge bounding $V_P(i)$ and $V_P(j) \iff \overline{p_i p_j} \cap P = \{p_i, p_j\}$ and $\overline{p_i p_j} \subseteq \partial\text{conv}(P)$, where the latter denotes the boundary of the convex hull of $P$.

**Proof.**
Denote by $b_{i,j}$ the bisector of $p_i$ and $p_j$, and let $C$ denote the family of circles centered at some point on $b_{i,j}$ and passing through $p_i$ (and $p_j$). There is an unbounded Voronoi edge bounding $V_P(i)$ and $V_P(j) \iff$ there is a ray $\rho \subset b_{i,j}$ such that $||r - p_k|| > ||r - p_i|| (= ||r - p_j||)$, for every $r \in \rho$ and every $p_k \in P$ with $k \notin \{i, j\}$. Equivalently, there is a ray $\rho \subset b_{i,j}$ such that for every point $r \in \rho$ the circle $C \in C$ centered at $r$ does not contain any point from $P$ in its interior.

The latter statement implies that the open halfplane $H$, whose bounding line passes through $p_i$ and $p_j$ and such that $H$ contains the infinite part of $\rho$, contains no point from $P$ in its interior. Therefore, $\overline{p_i p_j}$ appears on $\partial\text{conv}(P)$ and $\overline{p_i p_j}$ does not contain any $p_k \in P$, for $k \neq i, j$.

Conversely, suppose that $\overline{p_i p_j}$ appears on $\partial\text{conv}(P)$ and $\overline{p_i p_j} \cap P = \{p_i, p_j\}$. Then some halfplane $H$ whose bounding line passes through $p_i$ and $p_j$ contains no point from
P in its interior. In particular, the existence of H together with \( \overline{p_i p_j} \cap P = \{p_i, p_j\} \) implies that there is some circle \( C, C \subseteq H \) such that \( C \cap P = \{p_i, p_j\} \). Denote by \( r_0 \) the center of \( C \) and let \( \rho \) denote the ray starting from \( r_0 \) along \( b_{i,j} \) such that the infinite part of \( \rho \) is contained in \( H \). Consider any circle \( D, D \cap \overline{C} = \{p_i, p_j\} \) such that the infinite part of \( \rho \) is contained in \( H \). As neither \( H \) nor \( C \) contain any point from \( P \) in their respective interior, neither does \( D \). This holds for every \( D \), and we have seen above that this statement is equivalent to the existence of an unbounded Voronoi edge bounding \( V_P(i) \) and \( V_P(j) \).

9.3 Duality

A \textit{straight-line dual} of a plane graph \( G \) is a graph \( G' \) defined as follows: Choose a point for each face of \( G \) and connect any two such points by a straight edge, if the corresponding faces share an edge of \( G \). Observe that this notion depends on the embedding; that is why the straight-line dual is defined for a plane graph rather than for an abstract graph. In general, \( G' \) may have edge crossings, which may also depend on the choice of representative points within the faces. However, for Voronoi diagrams is a particularly natural choice of representative points such that \( G' \) is plane: the points from \( P \).

**Theorem 9.10 (Delaunay [2])** The straight-line dual of \( VD(P) \) for a set \( P \subset \mathbb{R}^2 \) of \( n \geq 3 \) points in general position (no three points from \( P \) are collinear and no four points from \( P \) are cocircular) is a triangulation: the unique Delaunay triangulation of \( P \).

**Proof.** By Lemma \[9.9\] the convex hull edges appear in the straight-line dual \( T \) of \( VD(P) \) and they correspond exactly to the unbounded edges of \( VD(P) \). All remaining edges of \( VD(P) \) are bounded, that is, both endpoints are Voronoi vertices. Consider some \( v \in VV(P) \). According to Corollary \[9.8(b)\], \( v \) is incident to exactly three Voronoi regions, which, therefore, form a triangle \( \triangle(v) \) in \( T \). By Corollary \[9.8(d)\], the circumcircle of \( \triangle(v) \) does not contain any point from \( P \) in its interior. Hence \( \triangle(v) \) appears in the (unique by Corollary \[6.17\]) Delaunay triangulation of \( P \).

Conversely, for any triangle \( p_ip_jp_k \) in the Delaunay triangulation of \( P \), by the empty circle property the circumcenter \( c \) of \( p_ip_jp_k \) has \( p_i, p_j, \) and \( p_k \) as its closest points from \( P \). Therefore, \( c \in VV(P) \) and—as above—the triangle \( p_ip_jp_k \) appears in \( T \). \( \square \)

It is not hard to generalize Theorem \[9.10\] to general point sets. In this case, a Voronoi vertex of degree \( k \) is mapped to a convex polygon with \( k \) cocircular vertices. Any triangulation of such a polygon yields a Delaunay triangulation of the point set.

**Corollary 9.11** \(|VE(P)| \leq 3n - 6 \) and \(|VV(P)| \leq 2n - 5\).

**Proof.** Every edge in \( VE(P) \) corresponds to an edge in the dual Delaunay triangulation. The latter is a plane graph on \( n \) vertices and thus has at most \( 3n - 6 \) edges and at most \( 2n - 4 \) faces by Lemma \[4.1\]. Only the bounded faces correspond to a vertex in \( VD(P) \). \( \square \)
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Figure 9.4: The Voronoi diagram of a point set and its dual Delaunay triangulation.

Corollary 9.12 For a set $P \subset \mathbb{R}^2$ of $n$ points, the Voronoi diagram of $P$ can be constructed in expected $O(n \log n)$ time and $O(n)$ space.

Proof. We have seen that a Delaunay triangulation $T$ for $P$ can be obtained using randomized incremental construction in the given time and space bounds. As $T$ is a plane graph, its number of vertices, edges, and faces all are linear in $n$. Therefore, the straight-line dual of $T$—which by Theorem 9.10 is the desired Voronoi diagram—can be computed in $O(n)$ additional time and space.

Exercise 9.13 Consider the Delaunay triangulation $T$ for a set $P \subset \mathbb{R}^2$ of $n \geq 3$ points in general position. Prove or disprove:

a) Every edge of $T$ intersects its dual Voronoi edge.

b) Every vertex of $VD(P)$ is contained in its dual Delaunay triangle.

9.4 Lifting Map

Recall the lifting map that we used in Section 6.3 to prove that the Lawson Flip Algorithm terminates. Denote by $\mathbb{U} : z = x^2 + y^2$ the unit paraboloid in $\mathbb{R}^3$. The lifting map $\ell : \mathbb{R}^2 \to \mathbb{U}$ with $\ell : p = (p_x, p_y) \mapsto (p_x, p_y, p_x^2 + p_y^2)$ is the projection of the $x/y$-plane onto $\mathbb{U}$ in direction of the $z$-axis.

For $p \in \mathbb{R}^2$ let $H_p$ denote the plane of tangency to $\mathbb{U}$ in $\ell(p)$. Denote by $h_p : \mathbb{R}^2 \to H_p$ the projection of the $x/y$-plane onto $H_p$ in direction of the $z$-axis (see Figure 9.5).

Lemma 9.14 $||\ell(q) - h_p(q)|| = ||p - q||^2$, for any points $p, q \in \mathbb{R}^2$. 

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Exercise 9.15  Prove Lemma 9.14.  Hint: First determine the equation of the tangent plane $H_p$ to $U$ in $\ell(p)$.

Theorem 9.16  For $p = (p_x, p_y) \in \mathbb{R}^2$ denote by $H_p$ the plane of tangency to the unit paraboloid $U = \{(x, y, z) : z = x^2 + y^2\} \subset \mathbb{R}^3$ in $\ell(p) = (p_x, p_y, p_x^2 + p_y^2)$. Let $\mathcal{H}(P) := \bigcap_{p \in P} H_p^+$ the intersection of all halfspaces above the planes $H_p$, for $p \in P$. Then the vertical projection of $\partial \mathcal{H}(P)$ onto the $x/y$-plane forms the Voronoi Diagram of $P$ (the faces of $\partial \mathcal{H}(P)$ correspond to Voronoi regions, the edges to Voronoi edges, and the vertices to Voronoi vertices).

Proof.  For any point $q \in \mathbb{R}^2$, the vertical line through $q$ intersects every plane $H_p$, $p \in P$. By Lemma 9.14 the topmost plane intersected belongs to the point from $P$ that is closest to $q$.  \qed

9.5 Point location in a Voronoi Diagram

One last bit is still missing in order to solve the post office problem optimally.

Theorem 9.17  Given a triangulation $T$ for a set $P \subset \mathbb{R}^2$ of $n$ points, one can build in $O(n)$ time an $O(n)$ size data structure that allows for any query point $q \in \text{conv}(P)$ to find in $O(\log n)$ time a triangle from $T$ containing $q$.

The data structure we will employ is known as Kirkpatrick's hierarchy. But before discussing it in detail, let us put things together in terms of the post office problem.

Corollary 9.18 (Nearest Neighbor Search)  Given a set $P \subset \mathbb{R}^2$ of $n$ points, one can build in expected $O(n \log n)$ time an $O(n)$ size data structure that allows for any query
point q ∈ conv(P) to find in O(\log n) time a nearest neighbor of q among the points from P.

**Proof.** First construct the Voronoi Diagram V of P in expected O(n \log n) time. It has exactly n convex faces. Every unbounded face can be cut by the convex hull boundary into a bounded and an unbounded part. As we are concerned with query points within conv(P) only, we can restrict our attention to the bounded parts.\(^1\) Any convex polygon can easily be triangulated in time linear in its number of edges (= number of vertices). As V has at most 3n − 6 edges and every edge appears in exactly two faces, V can be triangulated in O(n) time overall. Label each of the resulting triangles with the point from p, whose Voronoi region contains it, and apply the data structure from Theorem 9.17.

### 9.5.1 Kirkpatrick’s Hierarchy

We will now develop the data structure for point location in a triangulation, as described in Theorem 9.17. The main idea is to construct a hierarchy \(T_0, \ldots, T_h\) of triangulations, such that

- \(T_0 = T\),
- the vertices of \(T_i\) are a subset of the vertices of \(T_{i-1}\), for \(i = 1, \ldots, h\), and
- \(T_h\) comprises a single triangle only.

**Search.** For a query point x the triangle from T containing x can be found as follows.

**Search**\((x \in \mathbb{R}^2)\)

1. For \(i = h, h - 1, \ldots, 0\): Find a triangle \(t_i\) from \(T_i\) that contains \(x\).
2. return \(t_0\).

This search is efficient under the following conditions.

(C1) Every triangle from \(T_i\) intersects only few \((\leq c)\) triangles from \(T_{i-1}\). (These will then be connected via the data structure.)

(C2) \(h\) is small \((\leq d \log n)\).

**Proposition 9.19** The search procedure described above needs \(\leq 3cd \log n = O(\log n)\) orientation tests.

**Proof.** For every \(T_i, 0 \leq i < h\), at most c triangles are tested as to whether or not they contain \(x\). Using three orientation tests one can determine whether or not a triangle contains a given point. \(\square\)

\(^1\)We even know how to decide in \(O(\log n)\) time whether or not a given point lies within conv(P), see Exercise 3.22.
Thinning. Removing a vertex \( v \) and all its incident edges from a triangulation creates a non-triangulated hole that forms a star-shaped polygon since all points are visible from \( v \) (the star-point).

**Lemma 9.20** A star-shaped polygon, given as a sequence of \( n \geq 3 \) vertices and a star-point, can be triangulated in \( O(n) \) time.

**Proof.** For \( n = 3 \) there is nothing to do. For \( n > 3 \), consider a star-shaped polygon \( P = (p_0, \ldots, p_{n-1}) \) and a star-point \( s \) of \( P \). Consider some convex vertex \( p_i \) of \( P \), that is, \( \angle p_{i+1}p_ip_{i-1} \leq \pi \), all indices taken mod \( n \). (Every simple polygon on \( n \geq 3 \) vertices has at least three convex vertices, on the convex hull.)

As \( P \) is star-shaped, the quadrilateral \( Q = p_{i-1}p_ip_{i+1}s \) is completely contained in \( P \). Therefore, if \( \angle p_{i-1}sp_{i+1} \leq \pi \) and hence \( Q \) is convex (Figure 9.6a), then we can add \( p_{i-1}p_{i+1} \) as a diagonal. In this way one triangle is cut off from \( P \), leaving a star-shaped polygon \( P' \) (with respect to \( s \)) on \( n-1 \) vertices. The polygon \( P' \) can then be triangulated recursively. If, on the other hand, \( \angle p_{i-1}sp_{i+1} > \pi \) (Figure 9.6b), we cannot safely add the edge \( p_{i-1}p_{i+1} \) because it might intersect other edges of \( P \) or even lie outside of \( P \). But we claim that in this case there exists another convex vertex \( p_j \) of \( P \), for which \( \angle p_{j-1}sp_{j+1} \leq \pi \) and therefore we can add the edge \( p_{j-1}p_{j+1} \) instead.

In fact, it is enough to choose \( p_j \) to be some convex vertex of \( P \) that is not a neighbor of \( p_i \): As \( \sum_{k=0}^{n-1} \angle p_{k-1}sp_k = 2\pi \) and \( \angle p_{i-1}sp_i + \angle p_isp_{i+1} = \angle p_{i-1}sp_{i+1} > \pi \), we have \( \angle p_{j-1}sp_j + \angle p_jsp_{j+1} = \angle p_{j-1}sp_{j+1} < \pi \).

It remains to show that such a vertex \( p_j \) exists. \( P \) has at least three convex vertices. One of them is \( p_i \). If the only other two convex vertices are \( p_{i-1} \) and \( p_{i+1} \), then we make the whole argument with \( p_{i-1} \) instead of \( p_i \) and find \( j = i + 1 \). Note that \( p_{i-1} \) and \( p_{i+1} \) are not neighbors because \( P \) is not a triangle.

As for the linear time bound, we simply scan the sequence of vertices as in Graham’s Scan. For every triple \( p_{i-1}p_ip_{i+1} \) of successive vertices it is tested (in constant time) whether \( p_{i-1}p_ip_{i+1}s \) forms a convex quadrilateral. If so, then the edge \( p_{i-1}p_{i+1} \) is added, effectively removing \( p_i \) from further consideration. Therefore, \( p_i \) can be charged for the (potential, if there are enough vertices left) additional test of the new triple \( p_xp_{i-1}p_{i+1} \) formed with the predecessor \( p_x \) of \( p_{i-1} \) in the current sequence. As shown for Graham’s Scan in Theorem 3.25, this results in a linear time algorithm overall.\(^2\)

\(^2\)Recall that the \( O(n \log n) \) time bound for Graham’s Scan was caused by the initial sorting only.
As a side remark, the *kernel* of a simple polygon, that is, the set of all star-points, can be constructed in linear time as well. We will get back to this question later in the course.

Our working plan is to obtain $T_i$ from $T_{i-1}$ by removing several *independent* (pairwise non-adjacent) vertices and re-triangulating. These vertices should

a) have small degree (otherwise the degree within the hierarchy gets too large, that is, we need to test too many triangles on the next level) and

b) be many (otherwise the height $h$ of the hierarchy gets too large).

The following lemma asserts the existence of a sufficiently large set of independent small-degree vertices in every triangulation.

**Lemma 9.21** In every triangulation of $n$ points in $\mathbb{R}^2$ there exists an independent set of at least $\lceil n/18 \rceil$ vertices of maximum degree 8. Moreover, such a set can be found in $O(n)$ time.

**Proof.** Let $T = (V, E)$ denote the graph of the triangulation, which we consider as an abstract graph in the following. We may suppose that $T$ is maximally planar, that is, all faces are triangles. (Otherwise triangulate the exterior face arbitrarily. An independent set in the resulting graph is also independent in $T$.) For $n = 3$ the statement is true. Let $n \geq 4$.

By the Euler formula we have $|E| = 3n - 6$, that is,

$$\sum_{v \in V} \deg_T(v) = 2|E| = 6n - 12 < 6n.$$  

Let $W \subseteq V$ denote the set of vertices of degree at most 8. Claim: $|W| > n/2$. Suppose $|W| \leq n/2$. As every vertex has degree at least three, we have

$$\sum_{v \in V} \deg_T(v) = \sum_{v \in W} \deg_T(v) + \sum_{v \in V \setminus W} \deg_T(v) \geq 3|W| + 9|V \setminus W|$$

$$= 3|W| + 9(n - |W|) = 9n - 6|W| \geq 9n - 3n = 6n,$$

in contradiction to the above.

Construct an independent set $U$ in $T$ as follows (greedily): As long as $W \neq \emptyset$, add an arbitrary vertex $v \in W$ to $U$ and remove $v$ and all its neighbors from $W$.

Obviously $U$ is independent and all vertices in $U$ have degree at most 8. At each selection step at most 9 vertices are removed from $W$. Therefore $|U| \geq \lceil (n/2)/9 \rceil = \lceil n/18 \rceil$.

**Proof.** (of Theorem 9.17)

Construct the hierarchy $T_0, \ldots, T_h$ with $T_0 = T$ as follows. Obtain $T_i$ from $T_{i-1}$ by removing an independent set $U$ as in Lemma 9.21 and re-triangulating the resulting holes. By
Lemma 9.20 and Lemma 9.21 every step is linear in the number $|T_i|$ of vertices in $T_i$. The total cost for building the data structure is thus

$$\sum_{i=0}^{h} \alpha |T_i| \leq \sum_{i=0}^{h} \alpha n(17/18)^i < \alpha n \sum_{i=0}^{\infty} (17/18)^i = 18\alpha n \in O(n),$$

for some constant $\alpha$. Similarly the space consumption is linear.

The number of levels amounts to $h = \log_{18/17} n < 12.2\log n$. Thus by Proposition 9.19 the search needs at most $3 \cdot 8 \cdot 12.2 \cdot 18.n = 292.7 \log n$ orientation tests. \hfill \square

**Improvements.** As the name suggests, the hierarchical approach discussed above is due to David Kirkpatrick [5]. The constant 292 that appears in the search time is somewhat large. There has been a whole line of research trying to improve it using different techniques.

- Sarnak and Tarjan [6]: $4 \log n$.
- Edelsbrunner, Guibas, and Stolfi [3]: $3 \log n$.
- Goodrich, Orletsky, and Ramaiyer [4]: $2 \log n$.
- Adamy and Seidel [1]: $1 \log n + 2\sqrt{\log n} + O(\sqrt{\log n})$.

**Exercise 9.22** Let $\{p_1, p_2, \ldots, p_n\}$ be a set of points in the plane, which we call obstacles. Imagine there is a disk of radius $r$ centered at the origin which can be moved around the obstacles but is not allowed to intersect them (touching the boundary is ok). Is it possible to move the disk out of these obstacles? See the example depicted in Figure 9.7 below.

*More formally, the question is whether there is a (continuous) path $\gamma : [0,1] \to \mathbb{R}^2$ with $\gamma(0) = (0,0)$ and $\|\gamma(1)\| \geq \max\{\|p_1\|, \ldots, \|p_n\|\}$, such that at any time $t \in [0,1]$ and $\|\gamma(t) - p_i\| \geq r$, for any $1 \leq i \leq n$. Describe an algorithm to decide this question and to construct such a path—if one exists—given arbitrary points $\{p_1, p_2, \ldots, p_n\}$ and a radius $r > 0$. Argue why your algorithm is correct and analyze its running time.*

**Exercise 9.23** This exercise is about an application from Computational Biology:

You are given a set of disks $P = \{a_1, \ldots, a_n\}$ in $\mathbb{R}^2$, all with the same radius $r_a > 0$. Each of these disks represents an atom of a protein. A water molecule is represented by a disc with radius $r_w > r_a$. A water molecule cannot intersect the interior of any protein atom, but it can be tangent to one. We say that an atom $a_i \in P$ is accessible if there exists a placement of a water molecule such that it is tangent to $a_i$ and does not intersect the interior of any other atom in $P$. Given $P$, find an $O(n \log n)$ time algorithm which determines all atoms of $P$ that are inaccessible.
Exercise 9.24 Let $P \subset \mathbb{R}^2$ be a set of $n$ points. Describe a data structure to find in $O(\log n)$ time a point in $P$ that is furthest from a given query point $q$ among all points in $P$.

Exercise 9.25 Show that the bounds given in Theorem 9.17 are optimal in the algebraic computation tree model.

Questions

35. **What is the Voronoi diagram of a set of points in $\mathbb{R}^2$?** Give a precise definition and explain/prove the basic properties: convexity of cells, why is it a subdivision of the plane?, Lemma 9.7, Lemma 9.9.

36. **What is the correspondence between the Voronoi diagram and the Delaunay triangulation for a set of points in $\mathbb{R}^2$?** Prove duality (Theorem 9.10) and explain where general position is needed.

37. **How to construct the Voronoi diagram of a set of points in $\mathbb{R}^2$?** Describe an $O(n \log n)$ time algorithm, for instance, via Delaunay triangulation.

38. **How can the Voronoi diagram be interpreted in context of the lifting map?** Describe the transformation and prove its properties to obtain a formulation of the Voronoi diagram as an intersection of halfspaces one dimension higher.

39. **What is the Post-Office Problem and how can it be solved optimally?** Describe the problem and a solution using linear space, $O(n \log n)$ preprocessing, and $O(\log n)$ query time.
40. How does Kirkpatrick’s hierarchical data structure for planar point location work exactly? Describe how to build it and how the search works, and prove the runtime bounds. In particular, you should be able to state and prove Lemma 9.20, Lemma 9.21 and Theorem 9.17.

References


Chapter 10

Trapezoidal Maps

In this section, we will see another application of randomized incremental construction in the abstract configuration space framework. At the same time, this will give us an efficient algorithm for solving the general problem of point location, as well as a faster algorithm for computing all intersections between a given set of line segments.

10.1 The Trapezoidal Map

To start with, let us introduce the concept of a trapezoidal map. We are given a set \( S = \{s_1, \ldots, s_n\} \) of line segments in the plane (not necessarily disjoint). We make several general position assumptions. We assume that no two segment endpoints and intersection points have the same \( x \)-coordinate. As an exception, we do allow several segments to share an endpoint. We also assume that no line segment is vertical, that any two line segments intersect in at most one point (which is a common endpoint, or a proper crossing), and that no three line segments have a common point. Finally, we assume that \( s_i \subseteq [0,1]^2 \) for all \( i \) (which can be achieved by scaling the coordinates of the segments accordingly).

**Definition 10.1** The trapezoidal map of \( S \) is the partition of \([0,1]^2\) into vertices, edges, and faces (called trapezoids), obtained as follows. Every segment endpoint and point of intersection between segments gets connected by two vertical extensions with the next feature below and above, where a feature is either another line segment or an edge of the bounding box \([0,1]^2\).

Figure [10.1] gives an example.

(The general-position assumptions are made only for convenience and simplicity of the presentation. The various degeneracies can be handled without too much trouble, though we will not get into the details.)

The trapezoids of the trapezoidal map are “true” trapezoids (quadrangles with two parallel vertical sides), and triangles (which may be considered as degenerate trapezoids).
10.2 Applications of trapezoidal maps

In this chapter we will see two applications of trapezoidal maps (there are others):

1. **Point location**: Given a set of $n$ segments in the plane, we want to preprocess them in order to answer point-location queries: given a point $p$, return the cell (connected component of the complement of the segments) that contains $p$ (see Figure 10.2). This is a more powerful alternative to Kirkpatrick’s algorithm that handles only triangulations, and which is treated in Section 9.5. The preprocessing constructs the trapezoidal map of the segments (Figure 10.3) in expected time $O(n \log n + K)$, where $K$ is the number of intersections between the input segments; the query time will be $O(\log n)$ in expectation.

2. **Line segment intersection**: Given a set of $n$ segments, we will report all segment intersections in expected time $O(n \log n + K)$. This is a faster alternative to the line-sweep algorithm we saw in Section 5.2, which takes time $O((n + K) \log n)$.

10.3 Incremental Construction of the Trapezoidal Map

We can construct the trapezoidal map by inserting the segments one by one, in random order, always maintaining the trapezoidal map of the segments inserted so far. In order to perform manipulations efficiently, we can represent the current trapezoidal map as a doubly-connected edge list (see Section 4.2), for example.
10.3. Incremental Construction of the Trapezoidal Map

Figure 10.2: The general point location problem defined by a set of (possibly intersecting) line segments. In this example, the segments partition the plane into 5 cells.

Suppose that we have already inserted segments $s_1, \ldots, s_{r-1}$, and that the resulting trapezoidal map $\mathcal{T}_{r-1}$ looks like in Figure 10.1. Now we insert segment $s_r$ (see Figure 10.4).

Here are the four steps that we need to do in order to construct $\mathcal{T}_r$.

1. **Find** the trapezoid $\Box_0$ of $\mathcal{T}_{r-1}$ that contains the left endpoint of $s_r$.

2. **Trace** $s_r$ through $\mathcal{T}_{r-1}$ until the trapezoid containing the right endpoint of $s_r$ is found. To get from the current trapezoid $\Box$ to the next one, traverse the boundary of $\Box$ until the edge is found through which $s_r$ leaves $\Box$.

3. **Split** the trapezoids intersected by $s_r$. A trapezoid $\Box$ may get replaced by

   - two new trapezoids (if $s_r$ intersects two vertical extensions of $\Box$);
   - three new trapezoids (if $s_r$ intersects one vertical extension of $\Box$);
   - four new trapezoids (if $s_r$ intersects no vertical extension of $\Box$).

4. **Merge** trapezoids by removing parts of vertical extensions that do not belong to $\mathcal{T}_r$ anymore.

Figure 10.5 illustrates the **Trace** and **Split** steps. $s_6$ intersects 5 trapezoids, and they are being split into 3, 3, 4, 3, and 3 trapezoids, respectively.
Figure 10.3: The trapezoidal map is a refinement of the partition of the plane into cells. For example, cell 3 is a union of five trapezoids.

The Merge step is shown in Figure 10.6. In the example, there are two vertical edges that need to be removed (indicated with a cross) because they come from vertical extensions that are cut off by the new segment. In both cases, the two trapezoids to the left and right of the removed edge are being merged into one trapezoid (drawn shaded).

10.4 Using trapezoidal maps for point location

Recall that in the point location problem we want to preprocess a given set $S$ of segments in order to answer subsequent point-location queries: $S$ partitions the plane into connected cells and we want to know, given a query point $q$, to which cell $q$ belongs, see Figure 10.2.

Note that the trapezoidal map of $S$ is a refinement of the partition of the plane into cells, in the sense that a cell might be partitioned into several trapezoids, but every trapezoid belongs to a single cell, see Figure 10.3. Thus, once the trapezoidal map of $S$ is constructed, we can easily "glue together" trapezoids that touch along their vertical sides, obtaining the original cells. Then we can answer point-location queries using the same routine that performs the Find step (whose implementation will be described below).
10.5 Analysis of the incremental construction

In order to analyze the runtime of the incremental construction, we insert the segments in random order, and we employ the configuration space framework. We also implement the Find step in such a way that the analysis boils down to conflict counting, just as for the Delaunay triangulation.

10.5.1 Defining The Right Configurations

Recall that a configuration space is a quadruple $S = (X, \Pi, D, K)$, where $X$ is the ground set, $\Pi$ is the set of configurations, $D$ is a mapping that assigns to each configuration its defining elements ("generators"), and $K$ is a mapping that assigns to each configuration its conflict elements ("killers").

It seems natural to choose $X = S$, the set of segments, and to define $\Pi$ as the set of all possible trapezoids that could appear in the trapezoidal map of some subset of segments. Indeed, this satisfies one important property of configuration spaces: for each configuration, the number of generators is constant.

**Lemma 10.2** For every trapezoid $\square$ in the trapezoidal map of $R \subseteq S$, there exists a set $D \subseteq R$ of at most four segments, such that $\square$ is in the trapezoidal map of $D$.

**Proof.** By our general position assumption, each non-vertical side of $\square$ is a subset of a unique segment in $R$, and each vertical side of $\square$ is induced by a unique (left or right) endpoint, or by the intersection of two unique segments. In the latter case, one of these
segments also contributes a non-vertical side, and in the former case, we attribute the endpoint to the “topmost” segment with that (left or right) endpoint. It follows that there is a set of at most four segments whose trapezoidal map already contains □.

But there is a problem with this definition of configurations. Recall that we can apply the general configuration space analysis only if

(i) the cost of updating \( T_{r-1} \) to \( T_r \) is proportional to the structural change, the number of configurations in \( T_r \setminus T_{r-1} \); and

(ii) the expected cost of all \texttt{Find} operations during the randomized incremental construction is proportional to the expected number of conflicts. (This is the “conflict counting” part.)

Here we see that already (i) fails. During the Trace step, we traverse the boundary of each trapezoid intersected by \( s_r \) in order to find the next trapezoid. Even if \( s_r \) intersects only a small number of trapezoids (so that the structural change is small), the traversals may take very long. This is due to the fact that a trapezoid can be incident to a large number of edges. Consider the trapezoid labeled □ in Figure 10.7. It has many incident vertical extensions from above. Tracing a segment through such a trapezoid takes time that we cannot charge to the structural change.

To deal with this, we slightly adapt our notion of configuration.

\textbf{Definition 10.3} Let \( \Pi \) be the set of all trapezoids together with at most one incident vertical edge (“trapezoids with tail”) that appear in the trapezoidal map of some
Lemma 10.4 For every trapezoid with tail □ in the trapezoidal map of \( R \subseteq S \), there exists a set \( D \subseteq R \) of at most six segments, such that □ is in the trapezoidal map of \( D \).

Proof. We already know from Lemma 10.2 that the trapezoid without tail has at most 4 generators. And since the tail is induced by a unique segment endpoint or by the intersection of a unique pair of segments, the claim follows. \( \square \)

Here is the complete specification of our configuration space \( S = (X, \Pi, D, K) \).
Figure 10.8: A trapezoid with tail is a trapezoid together with at most one vertical edge attached to its upper or its lower segment.

Definition 10.5 Let $X = S$ be the set of segments, and $\Pi$ the set of all trapezoids with tail. For each trapezoid with tail $\square$, $D(\square)$ is the set of at most 6 generators. $K(\square)$ is the set of all segments that intersect $\square$ in the interior of the trapezoid, or cut off some part of the tail, or replace the topmost generator of the left or right side, see Figure 10.9.

Then $S = (X, \Pi, D, K)$ is a configuration space of dimension at most 6, by Lemma 10.4. The only additional property that we need to check is that $D(\square) \cap K(\square) = \emptyset$ for all trapezoids with tail, but this is clear since no generator of $\square$ properly intersects the trapezoid of $\square$ or cuts off part of its tail.

Figure 10.9: A trapezoid with tail $\square$ is in conflict with a segment $s$ (dashed) if $s$ intersects $\square$ in the interior of the trapezoid (left), or cuts off part of the tail (middle), or is a new topmost segment generating a vertical side.

10.5.2 Update Cost

Now we can argue that the update cost can be bounded by the structural change. We employ the same trick as for Delaunay triangulations. We prove that the update cost is in each step $r - 1 \rightarrow r$ proportional to the number of configurations that are being destroyed. Over the whole algorithm, we cannot destroy more configurations than we create, so the bound that we get is also a bound in terms of the overall structural change.
Lemma 10.6 In updating $\mathcal{T}_{r-1}$ to $\mathcal{T}_r$, the steps Trace, Split, and Merge can be performed in time proportional to the number of trapezoids with tail in $\mathcal{T}_{r-1} \setminus \mathcal{T}_r$.

Proof. By definition, the complexity (number of edges) of a trapezoid is proportional to the number of trapezoids with tail that share this trapezoid. This means, the cost of traversing the trapezoid can be charged to the trapezoids with tail containing it, and all of them will be destroyed (this includes the trapezoids with tail that just change their left or right generator; in the configuration space, this is also a destruction). This takes care of the Trace step. The Split and Merge steps can be done within the same asymptotic time bounds since they can be performed by traversing the boundaries of all intersected trapezoids a constant number of times each. For efficiently doing the manipulations on the trapezoidal map, we can for example represent it using a doubly-connected edge list. □

We can now employ the general configuration space analysis to bound the expected structural change throughout the randomized incremental construction; as previously shown, this asymptotically bounds the expected cost of the Trace, Split, and Merge steps throughout the algorithm. Let us recall the general bound.

Theorem 10.7 Let $\mathcal{S} = (X, \Pi, D, K)$ be a configuration space of fixed dimension with $|X| = n$. The expected number of configurations that are created throughout the algorithm is bounded by

$$O \left( \sum_{r=1}^{n} \frac{t_r}{r} \right),$$

where $t_r$ is the expected size of $\mathcal{T}_r$, the expected number of active configurations after $r$ insertion steps.

10.5.3 The History Graph

Here is how we realize the Find step (as well as the point-location queries for our point-location application). It is a straightforward history graph approach as for Delaunay triangulations. Every trapezoid that we ever create is a node in the history graph; whenever a trapezoid is destroyed, we add outgoing edges to its (at most four) successor trapezoids. Note that trapezoids are destroyed during the steps Split and Merge. In the latter step, every destroyed trapezoid has only one successor trapezoid, namely the one it is merged into. It follows that we can prune the nodes of the "ephemeral" trapezoids that exist only between the Split and Merge steps. What we get is a history graph of degree at most 4, such that every non-leaf node corresponds to a trapezoid in $\mathcal{T}_{r-1} \setminus \mathcal{T}_r$, for some $r$. 121
10.5.4 Cost of the Find step

We can use the history graph for point location during the Find step. Given a segment endpoint p, we start from the bounding box (the unique trapezoid with no generators) that is certain to contain p. Since for every trapezoid in the history graph, its area is covered by the at most four successor trapezoids, we can simply traverse the history graph along directed edges until we reach a leaf that contains p. This leaf corresponds to the trapezoid of the current trapezoidal map containing p. By the outdegree-4-property, the cost of the traversal is proportional to the length of the path that we traverse.

Here is the crucial observation that allows us to reduce the analysis of the Find step to “conflict counting”. Note that this is precisely what we also did for Delaunay triangulations, except that there, we had to deal explicitly with “ephemeral” triangles.

Recall Definition 8.5, according to which a conflict is a pair \((\square, s)\) where \(\square\) is a trapezoid with tail, contained in some intermediate trapezoidal map, and \(s \in K(\square)\).

Lemma 10.8 During a run of the incremental construction algorithm for the trapezoidal map, the total number of history graph nodes traversed during all Find steps is bounded by the number of conflicts during the run.

Proof. Whenever we traverse a node (during insertion of segment \(s_r\), say), the node corresponds to a trapezoid \(\square\) (which we also consider as a trapezoid with tail) in some set \(T_s, s < r\), such that \(p \in \square\), where \(p\) is the left endpoint of the segment \(s_r\). We can therefore uniquely identify this edge with the conflict \((\square, s_r)\). The statement follows.

Now we can use the configuration space analysis that precisely bounds the expected number of conflicts, and therefore the expected cost of the Find steps over the whole algorithm. Let us recapitulate the bound.

Theorem 10.9 Let \(S = (X, \Pi, D, K)\) be a configuration space of fixed dimension \(d\) with \(|X| = n\). The expected number of conflicts during randomized incremental construction of \(T_n\) is bounded by

\[
O \left( n \sum_{r=1}^{n} \frac{t_r}{r^2} \right),
\]

where \(t_r\) is as before the expected size of \(T_r\).

10.5.5 Applying the General Bounds

Let us now apply Theorem 10.7 and Theorem 10.9 to our concrete situation of trapezoidal maps. What we obviously need to determine for that is the quantity \(t_r\), the expected number of active configurations after \(r\) insertion steps.

Recall that the configurations are the trapezoids with tail that exist at this point. The first step is easy.
Observation 10.10 In every trapezoidal map, the number of trapezoids with tail is proportional to the number vertices.

Proof. Every trapezoid with tail that actually has a tail can be charged to the vertex of the trapezoidal map on the “trapezoid side” of the tail. No vertex can be charged twice in this way. The trapezoids with no tail are exactly the faces of the trapezoidal map, and since the trapezoidal map is a planar graph, their number is also proportional to the number vertices. □

Using this observation, we have therefore reduced the problem of computing $t_r$ to the problem of computing the expected number of vertices in $T_r$. To count the latter, we note that every segment endpoint and every segment intersection generates 3 vertices: one at the point itself, and two where the vertical extensions hit another feature. Here, we are sweeping the 4 bounding box vertices under the rug.

Observation 10.11 In every trapezoidal map of $r$ segments, the number of vertices is

$$6r + 3k,$$

where $k$ is the number of pairwise intersections between the $r$ segments.

So far, we have not used the fact that we have a random insertion order, but this comes next.

Lemma 10.12 Let $K$ be the total number of pairwise intersections between segments in $S$, and let $k_r$ be the random variable for the expected number of pairwise intersections between the first $r$ segments inserted during randomized incremental construction. Then

$$k_r = K \binom{n-2}{r-2} = K \frac{r(r-1)}{n(n-1)}.$$

Proof. Let us consider the intersection point of two fixed segments $s$ and $s'$. This intersection point appears in $T_r$ if and only both $s$ and $s'$ are among the first $r$ segments. There are $\binom{n}{r}$ ways of choosing the set of $r$ segments (and all choices have the same probability); since the number of $r$-element sets containing $s$ and $s'$ is $\binom{n-2}{r-2}$, the probability for the intersection point to appear is

$$\frac{\binom{n-2}{r-2}}{\binom{n}{r}} = \frac{r(r-1)}{n(n-1)}.$$

Summing this up over all $K$ intersection points, and using linearity of expectation, the statement follows. □

From Observation 10.10 Observation 10.11 and Lemma 10.12, we obtain the following
Corollary 10.13  The expected number $t_r$ of active configurations after $r$ insertion steps is
\[
t_r = O \left( r + K \frac{r(r - 1)}{n^2} \right).
\]

Plugging this into Theorem 10.7 and Theorem 10.9, we obtain the following final

Theorem 10.14  Let $S$ be a set of $n$ segments in the plane, with a total of $K$ pairwise intersections. The randomized incremental construction computes the trapezoidal map of $S$ in time
\[
O(n \log n + K).
\]

Proof.  We already know that the expected update cost (subsuming steps Trace, Split, and Merge) is proportional to the expected overall structural change, which by Theorem 10.7 is
\[
O \left( \sum_{r=1}^{n} \frac{t_r}{r} \right) = O(n) + O \left( \frac{K}{n^2} \sum_{r=1}^{n} r \right) = O(n + K).
\]

We further know that the expected point location cost (subsuming step Find) is proportional to the overall expected number of conflicts which by Theorem 10.9 is
\[
O \left( n \sum_{r=1}^{n} \frac{t_r}{r^2} \right) = O(n \log n) + O \left( \frac{K}{n} \sum_{r=1}^{n} 1 \right) = O(n \log n + K).
\]

\[\square\]

10.6 Analysis of the point location

Finally, we return to the application of trapezoidal maps for point location. We make precise what we mean by saying that “point-location queries are handled in $O(\log n)$ expected time", and we prove our claim.

Lemma 10.15  Let $S = \{s_1, \ldots, s_n\}$ be any set of $n$ segments. Then there exists a constant $c > 0$ such that, with high probability (meaning, with probability tending to 1 as $n \to \infty$), the history graph produced by the random incremental construction answers every possible point-location query in time at most $c \log n$.

Note that our only randomness assumption is over the random permutation of $S$ chosen at the beginning of the incremental construction. We do not make any randomness assumption on the given set of segments.

The proof of Lemma 10.15 is by a typical application of Chernoff's bound followed by the union bound.

Recall (or please meet) Chernoff’s bound:
Lemma 10.16 Let \( X_1, X_2, \ldots, X_n \) be independent 0/1 random variables, and let \( X = X_1 + \cdots + X_n \). Let \( p_i = \Pr[X_i = 1] \), and let \( \mu = E[X] = p_1 + \cdots + p_n \). Then,

\[
\begin{align*}
\Pr[X < (1 - \delta)\mu] &< \left( \frac{e^{-\delta}}{(1 - \delta)^{1-\delta}} \right)^\mu \quad \text{for every } 0 < \delta < 1; \\
\Pr[X > (1 + \delta)\mu] &< \left( \frac{e^{\delta}}{(1 + \delta)^{1+\delta}} \right)^\mu \quad \text{for every } \delta > 0.
\end{align*}
\]

The important thing to note is that \( e^{-\delta}/(1 - \delta)^{1-\delta} \) as well as \( e^{\delta}/(1 + \delta)^{1+\delta} \) are strictly less than 1 for every fixed \( \delta > 0 \), and decrease with increasing \( \delta \).

Now back to the proof of Lemma 10.15.

**Proof.** First note that, even though there are infinitely many possible query points, there is only a finite number of combinatorially distinct possible queries: If two query points lie together in every trapezoid (either both inside or both outside), among all possible trapezoids defined by segments of \( S \), then there is no difference in querying one point versus querying the other, as far as the algorithm is concerned. Since there are \( O(n^4) \) possible trapezoids (recall that each trapezoid is defined by at most four segments), there are only \( O(n^4) \) queries we have to consider.

Fix a query point \( q \). We will show that there exists a large enough constant \( c > 0 \), such that only with probability at most \( O(n^{-5}) \) does the query on \( q \) take more than \( c \log n \) steps.

Let \( s_1, s_2, \ldots, s_n \) be the random order of the segments chosen by the algorithm, and for \( 1 \leq r \leq n \) let \( T_r \) be the trapezoidal map generated by the first \( r \) segments. Note that for every \( r \), the point \( q \) belongs to exactly one trapezoid of \( T_r \). The question is how many times the trapezoid containing \( q \) changes during the insertion of the segments, since these are exactly the trapezoids of the history graph that will be visited when we do a point-location query on \( q \).

For \( 1 \leq r \leq n \), let \( A_r \) be the event that the trapezoid containing \( q \) changes from \( T_{r-1} \) to \( T_r \). What is the probability of \( A_r \)? As in Section 8.3, we "run the movie backwards": To obtain \( T_{r-1} \) from \( T_r \), we delete a random segment from among \( s_1, \ldots, s_r \); the probability that the trapezoid containing \( q \) in \( T_r \) is destroyed is at most \( 4/r \), since this trapezoid is defined by at most four segments. Thus, \( \Pr[A_r] \leq 4/r \), independently of every other \( A_s \), \( s \neq r \).

For each \( r \) let \( X_r \) be a random variable equal to 1 if \( A_r \) occurs, and equal to 0 otherwise. We are interested in the quantity \( X = X_1 + \cdots + X_r \). Then \( \mu = E[X] = \sum_{r=1}^n \Pr[A_r] = 4 \ln n + O(1) \). Applying Chernoff's bound with \( \delta = 2 \) (it is just a matter of choosing \( \delta \) large enough), we get

\[
\Pr[X > (1 + \delta)\mu] < 0.273^\mu = O(0.273^{4\ln n}) = O(n^{-5.19}),
\]

so we can take our \( c \) to be anything larger than \( 4(1 + \delta) = 12 \).

Thus, for every fixed query \( q \), the probability of a "bad event" (a permutation that results in a long query time) is \( O(n^{-5}) \). Since there are only \( O(n^4) \) possible choices for
q, by the union bound the probability of some \( q \) having a bad event is \( O(1/n) \), which tends to zero with \( n \).

### 10.7 The trapezoidal map of a simple polygon

An important special case of the trapezoidal map is obtained when the input segments form a simple polygon; see Figure [10.10]. In this case, we are mostly interested in the part of the trapezoidal map inside the polygon, since that part allows us to obtain a triangulation of the polygon in linear time.

![Figure 10.10: The trapezoidal map inside a simple polygon](image)

To get a triangulation, we first go through all trapezoids; we know that each trapezoid must have one polygon vertex on its left and one on its right boundary (due to general position, there is actually exactly one vertex on each of these two boundaries). Whenever the segment connecting these two vertices is not an edge of the polygon, we have a diagonal, and we insert this diagonal. Once we have done this for all trapezoids, it is easily seen that we have obtained a subdivision of the polygon into \( x \)-monotone polygons, each of which can be triangulated in linear time; see Exercise [10.24]. This immediately allows us to improve over the statement of Exercise [2.16].

**Corollary 10.17** A simple polygon with \( n \) vertices can be triangulated in expected time \( O(n \log n) \).

**Proof.** By Theorem [10.14], the trapezoidal decomposition induced by the segments of a simple polygon can be computed in expected time \( O(n \log n) \), since there are no intersections between the segments. Using the above linear-time triangulation algorithm from the trapezoidal map, the result follows. \( \square \)
The goal of this section is to further improve this bound and show the following result.

**Theorem 10.18** Let $S = \{s_1, s_2, \ldots, s_n\}$ be the set of edges of an $n$-vertex simple polygon, in counterclockwise order around the polygon. The trapezoidal map induced by $S$ (and thus also a triangulation of $S$) can be computed in expected time $O(n \log^* n)$.

Informally speaking, the function $\log^* n$ is the number of times we have to iterate the operation of taking (binary) logarithms, before we get from $n$ down to 1. Formally, we define

$$\log^{(h)}(n) = \begin{cases} n, & \text{if } h = 0 \\ \log^{(h-1)}(\log n), & \text{otherwise} \end{cases}$$

as the $h$-times iterated logarithm, and for $n \geq 1$ we set

$$\log^* n = \max\{h : \log^{(h)} n \geq 1\}.$$

For example, we have

$$\log^*(2^{65536}) = 5,$$

meaning that for all practical purposes, $\log^* n \leq 5$; a bound of $O(n \log^* n)$ is therefore very close to a linear bound.

**History flattening.** Recall that the bottleneck in the randomized incremental construction is the Find step. Using the special structure we have (the segments form a simple polygon in this order), we can speed up this step. Suppose that at some point during incremental construction, we have built the trapezoidal map of a subset of $r$ segments, along with the history graph. We now flatten the history by removing all trapezoids that are not in $T_r$, the current trapezoidal map. To allow for point location also in the future, we need an “entry point” into the flattened history, for every segment not inserted so far (the old entry point for all segments was the bounding unit square $[0,1]^2$).

**Lemma 10.19** Let $S$ be the set of edges of an $n$-vertex simple polygon, in counterclockwise order around the polygon. For $R \subseteq S$, let $T(R)$ be the trapezoidal map induced by $R$. In time proportional to $n$ plus the number of conflicts between trapezoids in $T(R)$ and segments in $S \setminus R$, we can find for all segment $s \in S$ a trapezoid of $T(R)$ that contains an endpoint of $s$.

**Proof.** In a trivial manner (and in time $O(n)$), we do this for the first segment $s_1$ and its first endpoint $p_1$. Knowing the trapezoid $\square_i$ containing $p_l$, the first endpoint of $s_i$, we trace the segment $s_i$ through $T(R)$ until $p_{l+1}$, its other endpoint and first endpoint of $s_{i+1}$ is found, along with the trapezoid $\square_{l+1}$ containing it. This is precisely what we also did in the Trace Step 2 of the incremental construction in Section 10.3.
By Lemma \[10.6\] (pretending that we are about to insert \(s_i\)), the cost of tracing \(s_i\) through \(T(R)\) is proportional to the size of \(T(R) \setminus T(R \cup \{s_i\})\), equivalently, the number of conflicts between trapezoids in \(T(R)\) and \(s_i\). The statement follows by adding up the costs for all \(i\).

This is exactly where the special structure of our segments forming a polygon helps. After locating \(p_i\), we can locate the next endpoint \(p_{i+1}\) in time proportional to the structural change that the insertion of \(s_i\) would cause. We completely avoid the traversal of old history trapezoids that would be necessary for an efficient location of \(p_{i+1}\) from scratch.

Next we show what Lemma \[10.19\] gives us in expectation.

**Lemma 10.20** Let \(S\) be the set of edges of an \(n\)-vertex simple polygon, in counterclockwise order around the polygon, and let \(T_r\) be the trapezoidal map obtained after inserting \(r\) segments in random order. In expected time \(O(n)\), we can find for each segment \(s\) not inserted so far a trapezoid of \(T_r\) containing an endpoint of \(s\).

**Proof.** According to Lemma \[10.19\] the expected time is bounded by \(O(n + \ell(r))\), where

\[
\ell(r) = \frac{1}{\binom{n}{r}} \sum_{R \subseteq S, |R| = r} \sum_{y \in S \setminus R} |\{ \square \in T(R) : y \in K(\square) \}|.
\]

This looks very similar to the bound on the quantity \(k(r)\) that we have derived in Section \[8.5\] to count the expected number of conflicts in general configuration spaces:

\[
k(r) \leq \frac{1}{\binom{n}{r}} \sum_{R \subseteq S, |R| = r} \frac{d}{r} \sum_{y \in S \setminus R} |\{ \Delta \in T(R) : y \in K(\Delta) \}|.
\]

Indeed, the difference is only an additional factor of \(\frac{d}{r}\) in the latter. For \(k(r)\), we have then computed the bound

\[
k(r) \leq k_1(r) - k_2(r) + k_3(r) \leq \frac{d}{r} (n-r)t_r - \frac{d}{r} (n-r)t_{r+1} + \frac{d^2}{r(r+1)} (n-r)t_{r+1},
\]

where \(t_r\) is the expected size of \(T_r\). Hence, to get a bound for \(\ell(r)\), we simply have to cancel \(\frac{d}{r}\) from all terms to obtain

\[
\ell(r) \leq (n-r)t_r - (n-r)t_{r+1} + \frac{d}{r+1} (n-r)t_{r+1} = O(n),
\]

since \(t_r \leq t_{r+1} = O(r)\) in the case of nonintersecting line segments. \(\square\)
10.7. The trapezoidal map of a simple polygon

The faster algorithm. We proceed as in the regular randomized incremental construction, except that we frequently and at well-chosen points flatten the history. Let us define

\[ N(h) = \left\lceil \frac{n}{\log^*(n)} \right\rceil, \quad 0 \leq h \leq \log^* n. \]

We have \( N(0) = 1, N(\log^* n) \leq n \) and \( N(\log^* n + 1) > n \). We insert the segments in random order, but proceed in \( \log^* n + 1 \) rounds. In round \( h = 1, \ldots, \log^* n + 1 \), we do the following.

(i) Flatten the history graph by finding for each segment \( s \) not inserted so far a trapezoid of the current trapezoidal map containing an endpoint of \( s \).

(ii) Insert the segments \( N(h - 1) \) up to \( N(h) - 1 \) in the random order, as usual, but starting from the flat history established in (i).

In the last round, we have \( N(h) - 1 \geq n \), so we stop with segment \( n \) in the random order.

From Lemma 10.20, we know that the expected cost of step (i) over all rounds is bounded by \( O(n \log^* n) \) which is our desired overall bound. It remains to prove that the same bound also deals with step (ii). We do not have to worry about the overall expected cost of performing the structural changes in the trapezoidal map: this will be bounded by \( O(n) \), using \( t_r = O(r) \) and Theorem 10.7. It remains to analyze the Find step, and this is where the history flattening leads to a speedup. Adapting Lemma 10.8 and its proof accordingly, we obtain the following.

**Lemma 10.22** During round \( h \) of the fast incremental construction algorithm for the trapezoidal map, the total number of history graph nodes traversed during all Find steps is bounded by \( N(h) - N(h - 1) \) plus the number of conflicts between trapezoids that are created during round \( h \), and segments inserted during round \( h \).

**Proof.** The history in round \( h \) only contains trapezoids that are active at some point in round \( h \). On the one hand, we have the “root nodes” present immediately after flattening the history, on the other hand the trapezoids that are created during the round. The term \( N(h) - N(h - 1) \) accounts for the traversals of the root nodes during round \( h \). As in the proof of Lemma 10.8, the traversals of other history graph nodes can be charged to the number of conflicts between trapezoids that are created during round \( h \) and segments inserted during round \( h \). 

To count the expected number of conflicts involving trapezoids created in round \( h \), we go back to the general configuration space framework once more. With \( k_h(r) \) being the expected number of conflicts created in step \( r \), restricted to the ones involving segments inserted in round \( h \), we need to bound

\[ \kappa(h) = \sum_{r=N(h-1)}^{N(h)-1} k_h(r). \]

\(^1\text{This amounts to } O(n) \text{ throughout the algorithm and can therefore be neglected}\)
As in (10.21), we get
\[ k_h(r) \leq \frac{d}{r} (N(h) - r)t_r - \frac{d}{r} (N(h) - r)t_{r+1} + \frac{d^2}{r(r+1)} (N(h) - r)t_{r+1}, \]
where we have replaced \( n \) with \( N(h) \), due to the fact that we do not need to consider segments in later rounds. Then \( t_r = O(r) \) yields
\[ \kappa(h) \leq O \left( N(h) \sum_{r=N(h)-1}^{N(h)-1} \frac{1}{r} \right) \]
\[ = O \left( N(h) \log \frac{N(h)}{N(h-1)} \right) \]
\[ = O \left( N(h) \log \frac{\log^{(h-1)} n}{\log^{(h)} n} \right) \]
\[ = O \left( N(h) \log^{(h)} n \right) = O(n). \]

It follows that step (ii) of the fast algorithm also requires expected linear time per round, and Theorem 10.18 follows.

Exercise 10.23 a) You are given a set of \( n \) pairwise disjoint line segments. Find an algorithm to answer vertical ray shooting queries in \( O(\log n) \) time. That is, preprocess the data such that given a query point \( q \) you can report in \( O(\log n) \) time which segment is the first above \( q \) (or if there are none). Analyze the running time and the space consumption of the preprocessing.

b) What happens if we allow intersections of the line segments? Explain in a few words how you have to adapt your solution and how the time and space complexity would change.

Exercise 10.24 Show that an \( n \)-vertex \( x \)-monotone polygon can be triangulated in time \( O(n) \). (As usual a polygon is given as a list of its vertices in counterclockwise order. A polygon \( P \) is called \( x \)-monotone if for all vertical lines \( \ell \), the intersection \( \ell \cap P \) has at most one component.)

Questions

41. What is the definition of a trapezoidal map?

42. How does the random incremental construction of a trapezoidal map proceed? What are the main steps to be executed at each iteration?

43. How can trapezoidal maps be used for the point location problem?
44. What is the configuration space framework? Recall Section 8.3.

45. What is a naive way of defining a configuration in the case of trapezoids, and why does it fail?

46. What is a more successful way of defining a configuration? Why do things work in this case?

47. What is the history graph, and how is it used to answer point location queries?

48. What is the performance of the random incremental construction of the trapezoidal map when used for the point-location problem? Be precise!

49. What probabilistic techniques are used in proving this performance bound?

50. How can you speed up the randomized incremental construction in the case where the input segments from a simple polygon? Sketch the necessary changes to the algorithm, and how they affect the analysis.
Chapter 11

Line Arrangements

During the course of this lecture we encountered several situations where it was convenient to assume that a point set is “in general position”. In the plane, general position usually amounts to no three points being collinear and/or no four of them being cocircular. This raises an algorithmic question: How can we test for $n$ given points whether or not three of them are collinear? Obviously, we can test all triples in $O(n^3)$ time. Can we do better? Yes, we can! Using a detour through the so-called dual plane, we will see that this problem can be solved in $O(n^2)$ time. However, the exact algorithmic complexity of this innocent-looking problem is not known. In fact, to determine this complexity is one of the major open problems in theoretical computer science.

We will get back to the complexity theoretic problems and ramifications at the end of this chapter. But first let us discuss how to obtain a quadratic time algorithm to test whether $n$ given points in the plane are in general position. This algorithm is a nice application of the projective duality transform, as defined below. Such transformations are very useful because they allow us to gain a new perspective on a problem by formulating it in a different but equivalent form. Sometimes such a dual form of the problem is easier to work with and—given that it is equivalent to the original primal form—any solution to the dual problem can be translated back into a solution to the primal problem.

So what is this duality transform about? Observe that points and hyperplanes in $\mathbb{R}^d$ are very similar objects, given that both can be described using $d$ coordinates/parameters. It is thus tempting to match these parameters to each other and so create a mapping between points and hyperplanes. In $\mathbb{R}^2$ hyperplanes are lines and the standard projective duality transform maps a point $p = (p_x, p_y)$ to the line $p^* : y = p_x x - p_y$ and a non-vertical line $g : y = mx + b$ to the point $g^* = (m, -b)$.

**Proposition 11.1** The standard projective duality transform is

- **incidence preserving**: $p \in g \iff g^* \in p^*$ and
- **order preserving**: $p$ is above $g \iff g^*$ is above $p^*$.

**Exercise 11.2** Prove Proposition 11.1.
Exercise 11.3 Describe the image of the following point sets under this mapping

a) a halfplane

b) $k \geq 3$ collinear points

c) a line segment

d) the boundary points of the upper convex hull of a finite point set.

Another way to think of duality is in terms of the parabola $P : y = \frac{1}{2}x^2$. For a point $p$ on $P$, the dual line $p^*$ is the tangent to $P$ at $p$. For a point $p$ not on $P$, consider the vertical projection $p'$ of $p$ onto $P$: the slopes of $p^*$ and $p'^*$ are the same, just $p^*$ is shifted by the difference in $y$-coordinates.

The question of whether or not three points in the primal plane are collinear transforms to whether or not three lines in the dual plane meet in a point. This question in turn we will answer with the help of line arrangements, as defined below.

11.1 Arrangements

The subdivision of the plane induced by a finite set $L$ of lines is called the arrangement $A(L)$. Observe that all cells of the subdivision are intersections of halfplanes and thus convex. A line arrangement is simple if no two lines are parallel and no three lines meet in a point. Although lines are unbounded, we can regard a line arrangement a bounded object by (conceptually) putting a sufficiently large box around that contains all vertices. Such a box can be constructed in $O(n \log n)$ time for $n$ lines.

Exercise 11.4 How?
Moreover, we can view a line arrangement as a planar graph by adding an additional vertex at "infinity", that is incident to all rays which leave this bounding box. For algorithmic purposes, we will mostly think of an arrangement as being represented by a doubly connected edge list (DCEL), cf. Section 4.2.

**Theorem 11.5** A simple arrangement \(A(L)\) of \(n\) lines in \(\mathbb{R}^2\) has \(\binom{n}{2}\) vertices, \(n^2\) edges, and \(\binom{n}{2} + n + 1\) faces/cells.

**Proof.** Since all lines intersect and all intersection points are pairwise distinct, there are \(\binom{n}{2}\) vertices.

The number of edges we count using induction on \(n\). For \(n = 1\) we have \(1^2 = 1\) edge.

By adding one line to an arrangement of \(n - 1\) lines we split \(n - 1\) existing edges into two and introduce \(n\) new edges along the newly inserted line. Thus, there are in total \((n - 1)^2 + 2n - 1 = n^2 - 2n + 1 + 2n - 1 = n^2\) edges.

The number \(f\) of faces can now be obtained from Euler’s formula \(v - e + f = 2\), where \(v\) and \(e\) denote the number of vertices and edges, respectively. However, in order to apply Euler’s formula we need to consider \(A(L)\) as a planar graph and take the symbolic “infinite” vertex into account. Therefore,

\[
f = 2 - \left(\binom{n}{2} + 1\right) + n^2 = 1 + \frac{1}{2}(2n^2 - n(n - 1)) = 1 + \frac{1}{2}(n^2 + n) = 1 + \binom{n}{2} + n.
\]

The complexity of an arrangement is simply the total number of vertices, edges, and faces (in general, cells of any dimension).

**Exercise 11.6** Consider a set of lines in the plane with no three intersecting in a common point. Form a graph \(G\) whose vertices are the intersection points of the lines and such that two vertices are adjacent if and only if they appear consecutively along one of the lines. Prove that \(\chi(G) \leq 3\), where \(\chi(G)\) denotes the chromatic number of the graph \(G\). In other words, show how to color the vertices of \(G\) using at most three colors such that no two adjacent vertices have the same color.

### 11.2 Construction

As the complexity of a line arrangement is quadratic, there is no need to look for a sub-quadratic algorithm to construct it. We will simply construct it incrementally, inserting the lines one by one. Let \(\ell_1, \ldots, \ell_n\) be the order of insertion.

At Step \(i\) of the construction, locate \(\ell_i\) in the leftmost cell of \(A((\ell_1, \ldots, \ell_{i-1}))\) it intersects. (The halfedges leaving the infinite vertex are ordered by slope.) This takes \(O(i)\) time. Then traverse the boundary of the face \(F\) found until the halfedge \(h\) is found where \(\ell_i\) leaves \(F\) (see Figure 11.2 for illustration). Insert a new vertex at this point, splitting \(F\) and \(h\) and continue in the same way with the face on the other side of \(h\).
Chapter 11. Line Arrangements

The insertion of a new vertex involves splitting two halfedges and thus is a constant time operation. But what is the time needed for the traversal? The complexity of \( \mathcal{A}(\{\ell_1, \ldots, \ell_{i-1}\}) \) is \( \Theta(i^2) \), but we will see that the region traversed by a single line has linear complexity only.

11.3 Zone Theorem

For a line \( \ell \) and an arrangement \( \mathcal{A}(L) \), the zone \( Z_{\mathcal{A}(L)}(\ell) \) of \( \ell \) in \( \mathcal{A}(L) \) is the set of cells from \( \mathcal{A}(L) \) whose closure intersects \( \ell \).

**Theorem 11.7** Given an arrangement \( \mathcal{A}(L) \) of \( n \) lines in \( \mathbb{R}^2 \) and a line \( \ell \) (not necessarily from \( L \)), the total number of edges in all cells of the zone \( Z_{\mathcal{A}(L)}(\ell) \) is at most \( 6n \).

**Proof.** Without loss of generality suppose that \( \ell \) is horizontal and that none of the lines from \( L \) is horizontal. (The first condition can be addressed by rotating the plane and the second by deciding that the left vertex of a horizontal edge is higher than the right vertex.)

For each cell of \( Z_{\mathcal{A}(L)}(\ell) \) split its boundary at its topmost vertex and at its bottommost vertex and orient all edges from bottom to top. Those edges that have the cell to their right are called left-bounding for the cell and those edges that have the cell to their left are called right-bounding. For instance, for the cell depicted to the right all left-bounding edges are shown blue and bold.

We will show that there are at most \( 3n \) left-bounding edges in \( Z_{\mathcal{A}(L)}(\ell) \) by induction on \( n \). By symmetry, the same bound holds also for the number of right-bounding edges.
11.3. Zone Theorem

in $Z_{\mathcal{A}(L)}(\ell)$. For $n = 1$, there is at most one (exactly one, unless $\ell$ is parallel to and lies below the only line in $L$) left-bounding edge in $Z_{\mathcal{A}(L)}(\ell)$ and $1 \leq 3n = 3$. Assume the statement is true for $n - 1$.

![Figure 11.3: At most three new left-boundary edges are created by adding $r$ to $\mathcal{A}(L\setminus\{r\})$.](image)

If no line from $L$ intersects $\ell$, then all lines in $L \cup \{\ell\}$ are parallel and there is at most $1 < 3n$ left-bounding edge in $Z_{\mathcal{A}(L)}(\ell)$. Else consider the rightmost line $r$ from $L$ intersecting $\ell$ and the arrangement $\mathcal{A}(L\setminus\{r\})$. By the induction hypothesis there are at most $3n - 3$ left-bounding edges in $Z_{\mathcal{A}(L\setminus\{r\})}(\ell)$. Adding $r$ back adds at most three new left-bounding edges: At most two edges (call them $\ell_0$ and $\ell_1$) of the rightmost cell of $Z_{\mathcal{A}(L\setminus\{r\})}(\ell)$ are intersected by $r$ and thereby split in two. Both of these two edges may be left-bounding and thereby increase the number of left-bounding edges by at most two. In any case, $r$ itself contributes exactly one more left-bounding edge to that cell. The line $r$ cannot contribute a left-bounding edge to any cell other than the rightmost: to the left of $r$, the edges induced by $r$ form right-bounding edges only and to the right of $r$ all other cells touched by $r$ (if any) are shielded away from $\ell$ by one of $\ell_0$ or $\ell_1$. Therefore, the total number of left-bounding edges in $Z_{\mathcal{A}(L)}(\ell)$ is bounded from above by $3 + 3n - 3 = 3n$. 

**Corollary 11.8** The arrangement of $n$ lines in $\mathbb{R}^2$ can be constructed in optimal $O(n^2)$ time and space.

**Proof.** Use the incremental construction described above. In Step $i$, for $1 \leq i \leq n$, we do a linear search among $i - 1$ elements to find the starting face and then traverse (part of) the zone of the line $\ell_i$ in the arrangement $\mathcal{A}((\ell_1, \ldots, \ell_{i-1}))$. By Theorem 11.7 the complexity of this zone and hence the time complexity of Step $i$ altogether is $O(i)$. Overall we obtain $\sum_{i=1}^{n} ci = O(n^2)$ time (and space), for some constant $c > 0$, which is optimal by Theorem 11.5. 

The corresponding bounds for hyperplane arrangements in $\mathbb{R}^d$ are $\Theta(n^d)$ for the complexity of a simple arrangement and $O(n^{d-1})$ for the complexity of a zone of a hyperplane.
Exercise 11.9 For an arrangement $A$ of a set of $n$ lines in $\mathbb{R}^2$, let $\mathcal{F} := \bigcup_{C \text{ is cell of } A} \overline{C}$ denote the union of the closure of all bounded cells. Show that the complexity (number of vertices and edges of the arrangement lying on the boundary) of $\mathcal{F}$ is $O(n)$.

### 11.4 The Power of Duality

The real beauty and power of line arrangements becomes apparent in context of projective point ↔ line duality. The following problems all can be solved in $O(n^2)$ time and space by constructing the dual arrangement.

**General position test.** Given $n$ points in $\mathbb{R}^2$, are any three of them collinear? (Dual: do three lines meet in a point?)

**Minimum area triangle.** Given $n$ points in $\mathbb{R}^2$, what is the minimum area triangle spanned by any three of them? For any vertex $l^*$ of the dual arrangement (primal: line $l$ through two points $p$ and $q$) find the closest point vertically above/below $l^*$ through which an input line passes (primal: closest line below/above and parallel to $l$ that passes through an input point). In this way one can find $O(n^2)$ candidate triangles by constructing the arrangement of the $n$ dual lines. The smallest among those candidates can be determined by a straightforward minimum selection (comparing the area of the corresponding triangles). Observe that vertical distance is not what determines the area of the corresponding triangle but orthogonal distance. However, the points that minimize these measures for any fixed line are the same...

Exercise 11.10 A set $P$ of $n$ points in the plane is said to be in $\varepsilon$-general position for $\varepsilon > 0$ if no three points of the form

$$p + (x_1, y_1), q + (x_2, y_2), r + (x_3, y_3)$$

are collinear, where $p, q, r \in P$ and $|x_i|, |y_i| < \varepsilon$, for $i \in \{1, 2, 3\}$. In words: $P$ remains in general position under changing point coordinates by less than $\varepsilon$ each.

Give an algorithm with runtime $O(n^2)$ for checking whether a given point set $P$ is in $\varepsilon$-general position.

### 11.5 Sorting all Angular Sequences.

Theorem 11.11 Consider a set $P$ of $n$ points in the plane. For a point $q \in P$ let $c_P(q)$ denote the circular sequence of points from $S \setminus \{q\}$ ordered counterclockwise around

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1For instance, maintain over the incremental construction for each vertex a vertically closest line. The number of vertices to be updated during insertion of a line $\ell$ corresponds to the complexity of the zone of $\ell$ in the arrangement constructed so far. Therefore maintaining this information comes at no extra cost asymptotically.
q (in order as they would be encountered by a ray sweeping around q). All \( c_p(q) \), \( q \in P \), collectively can be obtained in \( O(n^2) \) time.

**Proof.** Assume without loss of generality that no two points in \( P \) have the same \( x \)-coordinate (else rotate the plane infinitesimally). Consider the projective dual \( P^* \) of \( P \). An angular sweep around a point \( q \in P \) in the primal plane corresponds to a traversal of the line \( q^* \) from left to right in the dual plane. (A collection of lines through a single point \( q \) corresponds to a collection of points on a single line \( q^* \) and slope corresponds to \( x \)-coordinate.) Clearly, the sequence of intersection points along all lines in \( P^* \) can be obtained by constructing the arrangement in \( O(n^2) \) time. In the primal plane, any such sequence corresponds to an order of the remaining points according to the slope of the connecting line; to construct the circular sequence of points as they are encountered around \( q \), we have to split the sequence obtained from the dual into those points that are to the left of \( q \) and those that are to the right of \( q \); concatenating both yields the desired sequence.

**Exercise 11.12 (Eppstein [? ])** Describe an \( O(n^2) \) time algorithm that given a set \( P \) of \( n \) points in the plane finds a subset of five points that form a strictly convex empty pentagon (or reports that there is none if that is the case). Empty means that the convex pentagon may not contain any other points of \( P \).

**Hint:** Start with a point \( p \in P \) that is extremal in one direction and try to find out whether there is a solution \( P' \) containing \( p \). For this, consider the star-shaped polygon that visits all points in radial order, as seen from \( p \).

**Remark:** It was shown by Harborth [? ] that every set of ten or more points in general position contains a subset of five points that form a strictly convex empty pentagon.

### 11.6 Segment Endpoint Visibility Graphs

A fundamental problem in motion planning is to find a short(est) path between two given positions in some domain, subject to certain constraints. As an example, suppose we are given two points \( p, q \in \mathbb{R}^2 \) and a set \( S \subseteq \mathbb{R}^2 \) of obstacles. What is the shortest path between \( p \) and \( q \) that avoids \( S \)?

**Observation 11.13** The shortest path (if it exists) between two points that does not cross a finite set of finite polygonal obstacles is a polygonal path whose interior vertices are obstacle vertices.

One of the simplest type of obstacle conceivable is a line segment. In general the plane may be disconnected with respect to the obstacles, for instance, if they form a closed curve. However, if we restrict the obstacles to pairwise disjoint line segments then there is always a free path between any two given points. Apart from start and goal position, by the above observation we may restrict our attention concerning shortest
paths to straight line edges connecting obstacle vertices, in this case, segment endpoints.

**Definition 11.14** Consider a set $S$ of $n$ disjoint line segments in $\mathbb{R}^2$. The segment endpoint visibility graph $V(S)$ is a geometric straight line graph defined on the segment endpoints. Two segment endpoints $p$ and $q$ are connected in $V(S)$ if and only if

- the line segment $pq$ is in $S$ or
- $pq \cap s \subseteq \{p, q\}$ for every segment $s \in S$.

![Figure 11.4: A set of disjoint line segments and their endpoint visibility graph.](image)

If all segments are on the convex hull, the visibility graph is complete. If they form parallel chords of a convex polygon, the visibility graph consists of copies of $K_4$, glued together along opposite edges and the total number of edges is linear only.

These graphs also appear in the context of the following question: Given a set of disjoint line segments, is it possible to connect them to form (the boundary of) a simple polygon? Is it easy to see that this is not possible in general: Just take three parallel chords of a convex polygon (Figure 11.5a). However, if we do not insist that the segments appear on the boundary, but allow them to be diagonals or epigonals, then it is always possible [5, 4]. In other words, the segment endpoint visibility graph of disjoint line segments is Hamiltonian, unless all segments are collinear. It is actually essential to allow epigonals and not only diagonals [9, 3] (Figure 11.5b).

![Figure 11.5: Sets of disjoint line segments that do not allow certain polygons.](image)

Constructing $V(S)$ for a given set $S$ of disjoint segments in a brute force way takes $O(n^3)$ time. (Take all pairs of endpoints and check all other segments for obstruction.)
Theorem 11.15 (Welzl [10]) The segment endpoint visibility graph of \( n \) disjoint line segments can be constructed in worst case optimal \( O(n^2) \) time.

**Proof.** For simplicity we assume general position, that is, no three endpoints are collinear and no two have the same \( x \)-coordinate. It is no problem to handle such degeneracies explicitly.

We have seen above how all sorted angular sequences can be obtained from the dual line arrangement in \( O(n^2) \) time. Topologically sweep the arrangement from left to right (corresponds to changing the slope of the primal rays from \(-\infty\) to \(+\infty\)) while maintaining for each segment endpoint \( p \) the segment \( s(p) \) it currently “sees” (if any). Initialize by brute force in \( O(n^2) \) time (direction vertically downwards). Each intersection of two lines corresponds to two segment endpoints “seeing” each other along the primal line whose dual is the point of intersection. In order to process an intersection, we only need that all preceding (located to the left) intersections of the two lines involved have already been processed. This order corresponds to a topological sort of the arrangement graph where all edges are directed from left to right. (Clearly, this graph is acyclic, that is, it does not contain a directed cycle.) A topological sort can be obtained, for instance, via (reversed) post order DFS in time linear in the size of the graph (number of vertices and edges), which in our case here is \( O(n^2) \).

When processing an intersection, there are four cases. Let \( p \) and \( q \) be the two points involved such that \( p \) is to the left of \( q \).

1. The two points belong to the same input segment \( \rightarrow \) output the edge \( pq \), no change otherwise.
2. \( q \) is obscured from \( p \) by \( s(p) \) \( \rightarrow \) no change.
3. \( q \) is endpoint of \( s(p) \) \( \rightarrow \) output \( pq \) and update \( s(p) \) to \( s(q) \).
4. Otherwise \( q \) is endpoint of a segment \( t \) that now obscures \( s(p) \) \( \rightarrow \) output \( pq \) and update \( s(p) \) to \( t \).

Thus any intersection can be processed in constant time and the overall runtime of this algorithm is quadratic. 

11.7 Ham Sandwich Theorem

Suppose two thieves have stolen a necklace that contains rubies and diamonds. Now it is time to distribute the prey. Both, of course, should get the same number of rubies and the same number of diamonds. On the other hand, it would be a pity to completely disintegrate the beautiful necklace. Hence they want to use as few cuts as possible to achieve a fair gem distribution.

To phrase the problem in a geometric (and somewhat more general) setting: Given two finite sets \( R \) and \( D \) of points, construct a line that bisects both sets, that is, in either
halfplane defined by the line there are about half of the points from $R$ and about half of the points from $D$. To solve this problem, we will make use of the concept of levels in arrangements.

**Definition 11.16** Consider an arrangement $\mathcal{A}(L)$ induced by a set $L$ of $n$ non-vertical lines in the plane. We say that a point $p$ is on the $k$-level in $\mathcal{A}(L)$ if there are at most $k - 1$ lines below and at most $n - k$ lines above $p$. The 1-level and the $n$-level are also referred to as lower and upper envelope, respectively.

![Figure 11.6: The 3-level of an arrangement.](image)

Another way to look at the $k$-level is to consider the lines to be real functions; then the lower envelope is the pointwise minimum of those functions, and the $k$-level is defined by taking pointwise the $k^{th}$-smallest function value.

**Theorem 11.17** Let $R, D \subset \mathbb{R}^2$ be finite sets of points. Then there exists a line that bisects both $R$ and $D$. That is, in either open halfplane defined by $\ell$ there are no more than $|R|/2$ points from $R$ and no more than $|D|/2$ points from $D$.

**Proof.** Without loss of generality suppose that both $|R|$ and $|D|$ are odd. (If, say, $|R|$ is even, simply remove an arbitrary point from $R$. Any bisector for the resulting set is also a bisector for $R$.) We may also suppose that no two points from $R \cup D$ have the same $x$-coordinate. (Otherwise, rotate the plane infinitesimally.)

Let $R^*$ and $D^*$ denote the set of lines dual to the points from $R$ and $D$, respectively. Consider the arrangement $\mathcal{A}(R^*)$. The median level of $\mathcal{A}(R^*)$ defines the bisecting lines for $R$. As $|R| = |R^*|$ is odd, both the leftmost and the rightmost segment of this level are defined by the same line $\ell_r$ from $R^*$, the one with median slope. Similarly there is a corresponding line $\ell_d$ in $\mathcal{A}(D^*)$.

Since no two points from $R \cup D$ have the same $x$-coordinate, no two lines from $R^* \cup D^*$ have the same slope, and thus $\ell_r$ and $\ell_d$ intersect. Consequently, being piecewise linear continuous functions, the median level of $\mathcal{A}(R^*)$ and the median level of $\mathcal{A}(D^*)$ intersect (see Figure 11.7 for an example). Any point that lies on both median levels corresponds to a primal line that bisects both point sets simultaneously. \qed
How can the thieves use Theorem 11.17? If they are smart, they drape the necklace along some convex curve, say, a circle. Then by Theorem 11.17 there exists a line that simultaneously bisects the set of diamonds and the set of rubies. As any line intersects the circle at most twice, the necklace is cut at most twice. It is easy to turn the proof given above into an $O(n^2)$ algorithm to construct a line that simultaneously bisects both sets.

You can also think of the two point sets as a discrete distribution of a ham sandwich that is to be cut fairly, that is, in such a way that both parts have the same amount of ham and the same amount of bread. That is where the name “ham sandwich cut” comes from. The theorem generalizes both to higher dimension and to more general types of measures (here we study the discrete setting only where we simply count points). These generalizations can be proven using the Borsuk-Ulam Theorem, which states that any continuous map from $S^d$ to $R^d$ must map some pair of antipodal points to the same point. For a proof of both theorems see, for instance, Matoušek’s book [8].

**Theorem 11.18** Let $P_1, \ldots, P_d \subset R^d$ be finite sets of points. Then there exists a hyperplane $H$ that simultaneously bisects all of $P_1, \ldots, P_d$. That is, in either open halfspace defined by $H$ there are no more than $|P_i|/2$ points from $P_i$, for every $i \in \{1, \ldots, d\}$.

This implies that the thieves can fairly distribute a necklace consisting of $d$ types of gems using at most $d$ cuts.

In the plane, a ham sandwich cut can be found in linear time using a sophisticated prune and search algorithm by Lo, Matoušek and Steiger [7]. But in higher dimension, the algorithmic problem gets harder. In fact, already for $R^3$ the complexity of finding a ham sandwich cut is wide open: The best algorithm known, from the same paper by Lo et al. [7], has runtime $O(n^{3/2}\log^2 n/\log^* n)$ and no non-trivial lower bound is known. If the dimension $d$ is not fixed, it is both NP-hard and W[1]-hard\(^2\) in $d$ to decide the

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\(^2\)Essentially this means that it is unlikely to be solvable in time $O(f(d)p(n))$, for an arbitrary function
following question [6]: Given $d \in \mathbb{N}$, finite point sets $P_1, \ldots, P_d \subset \mathbb{R}^d$, and a point $p \in \bigcup_{i=1}^d P_i$, is there a ham sandwich cut through $p$?

**Exercise 11.19** The goal of this exercise is to develop a data structure for halfspace range counting.

a) Given a set $P \subset \mathbb{R}^2$ of $n$ points in general position, show that it is possible to partition this set by two lines such that each region contains at most $\left\lceil \frac{n}{4} \right\rceil$ points.

b) Design a data structure of size $O(n)$, which can be constructed in time $O(n \log n)$ and allows you, for any halfspace $h$, to output the number of points $|P \cap h|$ of $P$ contained in this halfspace $h$ in time $O(n^\alpha)$, for some $0 < \alpha < 1$.

**Exercise 11.20** Prove or disprove the following statement: Given three finite sets $A, B, C$ of points in the plane, there is always a circle or a line that bisects $A, B$ and $C$ simultaneously (that is, no more than half of the points of each set are inside or outside the circle or on either side of the line, respectively).

### 11.8 3-Sum

The 3-Sum problem is the following: Given a set $S$ of $n$ integers, does there exist a three-tuple of elements from $S$ that sum up to zero? By testing all three-tuples this can obviously be solved in $O(n^3)$ time. If the tuples to be tested are picked a bit more cleverly, we obtain an $O(n^2)$ algorithm.

Let $(s_1, \ldots, s_n)$ be the sequence of elements from $S$ in increasing order. Then we test the tuples as follows.

For $i = 1, \ldots, n$ {
  
  j = i, k = n.

  While $k \geq j$ {
    If $s_i + s_j + s_k = 0$ then exit with triple $s_i, s_j, s_k$.
    If $s_i + s_j + s_k > 0$ then $k = k - 1$ else $j = j + 1$.
  }
}

The runtime is clearly quadratic (initial sorting can be done in $O(n \log n)$ time). Regarding the correctness observe that the following is an invariant that holds at the start of every iteration of the inner loop: $s_i + s_x + s_k < 0$, for all $i \leq x < j$, and $s_i + s_j + s_x > 0$, for all $k < x \leq n$.

\[ ^f \text{and a polynomial } p. \]

\[ ^3 \text{That is, an element of } S \text{ may be chosen twice or even three times, although the latter makes sense for the number } 0 \text{ only. :(-) } \]
Interestingly, this is essentially the best algorithm known for 3-Sum. It is widely believed that the problem cannot be solved in sub-quadratic time, but so far this has been proved in some very restricted models of computation only, such as the linear decision tree model [1].

11.9 3-Sum hardness

There is a whole class of problems that are equivalent to 3-Sum up to sub-quadratic time reductions [2]; such problems are referred to as 3-Sum-hard.

**Definition 11.21** A problem $P$ is 3-Sum-hard if and only if every instance of 3-Sum of size $n$ can be solved using a constant number of instances of $P$—each of $O(n)$ size—and $o(n^2)$ additional time.

For instance, it is not hard to show that the following variation of 3-Sum—let us denote it by 3-Sum$^\circ$—is 3-Sum hard: Given a set $S$ of $n$ integers, does there exist a three-element subset of $S$ whose elements sum up to zero?

As another example, consider the Problem GeomBase: Given $n$ points on the three horizontal lines $y = 0$, $y = 1$, and $y = 2$, is there a non-horizontal line that contains at least three of them?

3-Sum can be reduced to GeomBase as follows. For an instance $S = \{s_1, \ldots, s_n\}$ of 3-Sum, create an instance $P$ of GeomBase in which for each $s_i$ there are three points in $P$: $(s_i, 0)$, $(-s_i/2, 1)$, and $(s_i, 2)$. If there are any three collinear points in $P$, there must be one from each of the lines $y = 0$, $y = 1$, and $y = 2$. So suppose that $p = (s_i, 0)$, $q = (-s_j/2, 1)$, and $r = (s_k, 2)$ are collinear. The inverse slope of the line through $p$ and $q$ is $-s_j/2 - s_i$ and the inverse slope of the line through $q$ and $r$ is $s_k + s_i/2$. The three points are collinear if and only if the two slopes are equal, that is, $-s_j/2 - s_i = s_k + s_i/2 \iff s_i + s_j + s_k = 0$.

A very similar problem is General Position, in which one is given $n$ arbitrary points and has to decide whether any three are collinear. For an instance $S$ of 3-Sum$^\circ$, create an instance $P$ of General Position by projecting the numbers $s_i$ onto the curve $y = x^3$, that is, $P = \{(a, a^3) \mid a \in S\}$.

Suppose three of the points, say, $(a, a^3)$, $(b, b^3)$, and $(c, c^3)$ are collinear. This is the case if and only if the slopes of the lines through each pair of them are equal. (Observe that $a$, $b$, and $c$ are pairwise distinct.)

\[
\begin{align*}
(b^3 - a^3)/(b - a) &= (c^3 - b^3)/(c - b) \iff \\
 b^2 + a^2 + ab &= c^2 + b^2 + bc \iff \\
 b &= (c^2 - a^2)/(a - c) \iff \\
 b &= -(a + c) \iff \\
 a + b + c &= 0.
\end{align*}
\]
Minimum Area Triangle is a strict generalization of General Position and, therefore, also 3-Sum-hard.

In Segment Splitting/Separation, we are given a set of \( n \) line segments and have to decide whether there exists a line that does not intersect any of the segments but splits them into two non-empty subsets. To show that this problem is 3-Sum-hard, we can use essentially the same reduction as for GeomBase, where we interpret the points along the three lines \( y = 0, y = 1, \) and \( y = 2 \) as sufficiently small “holes”. The parts of the lines that remain after punching these holes form the input segments for the Splitting problem. Horizontal splits can be prevented by putting constant size gadgets somewhere beyond the last holes, see the figure below. The set of input segments for the segment splitting problem requires sorting the points along each of the three horizontal lines, which can be done in \( O(n \log n) = o(n^2) \) time. It remains to specify what “sufficiently small” means for the size of those holes. As all input numbers are integers, it is not hard to see that punching a hole of \( (x - 1/4, x + 1/4) \) around each input point \( x \) is small enough.

In Segment Visibility, we are given a set \( S \) of \( n \) horizontal line segments and two segments \( s_1, s_2 \in S \). The question is: Are there two points, \( p_1 \in s_1 \) and \( p_2 \in s_2 \) which can see each other, that is, the open line segment \( p_1 p_2 \) does not intersect any segment from \( S \)? The reduction from 3-Sum is the same as for Segment Splitting, just put \( s_1 \) above and \( s_2 \) below the segments along the three lines.

In Motion Planning, we are given a robot (line segment), some environment (modeled as a set of disjoint line segments), and a source and a target position. The question is: Can the robot move (by translation and rotation) from the source to the target position, without ever intersecting the “walls” of the environment?

To show that Motion Planning is 3-Sum-hard, employ the reduction for Segment Splitting from above. The three “punched” lines form the doorway between two rooms, each modeled by a constant number of segments that cannot be split, similar to the boundary gadgets above. The source position is in one room, the target position in the other, and to get from source to target the robot has to pass through a sequence of three collinear holes in the door (suppose the doorway is sufficiently small compared to the length of the robot).

Exercise 11.22 The 3-Sum’ problem is defined as follows: given three sets \( S_1, S_2, S_3 \) of \( n \) integers each, are there \( a_1 \in S_1, a_2 \in S_2, a_3 \in S_3 \) such that \( a_1 + a_2 + a_3 = 0 \)? Prove that the 3-Sum’ problem and the 3-Sum problem as defined in the lecture
(S_1 = S_2 = S_3) are equivalent, more precisely, that they are reducible to each other in subquadratic time.

Questions

51. How can one construct an arrangement of lines in \( \mathbb{R}^2 \)? Describe the incremental algorithm and prove that its time complexity is quadratic in the number of lines (incl. statement and proof of the Zone Theorem).

52. How can one test whether there are three collinear points in a set of \( n \) given points in \( \mathbb{R}^2 \)? Describe an \( O(n^2) \) time algorithm.

53. How can one compute the minimum area triangle spanned by three out of \( n \) given points in \( \mathbb{R}^2 \)? Describe an \( O(n^2) \) time algorithm.

54. What is a ham sandwich cut? Does it always exist? How to compute it? State and prove the theorem about the existence of a ham sandwich cut in \( \mathbb{R}^2 \) and describe an \( O(n^2) \) algorithm to compute it.

55. What is the endpoint visibility graph for a set of disjoint line segments in the plane and how can it be constructed? Give the definition and explain the relation to shortest paths. Describe the \( O(n^2) \) algorithm by Welzl, including full proofs of Theorem 11.11 and Theorem 11.15.

56. Is there a subquadratic algorithm for General Position? Explain the term 3-Sum hard and its implications and give the reduction from 3-Sum to General Position.

57. Which problems are known to be 3-Sum-hard? List at least three problems (other than 3-Sum) and briefly sketch the corresponding reductions.

References


Chapter 12

Davenport-Schinzel Sequences

The complexity of a simple arrangement of \( n \) lines in \( \mathbb{R}^2 \) is \( \Theta(n^2) \) and so every algorithm that uses such an arrangement explicitly needs \( \Omega(n^2) \) time. However, there are many scenarios in which we do not need the whole arrangement but only some part of it. For instance, to construct a ham sandwich cut for two sets of points in \( \mathbb{R}^2 \) one needs the median levels of the two corresponding line arrangements only. As mentioned in the previous section, the relevant information about these levels can actually be obtained in linear time. Similarly, in a motion planning problem where the lines are considered as obstacles we are only interested in the cell of the arrangement we are located in. There is no way to ever reach any other cell, anyway.

This chapter is concerned with analyzing the complexity—that is, the number of vertices and edges—of a single cell in an arrangement of \( n \) curves in \( \mathbb{R}^2 \). In case of a line arrangement this is mildly interesting only: Every cell is convex and any line can appear at most once along the cell boundary. On the other hand, it is easy to construct an example in which there is a cell \( C \) such that every line appears on the boundary \( \partial C \).

But when we consider arrangement of line segments rather than lines, the situation changes in a surprising way. Certainly a single segment can appear several times along the boundary of a cell, see the example in Figure 12.1. Make a guess: What is the maximal complexity of a cell in an arrangement of \( n \) line segments in \( \mathbb{R}^2 \)?

Figure 12.1: A single cell in an arrangement of line segments.
You will find out the correct answer soon, although we will not prove it here. But my guess would be that it is rather unlikely that your guess is correct, unless, of course, you knew the answer already. :-) 

For a start we will focus on one particular cell of any arrangement that is very easy to describe: the lower envelope or, intuitively, everything that can be seen vertically from below. To analyze the complexity of lower envelopes we use a combinatorial description using strings with forbidden subsequences, so-called Davenport-Schinzel sequences. These sequences are of independent interest, as they appear in a number of combinatorial problems [2] and in the analysis of data structures [7]. The techniques used apply not only to lower envelopes but also to arbitrary cells of arrangements.

**Definition 12.1** An \((n, s)\)-Davenport-Schinzel sequence, for \(n, s \in \mathbb{N}\), is a sequence over an alphabet \(A\) of size \(n\) in which

- no two consecutive characters are the same and
- there is no alternating subsequence of the form \(\ldots a \ldots b \ldots a \ldots b \ldots\) of \(s + 2\) characters, for any \(a, b \in A\).

Let \(\lambda_s(n)\) be the length of a longest \((n, s)\)-Davenport-Schinzel sequence.

For example, \(abcbacb\) is a \((3, 4)\)-DS sequence but not a \((3, 3)\)-DS sequence because it contains the subsequence \(bcbcb\).

**Proposition 12.2** \(\lambda_s(m) + \lambda_s(n) \leq \lambda_s(m + n)\).

**Proof.** On the left hand side, we consider two Davenport-Schinzel sequences, one over an alphabet \(A\) of size \(m\) and another over an alphabet \(B\) of size \(n\). We may suppose that \(A \cap B = \emptyset\) (for each character \(x \in A \cap B\) introduce a new character \(x'\) and replace all occurrences of \(x\) in the second sequence by \(x'\) ). Concatenating both sequences yields a Davenport-Schinzel sequence over the alphabet \(A \cup B\) of size \(m + n\).

Let us now see how Davenport-Schinzel sequences are connected to lower envelopes. Consider a collection \(\mathcal{F} = \{f_1, \ldots, f_n\}\) of real-valued continuous functions that are defined on a common interval \(I \subset \mathbb{R}\). The lower envelope \(L_\mathcal{F}\) of \(\mathcal{F}\) is defined as the pointwise minimum of the functions \(f_i, 1 \leq i \leq n\), over \(I\). Suppose that any pair \(f_i, f_j, 1 \leq i < j \leq n\), intersects in at most \(s\) points. Then \(I\) can be decomposed into a finite sequence \(I_1, \ldots, I_\ell\) of (maximal connected) pieces on each of which a single function from \(\mathcal{F}\) defines \(L_\mathcal{F}\). Define the sequence \(\phi(\mathcal{F}) = (\phi_1, \ldots, \phi_\ell)\), where \(f_{\phi_i}\) is the function from \(\mathcal{F}\) which defines \(L_\mathcal{F}\) on \(I_i\).

**Observation 12.3** \(\phi(\mathcal{F})\) is an \((n, s)\)-Davenport-Schinzel sequence.

In the case of line segments the above statement does not hold because a set of line segments is in general not defined on a common real interval.
**Proposition 12.4** Let $F$ be a collection of $n$ real-valued continuous functions, each of which is defined on some real interval. If any two functions from $F$ intersect in at most $s$ points then $\phi(F)$ is an $(n, s+2)$-Davenport-Schinzel sequence.

**Proof.** Let us first argue that we may suppose that the functions in $F$ are piecewise quadratic. Denote by $P$ the set of points that are vertices of the arrangement induced by the graphs of the functions in $F$. In other words, $P$ is the set of all endpoints of functions and intersection points of two functions from $F$. Let $x_1, \ldots, x_m$ denote the sequence of $x$-coordinates of points from $P$, sorted in increasing order.

Consider the set $F_i \subseteq F$ of functions that are defined on the interval $[x_i, x_{i+1}]$, for $i \in \{1, \ldots, m-1\}$. By definition of $P$, no two functions intersect within the interval $(x_i, x_{i+1})$, that is, throughout $(x_i, x_{i+1})$ the functions from $F_i$ maintain the same total order (y-order). We describe how to replace each function in $F_i$ on $[x_i, x_{i+1}]$ by a quadratic function such that this order and therefore the combinatorics of the induced arrangement remains unchanged.

Consider a function $f \in F_i$. If $f$ is the only function in $F_i$ that passes through both points $(x_i, f(x_i))$ and $(x_{i+1}, f(x_{i+1}))$, then replace $f$ by the line segment connecting these two points. Otherwise, consider the set $\mathcal{F}$ of all functions in $F_i$ that pass through both points, and replace each function by a parabolic arc connecting the two points. These parabolic arcs can be chosen so that they are pairwise disjoint within the open interval $(x_i, x_{i+1})$ and all of them are infinitesimally close to the line segment between $(x_i, f(x_i))$ and $(x_{i+1}, f(x_{i+1}))$, while maintaining the total order among the functions in $\mathcal{F}$ within the interval $(x_i, x_{i+1})$. The described replacement does not introduce any new intersection points and therefore the combinatorics of the induced arrangement remains the same.

Now we are ready to prove the actual statement. The idea is to extend the functions to be defined on all of $\mathbb{R}$, without changing $\phi(F)$ too much, and then resort to Observation 12.3.

Consider any function $f \in F$ defined on an interval $[a, b]$. Extend $f$ to $\mathbb{R}$ using almost vertical rays pointing upward; from $a$ use a ray of sufficiently small slope, from $b$ use a ray of sufficiently large slope. For all functions use the same slope on these two extensions such that no extensions in the same direction intersect. By sufficiently small/large we mean that for any extension ray there is no endpoint of a function nor an intersection point of two functions in the open angular wedge bounded by the extension ray and the vertical ray starting from the same source. (There may be such points on the vertical ray, but not in the open wedge between the two rays.) As any two functions intersect only a finite number of times, there is only a finite number of endpoints and intersection points to consider, and so “sufficiently small/large” is well-defined.

Denote the resulting collection of functions totally defined on $\mathbb{R}$ by $F'$. If the extension rays are sufficiently close to vertical then $\phi(F') = \phi(F)$. (Recall that by our reasoning from above we may regard each function as a parabolic arc or a line segment locally.)

For any $f \in F'$ a single extension ray can create at most one additional intersection with any $g \in F'$. (Let $[a_f, b_f]$ and $[a_g, b_g]$ be the intervals on which the function $f$ and
was defined originally. Consider the ray $r$ extending $f$ from $a_f$ to the left. If $a_f \in [a_g, b_g]$ then $r$ may create a new intersection with $g$, if $a_f > b_g$ then $r$ creates a new intersection with the right extension of $g$ from $b_g$, and if $a_f < a_g$ then $r$ does not create any new intersection with $g$.

On the other hand, for any pair $s, t$ of segments, neither the left extension of the leftmost segment endpoint nor the right extension of the rightmost segment endpoint can introduce an additional intersection. Therefore, any pair of segments in $F'$ intersects at most $s + 2$ times and the claim follows.

Next we will give an upper bound on the length of Davenport-Schinzel sequences for small $s$.

\textbf{Lemma 12.5} $\lambda_1(n) = n$, $\lambda_2(n) = 2n - 1$, and $\lambda_3(n) \leq 2n(1 + \log n)$.

\textbf{Proof.} $\lambda_1(n) = n$ is obvious. $\lambda_2(n) = 2n - 1$ is given as an exercise. We prove $\lambda_3(n) \leq 2n(1 + \log n) = O(n \log n)$.

For $n = 1$ it is $\lambda_3(1) = 1 \leq 2$. For $n > 1$ consider any $(n, 3)$-DS sequence $\sigma$ of length $\lambda_3(n)$. Let $a$ be a character that appears least frequently in $\sigma$. Clearly $a$ appears at most $\lambda_3(n)/n$ times in $\sigma$. Delete all appearances of $a$ from $\sigma$ to obtain a sequence $\sigma'$ on $n - 1$ symbols. But $\sigma'$ is not necessarily a DS sequence because there may be consecutive appearances of a character $b$ in $\sigma'$, in case that $\sigma = \ldots a \ldots bab \ldots a \ldots$, in contradiction to $\sigma$ being an $(n, 3)$-DS sequence.

Claim: There are at most two pairs of consecutive appearances of the same character in $\sigma'$. Indeed, such a pair can be created around the first and last appearance of $a$ in $\sigma$ only. If any intermediate appearance of $a$ creates a pair $bb$ in $\sigma'$ then $\sigma = \ldots a \ldots bab \ldots a \ldots$, in contradiction to $\sigma$ being an $(n, 3)$-DS sequence.

Therefore, one can remove at most two characters from $\sigma'$ to obtain a $(n - 1, 3)$-DS sequence $\overline{\sigma}$. As the length of $\overline{\sigma}$ is bounded by $\lambda_3(n - 1)$, we obtain $\lambda_3(n) \leq \lambda_3(n - 1) + \lambda_3(n)/n + 2$. Reformulating yields

$$\frac{\lambda_3(n)}{n} \leq \frac{\lambda_3(n - 1)}{n - 1} + \frac{2}{n - 1} \leq \frac{1}{f(1)} + 2 \sum_{i=1}^{n-1} \frac{1}{i} = 1 + 2H_{n-1}$$

and together with $2H_{n-1} < 1 + 2\log n$ we obtain $\lambda_3(n) \leq 2n(1 + \log n)$. \qed
Bounds for higher-order Davenport-Schinzel sequences. As we have seen, \( \lambda_1(n) \) (no ab\( \alpha \)) and \( \lambda_2(n) \) (no ab\( \alpha \beta \alpha \)) are both linear in \( n \). It turns out that for \( s \geq 3 \), \( \lambda_s(n) \) is slightly superlinear in \( n \) (taking \( s \) fixed). The bounds are known almost exactly, and they involve the inverse Ackermann function \( \alpha(n) \), a function that grows extremely slowly.

To define the inverse Ackermann function, we first define a hierarchy of functions \( \alpha_1(n) \), \( \alpha_2(n) \), \( \alpha_3(n) \), \ldots where, for every fixed \( k \), \( \alpha_k(n) \) grows much more slowly than \( \alpha_{k-1}(n) \):

We first let \( \alpha_1(n) = \lfloor n/2 \rfloor \). Then, for each \( k \geq 2 \), we define \( \alpha_k(n) \) to be the number of times we must apply \( \alpha_{k-1} \), starting from \( n \), until we get a result not larger than 1. In other words, \( \alpha_k(n) \) is defined recursively by:

\[
\alpha_k(n) = \begin{cases} 
0, & \text{if } n \leq 1; \\
1 + \alpha_k(\alpha_{k-1}(n)), & \text{otherwise}.
\end{cases}
\]

Thus, \( \alpha_2(n) = \lceil \log_2 n \rceil \), and \( \alpha_3(n) = \log^* n \).

Now fix \( n \), and consider the sequence \( \alpha_1(n) \), \( \alpha_2(n) \), \( \alpha_3(n) \), \ldots. For every fixed \( n \), this sequence decreases rapidly until it settles at 3. We define \( \alpha(n) \) (the inverse Ackermann function) as the function that, given \( n \), returns the smallest \( k \) such that \( \alpha_k(n) \) is at most 3:

\[
\alpha(n) = \min\{k \mid \alpha_k(n) \leq 3\}.
\]

We leave as an exercise to show that for every fixed \( k \) we have \( \alpha_k(n) = o(\alpha_{k-1}(n)) \) and \( \alpha(n) = o(\alpha_k(n)) \).

Coming back to the bounds for Davenport-Schinzel sequences, for \( \lambda_3(n) \) (no ab\( \alpha \bar{a} \)) it is known that \( \lambda_3(n) = \Theta(n\alpha(n)) \) [4]. In fact it is known that \( \lambda_3(n) = 2n\alpha(n) \pm O(n\sqrt{\alpha(n)}) \) [5] [6]. For \( \lambda_4(n) \) (no ab\( \alpha \bar{a} \bar{b} \alpha \)) we have \( \lambda_4(n) = \Theta(n \cdot 2^{\alpha(n)}) \) [3].

For higher-order sequences the known upper and lower bounds are almost tight, and they are of the form \( \lambda_s(n) = n \cdot 2^{\text{poly}(\alpha(n))} \), where the degree of the polynomial in the exponent is roughly \( s/2 \) [3] [6].

Realizing DS sequences as lower envelopes. There exists a construction of a set of \( n \) segments in the plane whose lower-envelope sequence has length \( \Omega(n\alpha(n)) \). (In fact, the lower-envelope sequence has length \( n\alpha(n) - O(n) \), with a leading coefficient of 1; it is an open problem to get a leading coefficient of 2, or prove that this is not possible.)

It is an open problem to construct a set of \( n \) parabolic arcs in the plane whose lower-envelope sequence has length \( \Omega(n \cdot 2^{\alpha(n)}) \).

Generalizations of DS sequences. Also generalizations of Davenport-Schinzel sequences have been studied, for instance, where arbitrary subsequences (not necessarily an alternating pattern) are forbidden. For a word \( \sigma \) and \( n \in \mathbb{N} \) define \( \text{Ex}(\sigma, n) \) to be the maximum length of a word over \( A = \{1, \ldots, n\}^* \) that does not contain a subsequence of the form \( \sigma \). For example, \( \text{Ex}(ab\bar{a}b, n) = \lambda_3(n) \). If \( \sigma \) consists of two letters only, say \( a \)
and b, then $\text{Ex}(\sigma, n)$ is super-linear if and only if $\sigma$ contains $ababa$ as a subsequence \cite{1}. This highlights that the alternating forbidden pattern is of particular interest.

**Exercise 12.6** Prove that $\lambda_2(n) = 2n - 1$.

**Exercise 12.7** Prove that $\lambda_s(n)$ is finite for all $s$ and $n$.

## 12.1 Constructing lower envelopes

**Theorem 12.8** Let $\mathcal{F} = \{f_1, \ldots, f_n\}$ be a collection of real-valued continuous functions defined on a common interval $I \subset \mathbb{R}$ such that no two functions from $\mathcal{F}$ intersect in more than $s$ points. Then the lower envelope $L_\mathcal{F}$ can be constructed in $O(\lambda_s(n) \log n)$ time. (Assuming that an intersection between any two functions can be constructed in constant time.)

**Proof.** Divide and conquer. For simplicity, assume that $n$ is a power of two. Split $\mathcal{F}$ into two equal parts $\mathcal{F}_1$ and $\mathcal{F}_2$ and construct $L_{\mathcal{F}_1}$ and $L_{\mathcal{F}_2}$ recursively. The resulting envelopes can be merged using line sweep by processing $2\lambda_s(n/2) + \lambda_s(n)$ events (the inequality $2\lambda_s(n/2) \leq \lambda_s(n)$ is by Proposition 12.2). Here the first term accounts for events generated by the vertices of the two envelopes to be merged. The second term accounts for their intersections, each of which generates a vertex of the resulting envelope. Observe that no sorting is required and the SLS structure is of constant size. Therefore, the sweep can be done in time linear in the number of events.

This yields the following recursion for the runtime $T(n)$ of the algorithm. $T(n) \leq 2T(n/2) + c\lambda_s(n)$, for some constant $c \in \mathbb{N}$. Using Proposition 12.2 it follows that $T(n) \leq c \sum_{i=1}^{\log n} 2^i \lambda_s(n/2^i) \leq c \sum_{i=1}^{\log n} \lambda_s(n) = O(\lambda_s(n) \log n)$. \qed

**Exercise 12.9** Show that every $(n, s)$-Davenport-Schinzel sequence can be realized as the lower envelope of $n$ continuous functions from $\mathbb{R}$ to $\mathbb{R}$, every pair of which intersect at most $s$ times.

**Exercise 12.10** Show that every Davenport-Schinzel sequence of order two can be realized as a lower envelope of $n$ parabolas.

It is a direct consequence of the symmetry in the definition that the property of being a Davenport-Schinzel sequence is invariant under permutations of the alphabet. For instance, $\sigma = bcacba$ is a $(3, 3)$-DS sequence over $A = \{a, b, c\}$. Hence the permutation $\pi = (ab)$ induces a $(3, 3)$-DS sequence $\pi(\sigma) = acbca$ and similarly $\pi' = (cba)$ induces another $(3, 3)$-DS sequence $\pi'(\sigma) = acbca$.

When counting the number of Davenport-Schinzel sequences of a certain type we want to count essentially distinct sequences only. Therefore we call two sequences over a common alphabet $A$ equivalent if and only if one can be obtained from the other by a permutation of $A$. Then two sequences are distinct if and only if they are not
equivalent. A typical way to select a representative from each equivalence class is to order the alphabet and demand that the first appearance of a symbol in the sequence follows that order. For example, ordering $A = \{a, b, c\}$ alphabetically demands that the first occurrence of $a$ precedes the first occurrence of $b$, which in turn precedes the first occurrence of $c$.

**Exercise 12.11** Let $P$ be a convex polygon with $n + 1$ vertices. Find a bijection between the triangulations of $P$ and the set of pairwise distinct $(n, 2)$-Davenport-Schinzel sequences of maximum length $(2n - 1)$. It follows that the number of distinct maximum $(n, 2)$-Davenport-Schinzel sequences is exactly $C_{n-1} = \frac{1}{n} \binom{2n-2}{n-1}$, which is the $(n - 1)$-st Catalan number.

**Questions**

58. What is an $(n, s)$ Davenport-Schinzel sequence and how does it relate to the lower envelope of real-valued continuous functions? Give the precise definition and some examples. Explain the relationship to lower envelopes and how to apply the machinery to partial functions like line segments.

59. What is the value of $\lambda_1(n)$ and $\lambda_2(n)$?

60. What is the asymptotic value of $\lambda_3(n)$, $\lambda_4(n)$, and $\lambda_s(n)$ for larger $s$?

61. What is the combinatorial complexity of the lower envelope of a set of $n$ lines/parabolas/line segments?

**References**


Chapter 13

Epsilon Nets

13.1 Motivation

Here is our scenario for this chapter. We are given a set $A$ of points in $\mathbb{R}^d$ and a family $\mathcal{R}$ of ranges $r \subseteq \mathbb{R}^d$, for example the set of all balls, halfspaces, or convex sets in $\mathbb{R}^d$. $A$ is huge and probably not even known completely; similarly, $\mathcal{R}$ may not be accessible explicitly (in the examples above, it is an uncountable set). Still, we want to learn something about $A$ and some $r \in \mathcal{R}$.

The situation is familiar, definitely, if we don’t insist on the geometric setting. For example, let $A$ be the set of consumers living in Switzerland, and let $\bar{r}$ be the subset of consumers who frequently eat a certain food product, say Lindt chocolate. We have similar subsets for other food products, and together, they form the family of ranges $\mathcal{R}$.

If we want to learn something about $\bar{r}$, e.g. the ratio $|\bar{r}|/|A|$ (the fraction of consumers frequently eating Lindt chocolate), then we typically sample a subset $S$ of $A$ and see what portion of $S$ lies in $\bar{r}$. We want to believe that

$$\frac{|\bar{r} \cap S|}{|S|} \text{ approximates } \frac{|\bar{r}|}{|A|},$$

and statistics tells us to what extent this is justified. In fact, consumer surveys are based on this approach: in our example, $S$ is a sample of consumers who are being asked about their chocolate preferences. After this, the quantity $|\bar{r} \cap S|/|S|$ is known and used to predict the “popularity” $|\bar{r}|/|A|$ of Lindt chocolate among Swiss consumers.

In this chapter, we consider a different kind of approximation. Suppose that we are interested in the most popular food products in Switzerland, the ones which are frequently eaten by more than an $\varepsilon$-fraction of all consumers, for some fixed $0 \leq \varepsilon \leq 1$. The goal is to find a small subset $N$ of consumers that “represent” all popular products. Formally, we want to find a set $N \subseteq A$ such that

for all $r$: $\frac{|r|}{|A|} > \varepsilon \Rightarrow r \cap N \neq \emptyset$. 

Such a subset is called an *epsilon net*. Obviously, $N = A$ is an epsilon net for all $\epsilon$, but as already mentioned above, the point here is to have a *small* set $N$.

Epsilon nets are very useful in many contexts that we won’t discuss here. But already in the food consumption example above, it is clear that a small representative set of consumers is a good thing to have; for example if you quickly need a statement about a particular popular food product, you know that you will find somebody in your representative set who knows the product.

The material of this chapter is classic and goes back to Haussler and Welzl [1].

### 13.2 Range spaces and $\epsilon$-nets.

Here is the formal framework. Let $X$ be a (possibly infinite) set and $\mathcal{R} \subseteq 2^X$. The pair $(X, \mathcal{R})$ is called a *range space* with $X$ its *points* and the elements of $\mathcal{R}$ its *ranges*.

**Definition 13.1** Let $(X, \mathcal{R})$ be a range space. Given $A \subseteq X$, finite, and $\epsilon \in \mathbb{R}$, $0 \leq \epsilon \leq 1$, a subset $N$ of $A$ is called an $\epsilon$-net of $A$ (w.r.t. $\mathcal{R}$) if

$$\text{for all } r \in \mathcal{R}: \ |r \cap A| > \epsilon |A| \Rightarrow r \cap N \neq \emptyset .$$

This definition is easy to write down, but it is not so easy to grasp, and this is why we will go through a couple of examples below. Note that we have a slightly more general setup here, compared to the motivating Section 13.1 where we had $X = A$.

### 13.2.1 Examples

Typical examples of range spaces in our geometric context are

- $(\mathbb{R}, \mathcal{H}_1)$ with $\mathcal{H}_1 := \{(-\infty, a] \mid a \in \mathbb{R}\} \cup \{(a, \infty) \mid a \in \mathbb{R}\}$ (*half-infinite intervals*),

- $(\mathbb{R}, \mathcal{I})$ with $\mathcal{I} := \{[a, b] \mid a, b \in \mathbb{R}, a \leq b\}$ (*intervals*),

and higher-dimensional counter-parts

- $(\mathbb{R}^d, \mathcal{H}_d)$ with $\mathcal{H}_d$ the closed *halfspaces* in $\mathbb{R}^d$ bounded by hyperplanes,

- $(\mathbb{R}^d, \mathcal{B}_d)$ with $\mathcal{B}_d$ the closed *balls* in $\mathbb{R}^d$,

- $(\mathbb{R}^d, \mathcal{S}_d)$ with $\mathcal{S}_d$ the $d$-dimensional *simplices* in $\mathbb{R}^d$, and

- $(\mathbb{R}^d, \mathcal{C}_d)$ with $\mathcal{C}_d$ the *convex sets* in $\mathbb{R}^d$.

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1In order to avoid confusion: A range space is nothing else but a set system, sometimes also called hypergraph. It is the context, where we think of $X$ as points and $\mathcal{R}$ as ranges in some geometric ambient space, that suggests the name at hand.
\(\varepsilon\)-Nets w.r.t. \((\mathbb{R}, \mathcal{H}_1)\) are particularly simple to obtain. For \(A \subseteq \mathbb{R}\), \(N := \{\min A, \max A\}\) is an \(\varepsilon\)-net for every \(\varepsilon\)—it is even a 0-net. That is, there are \(\varepsilon\)-nets of size 2, independent from \(|A|\) and \(\varepsilon\).

The situation gets slightly more interesting for the range space \((\mathbb{R}, \mathcal{I})\) with intervals. Given \(\varepsilon\) and \(A\) with elements

\[a_1 < a_2 < \cdots < a_n,\]

we observe that an \(\varepsilon\)-net must contain at least one element from any contiguous sequence \(\{a_i, a_{i+1}, \ldots, a_{i+k-1}\}\) of \(k > \varepsilon n\) (i.e. \(k \geq \lceil \varepsilon n \rceil + 1\)) elements in \(A\). In fact, this is a necessary and sufficient condition for \(\varepsilon\)-nets w.r.t. intervals. Hence,

\[\{a_{\lceil \varepsilon n \rceil + 1}, a_{2\lceil \varepsilon n \rceil + 2}, \ldots\}\]

is an \(\varepsilon\)-net of size \(\left\lfloor \frac{n}{\lceil \varepsilon n \rceil + 1} \right\rfloor \leq \left\lfloor \frac{1}{\varepsilon} \right\rfloor - 1\). So while the size of the net depends now on \(\varepsilon\), it is still independent of \(|A|\).

### 13.2.2 No point in a large range.

Let us start with a simple exercise, showing that large ranges are easy to “catch”. Assume that \(|r \cap A| > \varepsilon |A|\) for some fixed \(r\) and \(\varepsilon\), \(0 \leq \varepsilon \leq 1\).

Now consider the set \(S\) obtained by drawing \(s\) elements uniformly at random from \(A\) (with replacement). We write

\[S \sim A^s,\]

indicating that \(S\) is chosen uniformly at random from the set \(A^s\) of \(s\)-element sequences over \(A\).

What is the probability that \(S \sim A^s\) fails to intersect with \(r\), i.e. \(S \cap r = \emptyset\)? For \(p := \frac{|r \cap A|}{|A|}\) (note \(p > \varepsilon\)) we get\(^3\)

\[\text{prob}(S \cap r = \emptyset) = (1 - p)^s < (1 - \varepsilon)^s \leq e^{-\varepsilon s}.\]

That is, if \(s = \frac{1}{\varepsilon}\) then this probability is at most \(e^{-1} \approx 0.368\), and if we choose \(s = \lambda\frac{1}{\varepsilon}\), then this probability decreases exponentially with \(\lambda\): It is at most \(e^{-\lambda}\).

For example, if \(|A| = 10000\) and \(|r \cap A| > 100\) (\(r\) contains more than 1% of the points in \(A\)), then a sample of 300 points is disjoint from \(r\) with probability at most \(e^{-3} \approx 0.05\).

\(^2\)The number \(L\) of elements in the set is the largest \(\ell\) such that \(\ell(\lceil \varepsilon n \rceil + 1) \leq n\), hence \(L = \left\lfloor \frac{n}{\lceil \varepsilon n \rceil + 1} \right\rfloor\). Since \(\lceil \varepsilon n \rceil + 1 > \varepsilon n\), we have \(\frac{n}{\lceil \varepsilon n \rceil + 1} < \frac{1}{\varepsilon}\), and so \(L < \frac{1}{\varepsilon}\), i.e. \(L \leq \left\lfloor \frac{1}{\varepsilon} \right\rfloor - 1\).

\(^3\)We make use of the inequality \(1 + x \leq e^x\) for all \(x \in \mathbb{R}\).
13.2.3 Smallest enclosing balls.

Here is a potential use of this for a geometric problem. Suppose \( A \) is a set of \( n \) points in \( \mathbb{R}^d \), and we want to compute the smallest enclosing ball of \( A \). In fact, we are willing to accept some mistake, in that, for some given \( \epsilon \), we want a small ball that contains all but at most \( \epsilon n \) points from \( A \). So let’s choose a sample \( S \) of \( \lambda \frac{1}{\epsilon} \) points drawn uniformly (with replacement) from \( A \) and compute the smallest enclosing ball \( B \) of \( S \). Now let \( r := \mathbb{R}^d \setminus B \), the complement of \( B \) in \( \mathbb{R}^d \), play the role of the range in the analysis above. Obviously \( r \cap S = \emptyset \), so it is unlikely that \( |r \cap A| > \epsilon |A| \), since—if so—the probability of \( S \cap r = \emptyset \) was at most \( e^{-\lambda} \).

It is important to understand that this was complete nonsense!

For the probabilistic analysis above we have to first choose \( r \) and then draw the sample—and not, as done in the smallest ball example, first draw the sample and then choose \( r \) based on the sample. That cannot possibly work, since we could always choose \( r \) simply as the complement \( \mathbb{R}^d \setminus S \)—then clearly \( r \cap S = \emptyset \) and \( |r \cap A| > \epsilon |A| \), unless \( |S| \geq (1 - \epsilon)|A| \).

While you hopefully agree on this, you might find the counterargument with \( r = \mathbb{R}^d \setminus S \) somewhat artificial, e.g. complements of balls cannot be that selective in ‘extracting’ points from \( A \). It is exactly the purpose of this chapter to understand to what extent this advocated intuition is justified or not.

13.3 Either almost all is needed or a constant suffices.

Let us reveal the spectrum of possibilities right away, although its proof will have to await some preparatory steps.

**Theorem 13.2** Let \((X, R)\) be an infinite range space. Then one of the following two statements holds.

1. For every \( n \in \mathbb{N} \) there is a set \( A_n \subseteq X \) with \( |A_n| = n \) such that for every \( \epsilon \), \( 0 \leq \epsilon \leq 1 \), an \( \epsilon \)-net must have size at least \((1 - \epsilon)n\).

2. There is a constant \( \delta \) depending on \((X, R)\), such that for every finite \( A \subseteq X \) and every \( \epsilon \), \( 0 < \epsilon \leq 1 \), there is an \( \epsilon \)-net of \( A \) w.r.t. \( R \) of size at most \( \frac{88}{\epsilon} \log_2 \frac{45}{\epsilon} \) (independent of the size of \( A \)).

That is, either we have always \( \epsilon \)-nets of size independent of \( |A| \), or we have to do the trivial thing, namely choosing all but \( \epsilon n \) points for an \( \epsilon \)-net. Obviously, the range spaces \((\mathbb{R}, \mathcal{H}_1)\) and \((\mathbb{R}, \mathcal{I})\) fall into category (2) of the theorem.

For an example for (1), consider \((\mathbb{R}^2, \mathcal{C}_2)\). For any \( n \in \mathbb{N} \), let \( A_n \) be a set of \( n \) points in convex position. For every \( N \subseteq A_n \), there is a range \( r \in \mathcal{C}_2 \), namely the convex hull of \( A_n \setminus N \), such that \( A_n \cap r = A_n \setminus N \) (hence, \( r \cap N = \emptyset \)); see Figure 13.1. Therefore,
N ⊆ A_n cannot be an ε-net of A_n w.r.t. C_2 if |A_n \ N| = n - |N| > εn. Consequently, an ε-net must contain at least n - εn = (1 - ε)n points.

Figure 13.1: If R consists of all convex sets in the plane, then only trivial epsilon nets exist: for every subset N (black points) of a set A_n in convex position, the range r = conv(A_n \ N) fails to intersect N.

So what distinguishes (R^2, C_2) from (R, H_1) and (R, I)? And which of the two cases applies to the many other range spaces we have listed above? Will all of this eventually tell us something about our attempt of computing a small ball covering all but at most εn out of n given points? This and more should be clear by the end of this chapter.

13.4 What makes the difference: VC-dimension

Given a range space (X, R) and A ⊆ X, we let

$$\mathcal{R}\mathcal{|}_A := \{r \cap A \mid r \in \mathcal{R}\},$$

the projection of R to A. Even if R is infinite, \(\mathcal{R}\mathcal{|}_A\) is always of size at most 2^n if A is an n-element set. The significance of projections in our context becomes clear if we rewrite Definition 13.1 in terms of projections: N ⊆ A is an ε-net if

for all r ∈ \(\mathcal{R}\mathcal{|}_A\): |r| > ε|A| \(\Rightarrow\) r ∩ N ≠ ∅.

All of a sudden, the conditions for an ε-net have become discrete, and they only depend on the finite range space (A, \(\mathcal{R}\mathcal{|}_A\)).

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4If we were satisfied with any abstract example for category (1), we could have taken (X, 2^X) for any infinite set X.
Note that, for $A$ a set of $n$ points in convex position in the plane, $|C_2|_A = 2^n$; we get every subset of $A$ by an intersection with a convex set (this is also the message of Figure 13.1). That is $|C_2|_A = 2^n$, the highest possible value.

For $A$ a set of $n$ points in $\mathbb{R}$, we can easily see that $|I|_X|_A| = (n+1)^2 = O(n^2)$. A similar argument shows that $|H_1|_X|_A| = 2n$. Now comes the crucial definition.

Definition 13.3 Given a range space $(X, \mathcal{R})$, a subset $A$ of $X$ is shattered by $\mathcal{R}$ if $|\mathcal{R}|_A = 2^{|A|}$. The VC-dimension $\text{VCdim}(X, \mathcal{R})$, is the cardinality (possibly infinite) of the largest subset of $X$ that is shattered by $\mathcal{R}$. If no set is shattered (i.e. not even the empty set which means that $\mathcal{R}$ is empty), we set the VC-dimension to $-1$.

We had just convinced ourselves that $(C_2, \mathcal{C}_2)$ has arbitrarily large sets that can be shattered. Therefore, $\text{VCdim}(C_2, \mathcal{C}_2) = \infty$.

Consider now $(\mathbb{R}, I)$. Two points $A = \{a, b\}$ can be shattered, since for each of the 4 subsets, $\emptyset$, $\{a\}$, $\{b\}$, and $\{a, b\}$, of $A$, there is an interval that generates that subset by intersection with $A$. However, for $A = \{a, b, c\}$ with $a < b < c$ there is no interval that contains $a$ and $c$ but not $b$. Hence, $\text{VCdim}(\mathbb{R}, I) = 2$.

Exercise 13.4 What is $\text{VCdim}(\mathbb{R}, H_1)$?

Exercise 13.5 Prove that if $\text{VCdim}(X, \mathcal{R}) = \infty$, then we are in case (1) of Theorem 13.2, meaning that only trivial epsilon nets always exist.

13.4.1 The size of projections for finite VC-dimension.

Here is the (for our purposes) most important consequence of finite VC dimension: there are only polynomially many ranges in every projection.

Lemma 13.6 (Sauer's Lemma) If $(X, \mathcal{R})$ is a range space of finite VC-dimension at most $\delta$, then

$$|\mathcal{R}|_A \leq \Phi_{\delta}(n) := \sum_{i=0}^{\delta} \binom{n}{i}$$

for all $A \subseteq X$ with $|A| = n$.

---

5Given $A$ as $a_1 < a_2 \cdots < a_n$ we can choose another $n+1$ points $b_i, 0 \leq i \leq n$, such that

$$b_0 < a_1 < b_2 < a_2 < b_3 < \cdots b_{n-1} < a_n < b_n.$$ Each nonempty intersection of $A$ with an interval can be uniquely written as $A \cap [b_i, b_j]$ for $0 \leq i < j \leq n$. This gives $\binom{n+1}{2}$ plus one for the empty set.

6"VC" in honor of the Russian statisticians V. N. Vapnik and A. Ya. Chervonenkis, who discovered the crucial role of this parameter in the late sixties.
13.4. What makes the difference: VC-dimension

Proof. First let us observe that \( \Phi : \mathbb{N}_0 \cup \{-1\} \times \mathbb{N}_0 \to \mathbb{N}_0 \) is defined by the recurrence\footnote{We recall that the binomial coefficients \( \binom{n}{k} \) (with \( k, n \in \mathbb{N}_0 \)) satisfy the recurrence \( \binom{n}{k} = 0 \) if \( n < k \), \( \binom{n}{0} = 1 \), and \( \binom{n}{k} = \binom{n-1}{k} + \binom{n-1}{k-1} \).}

\[
\Phi_{\delta}(n) = \begin{cases} 
0 & \delta = -1, \\
1 & n = 0 \text{ and } \delta \geq 0, \text{ and} \\
\Phi_{\delta}(n-1) + \Phi_{\delta-1}(n-1) & \text{otherwise.}
\end{cases}
\]

Second, we note that the VC-dimension cannot increase by passing from \((A, R)\) to a projection \((A', R)\), \(R := R|_A\). Hence, it suffices to consider the finite range space \((A, R)\)—which is of VC-dimension at most \(\delta\)—and show \(|R| \leq \Phi_\delta(n)\) (since \(\Phi\) is monotone in \(\delta\)).

Now we proceed to a proof by induction of this inequality. If \(A = \emptyset\) or \(R = \emptyset\) the statement is trivial. Otherwise, we consider the two ‘derived’ range spaces for some fixed \(x \in A\):

\[
(A \setminus \{x\}, R - x), \quad \text{with } R - x := \{r \setminus \{x\} \mid r \in R\}
\]

(note \(R - x = R|_{A \setminus \{x\}}\)) and

\[
(A \setminus \{x\}, R^{(x)}), \quad \text{with } R^{(x)} := \{r \in R \mid x \not\in r, r \cup \{x\} \in R\}.
\]

Observe that the ranges in \(R^{(x)}\) are exactly those ranges in \(R - x\) that have two preimages under the map \(R \ni r \mapsto r \setminus \{x\} \in R - x\), all other ranges have a unique preimage. Consequently,

\[
|R| = |R - x| + |R^{(x)}|.
\]

We have \(|R - x| \leq \Phi_\delta(n - 1)\). If \(A' \subseteq A \setminus \{x\}\) is shattered by \(R^{(x)}\), then \(A' \cup \{x\}\) is shattered by \(R\). Hence, \((A \setminus \{x\}, R^{(x)})\) has VC-dimension at most \(\delta - 1\) and \(|R^{(x)}| \leq \Phi_{\delta-1}(n - 1)\). Summing up, it follows that

\[
|R| \leq \Phi_\delta(n - 1) + \Phi_{\delta-1}(n - 1) = \Phi_\delta(n)
\]

which yields the assertion of the lemma. \(\square\)

In order to see that the bound given in the lemma is tight, consider the range space

\[
\left( X_i \bigcup_{i=0}^{\delta} \binom{X}{i} \right).
\]

Obviously, a set of more than \(\delta\) elements cannot be shattered (hence, the VC-dimension is at most \(\delta\)), and for any finite \(A \subseteq X\), the projection of the ranges to \(A\) is \(\bigcup_{i=0}^{\delta} \binom{A}{i}\)—with cardinality \(\Phi_{\delta}(|A|)\).
We note that a rough, but for our purposes good enough estimate for $\Phi$ is given by

$$\Phi_\delta(n) \leq n^\delta$$

for $\delta \geq 2$.

We have seen now that the maximum possible size of projections either grows exponentially ($2^n$ in case of infinite VC-dimension) or it is bounded by a polynomial $n^\delta$ in case of finite VC-dimension $\delta$). The latter is the key to the existence of small $\varepsilon$-nets. Before shedding light on this, let us better understand when the VC-dimension is finite.

## 13.5 VC-dimension of Geometric Range Spaces

### 13.5.1 Halfspaces.

Let us investigate the VC-dimension of $(\mathbb{R}^2, \mathcal{H}_2)$. It is easily seen that three points in the plane can be shattered by halfplanes, as long a they do not lie on a common line. Hence, the VC-dimension is at least 3. Now consider 4 points. If three of them lie on a common line, there is no way to separate the middle point on this line from the other two by a halfplane. So let us assume that no three points lie on a line. Either three of them are vertices of a triangle that contains the fourth point—then we cannot possibly separate the fourth point from the remaining three points by a halfplane. Or the four points are vertices of a convex quadrilateral—then there is no way of separating the endpoints from a diagonal from the other two points. Consequently, four points cannot be shattered, and $\text{VCdim}(\mathbb{R}^2, \mathcal{H}_2) = 3$ is established.

The above argument gets tedious in higher dimensions, if it works in a rigorous way at all. Fortunately, we can employ a classic: By Radon’s Lemma (Theorem 3.8), every set $A$ of $\geq d + 2$ points in $\mathbb{R}^d$ can be partitioned into two disjoint subsets $A_1$ and $A_2$ such that $\text{conv}(A_1) \cap \text{conv}(A_2) \neq \emptyset$. We get, as an easy implication, that a set $A$ of at least $d + 2$ points in $\mathbb{R}^d$ cannot be shattered by halfspaces. Indeed, let $A_1 \cup A_2$ be a partition as guaranteed by Radon’s Lemma. Now every halfspace containing $A_1$ must contain at least one point of $A_2$, hence $h \cap A = A_1$ is impossible for a halfspace $h$ and thus $A$ is not shattered by $\mathcal{H}_d$. Moreover, it is easily seen that the vertex set of a $d$-dimensional simplex (there are $d + 1$ vertices) can be shattered by halfspaces (each subset of the vertices forms a face of the simplex and can thus be separated from the rest by a hyperplane). We summarize that

$$\text{VCdim}(\mathbb{R}^d, \mathcal{H}_d) = d + 1.$$

Let us now consider the range space $(\mathbb{R}^d, \mathcal{H}_d)$, where $\mathcal{H}_d$ denotes the set of all closed halfspaces below non-vertical hyperplanes\footnote{A hyperplane is called non-vertical if it can be specified by a linear equation $\sum_{i=1}^d \lambda_i x_i = \lambda_{d+1}$ with $\lambda_d \neq 0$; see also Section 1.2.}—we call these lower halfspaces. Since $\mathcal{H}_d \subseteq

\begin{align*}
8 & \text{A better estimate, at least for } \delta \geq 3, \text{ is given by } \Phi_\delta(n) < \left(\frac{\varepsilon}{2}\right)^d \text{ for all } n, d \in \mathbb{N}, d \leq n. \\
9 & \text{A hyperplane is called non-vertical if it can be specified by a linear equation } \sum_{i=1}^d \lambda_i x_i = \lambda_{d+1} \text{ with } \lambda_d \neq 0; \text{ see also Section 1.2.}
\end{align*}
the VC-dimension of \((\mathbb{R}^d, \mathcal{H}_d)\) is at most \(d + 1\), but, in fact, it not too difficult to show
\[
\text{VCdim}(\mathbb{R}^d, \mathcal{H}_d) = d.
\]
(13.7)
(\text{Check this claim at least for } d = 2.) This range space is a geometric example where the bound of Sauer’s Lemma is attained. Indeed, for any set \(A\) of \(n\) points in \(\mathbb{R}^d\) in general position\(^{10}\), it can be shown that
\[
|\mathcal{H}_d|_A = \Phi_d(n).
\]

### 13.5.2 Balls.

It is easy to convince oneself that the VC-dimension of disks in the plane is 3: Three points not on a line can be shattered and four points cannot. Obviously not, if one of the points is in the convex hull of the other, and for four vertices of a convex quadrilateral, it is not possible for both diagonals to be separated from the endpoints of the respective other diagonal by a circle (if you try to draw a picture of this, you see that you get two circles that intersect four times, which we know is not be the case).

A more rigorous argument which works in all dimensions is looming with the help of (13.7), if we employ the following transformation called \textit{lifting map} that we have already encountered for \(d = 2\) in Section 6.3:
\[
\mathbb{R}^d \rightarrow \mathbb{R}^{d+1}
\]
\[
(x_1, x_2, \ldots, x_d) = p \mapsto \ell(p) = (x_1, x_2, \ldots, x_d, x_1^2 + x_2^2 + \cdots + x_d^2)
\]
(For a geometric interpretation, this is a vertical projection of \(\mathbb{R}^d\) to the unit paraboloid \(x_{d+1} = x_1^2 + x_2^2 + \cdots + x_d^2\) in \(\mathbb{R}^{d+1}\).) The remarkable property of this transformation is that it maps balls in \(\mathbb{R}^d\) to halfspaces in \(\mathbb{R}^{d+1}\) in the following sense.

Consider a ball \(B_d(c, \rho)\) \((c = (c_1, c_2, \ldots, c_d) \in \mathbb{R}^d\) the center, and \(\rho \in \mathbb{R}^+\) the radius). A point \(p = (x_1, x_2, \ldots, x_d)\) lies in this ball if and only if
\[
\sum_{i=1}^{d} (x_i - c_i)^2 \leq \rho^2 \iff \sum_{i=1}^{d} (x_i^2 - 2x_i c_i + c_i^2) \leq \rho^2
\]
\[
\iff \left( \sum_{i=1}^{d} (-2c_i)x_i \right) + (x_1^2 + x_2^2 + \cdots + x_d^2) \leq \rho^2 - \sum_{i=1}^{d} c_i^2
\]
this equivalently means that \(\ell(p)\) lies below the non-vertical hyperplane (in \(\mathbb{R}^{d+1}\))
\[
h = h(c, \rho) = \left\{ x \in \mathbb{R}^{d+1} \bigg| \sum_{i=1}^{d+1} h_i x_i = h_{d+2} \right\}
\]
with
\[
(h_1, h_2, \ldots, h_d, h_{d+1}, h_{d+2}) = \left( -2c_1, -2c_2, \ldots, -2c_d, 1, \rho^2 - \sum_{i=1}^{d} c_i^2 \right).
\]

\(^{10}\)No \(i + 2\) on a common \(i\)-flat for \(i \in \{1, 2, \ldots, d - 1\}\); in particular, no \(d + 1\) points on a common hyperplane.
It follows that a set $A \subseteq \mathbb{R}^d$ is shattered by $\mathcal{B}_d$ (the set of closed balls in $\mathbb{R}^d$) if and only if $\ell(A) := \{\ell(p) \mid p \in A\}$ is shattered by $\mathcal{H}_{d+1}$. Assuming (13.7), this readily yields

$$\text{VCdim}(\mathbb{R}^d, \mathcal{B}_d) = d + 1.$$  

The lifting map we have employed here is a special case of a more general paradigm called \textit{linearization} which maps non-linear conditions to linear conditions in higher dimensions.

We have clarified the VC-dimension for all examples of range spaces that we have listed in Section 13.2.1, except for the one involving simplices. The following exercise should help to provide an answer.

**Exercise 13.8** For a range space $(X, \mathcal{R})$ of VC-dimension $d$ and a number $k \in \mathbb{N}$, consider the range space $(X, I)$ where ranges form intersections of at most $k$ ranges from $\mathcal{R}$, that is, $I = \{\bigcap_{i=1}^{k} R_i \mid \forall i : R_i \in \mathcal{R}\}$. Give an upper bound for the VC-dimension of $(X, I)$.

### 13.6 Small $\varepsilon$-Nets, an Easy Warm-up Version

Let us prove a first bound on the size of $\varepsilon$-nets when the VC-dimension is finite.

**Theorem 13.9** Let $n \in \mathbb{N}$ and $(X, \mathcal{R})$ be a range space of VC-dimension $d \geq 2$. Then for any $A \subseteq X$ with $|A| = n$ and any $\varepsilon \in \mathbb{R}^+$ there exists an $\varepsilon$-net $N$ of $A$ w.r.t. $\mathcal{R}$ with $|N| \leq \lceil \frac{d \ln n}{\varepsilon} \rceil$.

**Proof.** We restrict our attention to the finite projected range space $(A, \mathcal{R})$, $\mathcal{R} := \mathcal{R}|_{A}$, for which we know $|\mathcal{R}| \leq \Phi_d(n) \leq n^d$. It suffices to show that there is a set $N \subseteq A$ with $|N| \leq \frac{d \ln n}{\varepsilon}$ which contains an element from each $r \in R_{\varepsilon} := \{r \in \mathcal{R} : |r| > \varepsilon n\}$.

Suppose, for some $s \in \mathbb{N}$ (to be determined), we let $N \sim A^s$. For each $r \in R_{\varepsilon}$, we know that $\text{prob}(r \cap N = \emptyset) < (1 - \varepsilon)^s \leq e^{-\varepsilon s}$. Therefore,

$$\text{prob}(N \text{ is not } \varepsilon\text{-net of } A) = \text{prob}(\exists r \in R_{\varepsilon} : r \cap N = \emptyset) \leq \text{prob}(\bigvee_{r \in R_{\varepsilon}} (r \cap N = \emptyset)) \leq \sum_{r \in R_{\varepsilon}} \text{prob}(r \cap N = \emptyset) < |R_{\varepsilon}| e^{-\varepsilon s} \leq n^d e^{-\varepsilon s}.$$  

It follows that if $s$ is chosen so that $n^d e^{-\varepsilon s} \leq 1$, then $\text{prob}(N \text{ is not } \varepsilon\text{-net of } A) < 1$ and there remains a positive probability for the event that $N$ is an $\varepsilon$-net of $A$. Now

$$n^d e^{-\varepsilon s} \leq 1 \iff n^d \leq e^{\varepsilon s} \iff d \ln n \leq \varepsilon s.$$  

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That is, for $s = \lceil \frac{d \ln n}{\varepsilon} \rceil$, the probability of obtaining an $\varepsilon$-net is positive, and therefore an $\varepsilon$-net of that size has to exist.\footnote{This line of argument “If an experiment produces a certain object with positive probability, then it has to exist”, as trivial as it is, admittedly needs some time to digest. It is called The Probabilistic Method, and was used and developed to an amazing extent by the famous Hungarian mathematician Paul Erdős starting in the thirties.}

If we are willing to invest a little more in the size of the random sample $N$, then the probability of being an $\varepsilon$-net grows dramatically. More specifically, for $s = \lceil \frac{d \ln n + \lambda}{\varepsilon} \rceil$, we have

$$n^d e^{-\varepsilon s} \leq n^d e^{-d \ln n - \lambda} = e^{-\lambda},$$

and, therefore, a sample of that size is an $\varepsilon$-net with probability at least $1 - e^{-\lambda}$.

We realize that we need $\frac{d \ln n}{\varepsilon}$ sample size to compensate for the (at most) $n^d$ subsets of $A$ which we have to hit—it suffices to ensure positive success probability. The extra $\frac{\lambda}{\varepsilon}$ allows us to boost the success probability.

Also note that if $A$ were shattered by $R$, then $R = 2^A$ and $|R| = 2^n$. Using this bound instead of $n^d$ in the proof above would require us to choose $s$ to be roughly $\frac{n \ln 2}{\varepsilon}$, a useless estimate which even exceeds $n$ unless $\varepsilon$ is large (at least $\ln 2 \approx 0.69$).

13.6.1 Smallest enclosing balls, again

It is time to rehabilitate ourselves a bit concerning the suggested procedure for computing a small ball containing all but at most $\varepsilon|A|$ points from an $n$-point set $A \subseteq \mathbb{R}^d$.

Let $(\mathbb{R}^d, B_d^{\text{compl}})$ be the range space whose ranges consist of all complements of closed balls in $\mathbb{R}^d$. This has the same VC-dimension $d + 1$ as $(\mathbb{R}^d, B_d)$. Indeed, if $A \cap r = A'$ for a ball $r$, then $A \cap (\mathbb{R}^d \setminus r) = A \setminus A'$, so $A$ is shattered by $B_d$ if and only if $A$ is shattered by $B_d^{\text{compl}}$.

Hence, if we choose a sample $N$ of size $s = \lceil \frac{(d + 1) \ln n + \lambda}{\varepsilon} \rceil$ then, with probability at least $1 - e^{-\lambda}$ this is an $\varepsilon$-net for $A$ w.r.t. $B_d^{\text{compl}}$. Let us quickly recall what this means: whenever the complement of some ball $B$ has empty intersection with $N$, then this complement contains at most an $\varepsilon|A|$ points of $A$. As a consequence, the smallest ball enclosing $N$ has at most $\varepsilon|A|$ points of $A$ outside, with probability at least $1 - e^{-\lambda}$.

13.7 Even Smaller $\varepsilon$-Nets

We still need to prove part 2 of Theorem 13.2, the existence of $\varepsilon$-nets whose size is independent of $A$. For this, we employ the same strategy as in the previous section, i.e. we sample elements from $A$ uniformly at random, with replacement; a refined analysis will show that—compared to the bound of Theorem 13.9—much less elements suffice. Here is the main technical lemma.
**Lemma 13.10** Let \((X, \mathcal{R})\) be a range space of VC-dimension \(\delta \geq 2\), and let \(A \subseteq X\) be finite. If \(x \sim A^m\) for \(m \geq 8/\varepsilon\), then for the set \(N_x\) of elements occurring in \(x\), we have

\[
\text{prob}(N_x \text{ is not an } \varepsilon\text{-net for } A \text{ w.r.t. } \mathcal{R}) \leq 2\Phi_\delta(2m)2^{-\frac{\varepsilon m}{2}}.
\]

Before we prove this, let us derive the bound of Theorem 13.2 from it. Let us recall the statement.

**Theorem 13.2 (2).** Let \((X, \mathcal{R})\) be a range space with VC-dimension \(\delta \geq 2\). Then for every finite \(A \subseteq X\) and every \(\varepsilon, 0 < \varepsilon \leq 1\), there is an \(\varepsilon\)-net of \(A\) w.r.t. \(\mathcal{R}\) of size at most \(8\delta \varepsilon \log_2 4\delta \varepsilon\) (independent of the size of \(A\)).

We want that

\[
2\Phi_\delta(2m)2^{-\frac{\varepsilon m}{2}} < 1,
\]

since then we know that an \(\varepsilon\)-net of size \(m\) exists. We have

\[
\begin{align*}
2\Phi_\delta(2m)2^{-\frac{\varepsilon m}{2}} &< 1 \\
\iff 2(2m)^\delta < 2^{\frac{\varepsilon m}{2}} \\
\iff 1 + \delta \log_2(2m) < \frac{\varepsilon m}{2} \\
\iff 2\delta \log_2 m < \frac{\varepsilon m}{2} \\
\iff \frac{4\delta}{\varepsilon} < \frac{m}{\log_2 m}.
\end{align*}
\]

In the second to last implication, we have used \(\delta \geq 2\) and \(m \geq 8\). Now we claim that the latter inequality is satisfied for \(m = 2\frac{4\delta}{\varepsilon} \log_2 4\delta \varepsilon\). To see this, we need that \(\frac{m}{\log_2 m} > \alpha\) for \(m = 2\alpha \log_2 \alpha\) and \(\alpha = \frac{4\delta}{\varepsilon}\). We compute

\[
\frac{m}{\log_2 m} = \frac{2\alpha \log_2 \alpha}{\log_2(2\alpha \log_2 \alpha)} = \frac{2\alpha \log_2 \alpha}{1 + \log_2 \alpha + \log_2 \log_2 \alpha} = \frac{2\log_2 \alpha}{1 + \log_2 \alpha + \log_2 \log_2 \alpha} \geq \alpha
\]

as long as

\[
\log_2 \alpha \geq 1 + \log_2 \log_2 \alpha \iff \log_2 \alpha \geq \log_2 2 \log_2 \alpha \iff \alpha \geq 2 \log_2 \alpha,
\]

which holds as long as \(\alpha \geq 2\), and this is satisfied for \(\alpha = \frac{4\delta}{\varepsilon}\).

**Proof.** (Lemma 13.10) By going to the projection \((A, \mathcal{R}|_A)\), we may assume that \(X = A\), so we have a finite range space over \(n\) elements (see Section 13.4). Fix \(\varepsilon \in \mathbb{R}^+\). For \(t \in \mathbb{N}\), a range \(r \in \mathcal{R}\) and \(x \in A^t\) (a \(t\)-vector of elements from \(A\)), we define

\[
\text{count}(r, x) = |\{i \in [t] : x_i \in r\}|.
\]
Now we consider two events over $A^{2m}$. The first one is the bad event of not getting an $\epsilon$-net when we choose $m$ elements at random from $A$ (plus another $m$ elements that we ignore for the moment): 

$$Q := \{ xy \in A^{2m} \mid x \in A^m, y \in A^m, \exists r \in R_\epsilon : \text{count}(r, x) = 0 \} .$$

Recall that $R_\epsilon = \{ r \in \mathbb{R} : |r| > \epsilon n \}$. Thus $\text{prob}(Q) = \text{prob}(N_x \text{ is not } \epsilon\text{-net})$ which is exactly what we want to bound.

The second auxiliary event looks somewhat weird at first: 

$$J := \{ xy \in A^{2m} \mid x \in A^m, y \in A^m, \exists r \in R_\epsilon : \text{count}(r, x) = 0 \text{ and } \text{count}(r, y) \geq \frac{\epsilon m}{2} \} .$$

This event satisfies $J \subseteq Q$ and contains pairs of sequences $x$ and $y$ with somewhat contradicting properties for some $r$. While the first part $x$ fails to contain any element from $r$, the second part $y$ has many elements from $r$.

**Claim 1.** $\text{prob}(J) \leq \text{prob}(Q) \leq 2 \text{prob}(J)$. 

The first inequality is a consequence of $J \subseteq Q$. To prove the second inequality, we show that $\text{prob}(J) / \text{prob}(Q) = \text{prob}(J \cap Q) / \text{prob}(Q) = \text{prob}(J \mid Q) \geq \frac{1}{2}$. 

So suppose that $xy \in Q$, with “witness” $r$, meaning that $r \in R_\epsilon$ and $\text{count}(r, x) = 0$. We show that $xy \in J$ with probability at least $1/2$, for every fixed such $x$ and $y$ chosen randomly from $A^m$. This entails the claim.

The random variable $\text{count}(r, y)$ is a sum of $m$ independent Bernoulli experiments with success probability $p := |r|/n > \epsilon$ (thus expectation $p$ and variance $p(1 - p)$). Using linearity of expectation and variance (the latter requires independence of the experiments), we get 

$$\text{E}(\text{count}(r, y)) = pm,$$

$$\text{Var}(\text{count}(r, y)) = p(1 - p)m.$$ 

Now we use Chebyshev’s inequality\textsuperscript{12} to bound the probability of the “bad” event that $\text{count}(r, y) < \frac{\epsilon m}{2}$. We have 

$$\text{prob} \left( \frac{\epsilon m}{2} \right) \leq \text{prob} \left( \frac{pm}{2} \right) \leq \text{prob} \left( \left| \text{count}(r, y) - \text{E}(\text{count}(r, y)) \right| > \frac{pm}{2} \right)$$

$$= \text{prob} \left( \left| \text{count}(r, y) - \text{E}(\text{count}(r, y)) \right| > \frac{1}{2} \text{Var}(\text{count}(r, y)) \right) \leq \frac{4(1 - p)^2}{p(1 - p)m} = \frac{4(1 - p)}{pm} \leq \frac{4}{\epsilon m} \leq \frac{1}{2},$$

\textsuperscript{12}prob(|X - E(X)| > kVar(X)) \leq \frac{k^2}{k^2 \text{Var}(X)}
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since \(m \geq \frac{8}{\varepsilon}\). Hence

\[
\text{prob}\left(\text{count}(r, y) \geq \frac{\varepsilon m}{2}\right) \geq \frac{1}{2},
\]

and the claim is proved.

Now the weird event \(J\) reveals its significance: we can nicely bound its probability. The idea is this: We enumerate \(A\) as \(A = \{a_1, a_2, \ldots, a_n\}\) and let the type of \(z \in A^n\) be the \(n\)-sequence

\[
\text{type}(z) = (\text{count}(a_1, z), \text{count}(a_2, z), \ldots, \text{count}(a_n, z)).
\]

For example, the type of \(z = (1, 2, 4, 2, 2, 4)\) w.r.t. \(A = \{1, 2, 3, 4\}\) is type \(z = (1, 3, 0, 2)\).

Now fix an arbitrary type \(\tau\). We will bound the conditional probability

\[
\text{prob}(xy \in J | xy \text{ has type } \tau) \leq \Phi_\delta(2m)2^{-\varepsilon m/2}.
\]

To analyze the probability in question, we need to sample \(z = xy\) uniformly at random from all sequences of type \(\tau\). This can be done as follows: take an arbitrary sequence \(z'\) of type \(\tau\), and then apply a random permutation \(\pi \in S_{2m}\) to obtain \(z = \pi(z')\), meaning

\[
z_i = z'_{\pi(i)} \text{ for all } i.
\]

Why does this work? First of all, \(\pi\) preserves the type, and it is easy to see that all sequences \(z\) of type \(\tau\) can be obtained in this way. Now we simply count the number of permutations that map \(z'\) to a fixed \(z\) and see that this number only depends on \(\tau\), so it is the same for all \(z\). Indeed, for every element \(a_i \in A\), there are \(\tau_i!\) many ways of mapping the \(a_i\)'s in \(z'\) to the \(a_i\)'s in \(z\). The number of permutations that map \(z'\) to \(z\) is therefore given by

\[
\prod_{i=1}^{n} \tau_i!.
\]

By these considerations,

\[
\text{prob}(xy \in J | xy \text{ has type } \tau) = \text{prob}(\pi(z') \in J).
\]

To estimate this, we let \(S\) be the set of distinct elements in \(z'\) (this is also a function of the type \(\tau\)). Since \((X, \mathcal{R})\) has VC-dimension \(\delta\), we know from Lemma 13.6 that \(|\mathcal{R}|_\delta \leq \Phi_\delta(2m)|\), so at most that many different subsets \(T\) of \(S\) can be obtained by intersections with ranges \(r \in \mathcal{R}\), and in particular with ranges \(r \in \mathcal{R}_\varepsilon\).

Now we look at some fixed such \(T\), consider a permutation \(\pi\) and write \(\pi(z') = xy\). We call \(T\) a witness for \(\pi\) if no element of \(x\) is in \(T\), but at least \(\varepsilon m/2\) elements of \(y\) are in \(T\). According to this definition,

\[
\pi(z') \in J \iff \pi \text{ has some witness } T \subseteq S.
\]

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By the union bound,
\[
\operatorname{prob}(\pi(z') \in J) = \operatorname{prob}(\exists T: T \text{ is a witness for } \pi) \leq \sum_T \operatorname{prob}(T \text{ is a witness for } \pi).
\]

Suppose that \(z'\) contains \(\ell \geq \epsilon m/2\) occurrences of elements of \(T\) (for smaller \(\ell\), \(T\) cannot be a witness). In order for \(\pi(z') = xy\) to be in \(J\), no element of \(T\) can appear in the first half \(x\), but all \(\ell\) occurrences of elements from \(T\) must fall into the second half \(y\). Therefore, the probability of \(T\) being a witness for a random permutation is
\[
\frac{\binom{m}{\ell}}{\binom{2m}{\ell}} = \frac{m(m - 1) \ldots (m - \ell + 1)}{2m(2m - 1) \ldots (2m - \ell + 1)} \leq 2^{-\ell} \leq 2^{-\frac{\epsilon m}{2}},
\]
since among all the \(\binom{2m}{\ell}\) equally likely ways in which \(\pi\) distributes the \(\ell\) occurrences in \(z\), exactly the \(\binom{m}{\ell}\) equally likely ways of putting them into \(y\) are good.\(^{13}\) Summing up over all at most \(\Phi(2m)\) sets \(T\), we get
\[
\operatorname{prob}(xy \in J | \ xy \text{ has type } \tau) \leq \Phi(2m)2^{-\frac{\epsilon m}{2}},
\]
for all \(\tau\).

The proof is finished by combining the two claims:
\[
\operatorname{prob}(N_x \text{ is not } \epsilon\text{-net}) = \operatorname{prob}(Q) \leq 2\operatorname{prob}(J) \leq 2\Phi(2m)2^{-\frac{\epsilon m}{2}}.
\]

\(\square\)

Questions

62. What is a range space? Give a definition and a few examples.
63. What is an epsilon net? Provide a formal definition, and explain in words what it means.
64. What is the VC dimension of a range space? Give a definition, and compute the VC dimension of a few example range spaces.
65. Which range spaces always have small epsilon nets (and how small), and which ones don't? Explain Theorem \[13.2\]
66. How can you compute small epsilon nets? Explain the general idea, and give the analysis behind the weaker bound of Theorem \[13.9\]
67. How can you compute smaller epsilon nets? Sketch the main steps of the proof of the stronger bound of Theorem \[13.2\]

\(^{13}\)This argument can be made more formal by explicitly counting the permutations that map \(\{1, 2, \ldots, \ell\}\) to the set \(\{1, 2, \ldots, m\}\). We leave this to the reader.
References