



On simplifying dot maps [☆]

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Abstract

Dot maps—drawings of point sets—are a well known cartographic method to visualize density functions over an area. We study the problem of simplifying a given dot map: given a set P of points in the plane, we want to compute a smaller set Q of points whose distribution approximates the distribution of the original set P .

We formalize this using the concept of ε -approximations, and we give efficient algorithms for computing the approximation error of a set Q of m points with respect to a set P of n points (with $m \leq n$) for certain families of ranges, namely unit squares, arbitrary squares, and arbitrary rectangles.

If the family \mathcal{R} of ranges is the family of all possible unit squares, then we compute the approximation error of Q with respect to P in $O(n \log n)$ time. If \mathcal{R} is the family of all possible rectangles, we present an $O(mn \log n)$ time algorithm. If \mathcal{R} is the family of all possible squares, then we present a simple $O(m^2n + n \log n)$ algorithm and an $O(n^2 \sqrt{n} \log n)$ time algorithm which is more efficient in the worst case.

Finally, we develop heuristics to compute good approximations, and we evaluate our heuristics experimentally.
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1. Introduction

An important component in the area of cartography is the ability to represent and visualize the distribution or density of some phenomenon such as the population distribution over a certain region. The most common technique to achieve this is the *dot map*, as shown in Fig. 1. The term *dot map* is self-explanatory—it refers to the use of dots or points placed on a map to represent a given distribution.

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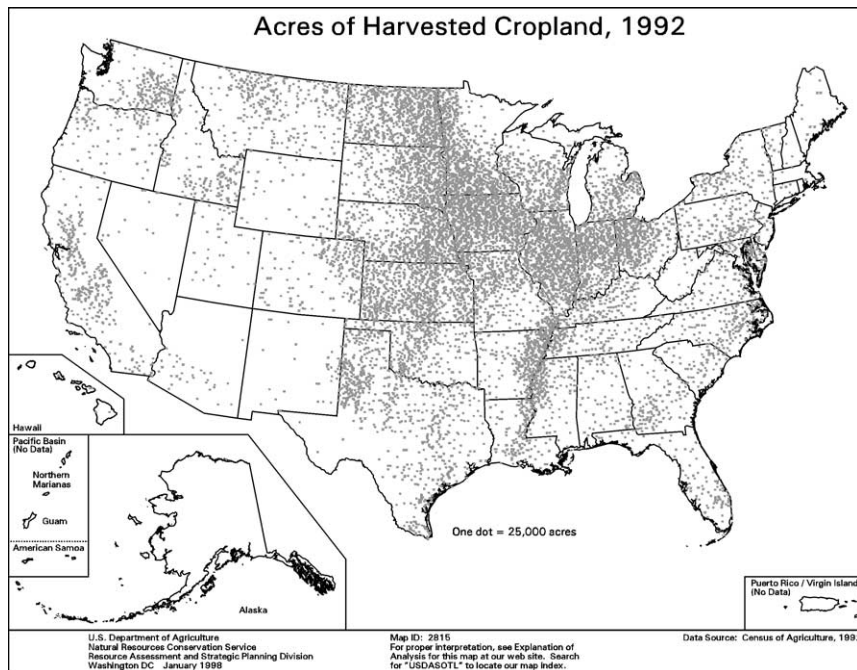


Fig. 1. Example of a dot map.

Dot maps are quite important and their use has been extensively studied in cartography—see for instance Chapter 8 of the book by Dent [6].

There are many issues involved in the use of dot maps as a tool for representing distributions. For example, the radius of the dots used, or the decision to allow or disallow dots to overlap are important visual considerations [6]. Depending on the application, it can also be important to take the topographic ‘background map’ into account: a dot map representing population density should not have dots inside lakes, in mountainous areas one may have to take altitude into account, and it may be important to ensure that dots are on the correct side of borders or other features [11]. In this paper, we concentrate on two related computational issues that purely deal with distribution issues; visual considerations and adherence to a background map are beyond the scope of this paper.

The first question we study is: Given a point set P representing a certain distribution, how can we automatically simplify it, that is, generate a smaller representative point set Q of a given size? This question arises when one wishes to scale a map: the number of points in the map has to decrease when the size of the map is decreased, otherwise it would become too cluttered. It may also arise in the generation of the initial dot map: “The printed dot map of the population distribution should be constructed at a larger scale based on more detailed information such as settlements and houses and then reduced to the final scale”, as Ditz [11] writes. The first question—How can we compute a good approximation?—immediately leads to the second: Given sets P and Q , how can we determine the quality of Q as an approximation to P ? To determine the quality of an approximation, we need a quantitative measure of similarity between dot maps. Our measure is inspired by interactive GISs where a user can use a dot map of, say, the population density, to estimate the population within a region [11]. This can either be a user-defined area—a square, for example—or a geographically meaningful region such as the area

within a certain distance from a river. This leads us to propose the notion of ε -approximations [17] as a quantitative measure of the quality of an approximation. A set Q of m points is called an ε -approximation of a set P of n points¹ with respect to a family \mathcal{R} of ranges, if for any range $r \in \mathcal{R}$ we have

$$||r \cap P|/n - |r \cap Q|/m| \leq \varepsilon.$$

In other words, if we approximate the number of points from P inside a range r by multiplying the number of points from Q inside the range by n/m , then we make an error of at most εn . This leads us to define $\Delta_{\mathcal{R}}(Q, P)$, the *approximation error of Q with respect to P* , for a family \mathcal{R} of ranges, as

$$\Delta_{\mathcal{R}}(Q, P) = \max_{r \in \mathcal{R}} ||r \cap P| - (n/m) \cdot |r \cap Q||.$$

The value n/m , which can be viewed as the weight of a point in Q as compared to a point in P , is called the *dot value* of the points in Q . We usually denote it by δ . In this paper we focus on squares and rectangles² as ranges. Of these types of ranges, squares are probably most natural in our application. Another natural range to consider would be discs.

1.1. Related work

ε -Approximations have been studied and used extensively in computational geometry—see for instance Chazelle’s book [5]—and various algorithms are known to compute ε -approximations of asymptotically optimal size for a set P and a given value of ε . Note that we want to solve a slightly different problem: in our case ε is not given, but the desired number of points in the approximation Q . Still, one may use the same type of algorithms. For instance, in many cases it turns out that random sampling is expected to produce an approximation of asymptotically optimal size. (One caveat is in place here: the optimality here refers to the worst-case size of an ε -approximation over all point sets P of n points, not to the minimum size needed for the given set P . These two sizes need not be the same.) Thus, for our problem we could simply take a random subset $Q \subset P$ of the desired size. Then, of course, one would want to check how good the sample is, that is, one needs an algorithm to compute the approximation error of given sets P and Q .

The use of ε -approximations to measure the similarity of two point sets is related to some statistical methods to derive a (continuous) density function from a given point set; see the book by Baily and Gatrell [2] for more information on statistical methods for spatial data analysis. For example, kernel estimation defines the density $\lambda(x)$ at a point x in the plane by summing the number of points within a region around the point x in a weighted manner; the shape of the region and the exact weighting scheme depends on the kernel used. Comparing two point sets—for example, to see whether the distribution of some feature of the population (number of cancer deaths, for instance) deviates from the population distribution itself—is then done by comparing the density functions $\lambda_1(x)$ and $\lambda_2(x)$ obtained for the two points sets. Usually one takes the quotient of these two values, but if one wants to bound the worst-case error this doesn’t work ($\lambda_2(x)$ may be zero) and one could take the absolute difference. The notion of ε -approximation with unit squares as ranges can be seen as a special case of this, where the kernel is a block function with value 1 inside the unit square centered at the point x and value 0 elsewhere (and

¹ Traditionally, in the definition of ε -approximation it is required that $Q \subset P$, but this is not necessary.

² In this paper squares and rectangles are always axis-parallel.

the other parameters of the kernel estimation method chosen suitably). The advantage of such a simple kernel function is that it is computationally easier, in the sense that it makes computing the approximation error easier. Recall that the motivation behind our use of ε -approximations is that we want to bound the maximum error when an approximating set Q is used to estimate the number of points from a set P inside a range. The size of such a range is not fixed in an interactive setting. Hence, we also look at squares of arbitrary sizes, which makes our error measure different from traditional kernel methods.

The approximation error as defined above is a generalization of the *bichromatic discrepancy* (or *combinatorial discrepancy*). Here one colors each point of a given set either red or blue and one is interested in the maximum difference, over all possible ranges of the given family, between the number of red points and the number of blue points inside a range. If we call the red point set P and the blue point set Q , and we define the dot value to be 1 (even when $|P| \neq |Q|$), then the bichromatic discrepancy equals the approximation error. Also, finding an optimal red-blue coloring of a given set P is identical to finding a subset $Q \subset P$ such that the discrepancy of Q with respect to P and dot value 2 is minimized. The concept of bichromatic discrepancy arises in computational learning theory, in particular in the so-called minimizing disagreement problem in agnostic PAC-learning [8,12]. Thus our algorithms to compute the approximation error of two given sets with respect to a family \mathcal{R} of ranges may be used to solve the minimizing disagreement problem when the class of hypotheses is \mathcal{R} —see the paper by Dobkin et al. [9] for details.

Finally, we note that our problem is related to that of computing the *area discrepancy* (or *continuous discrepancy*) of a point set P . This is a measure of how uniform that point set is, and it has applications in computer graphics [7,9,16].

1.2. Our results

Computing the approximation error of a set Q of m points with respect to a set P of n points, with $m \leq n$, is the topic of Section 2. We obtain the following results. If \mathcal{R} is the family of all possible unit squares, then we can compute the approximation error of Q with respect to P in $O(n \log n)$ time. If \mathcal{R} is the family of all possible rectangles, then we present two algorithms, a simple $O(m^2n + n \log n)$ algorithm and a more efficient $O(mn \log n)$ time algorithm. This is a slight improvement over an algorithm of Dobkin et al. [9] when m is $o(n)$. Their algorithm runs in $O(n^2 \log n)$ time regardless of how small m is. If \mathcal{R} is the family of all possible squares, then we present a simple $O(m^2n + n \log n)$ algorithm and an $O(n^2 \sqrt{n} \log n)$ time algorithm which is more efficient in the worst case.

We turn our attention in Section 3 to the experimental component of the paper. The goal is to develop heuristics to generate for a given set P an approximation Q of the desired size with as small an error as possible. We concentrate on the case of square ranges, as this seems most relevant to our application. Our heuristics use as a subroutine an algorithm to compute the error for given P and Q . Unfortunately, our algorithm for arbitrary squares is rather slow, and some of the heuristics call this subroutine many times. Hence, we first show experimentally that the exact error with respect to squares can be approximated well by computing the error with respect to fixed-size squares for a number of different sizes. After having established this, we compare various heuristics to find a good approximation of a given point set P . One of our findings is that taking the best approximation out of a large collection of random samples does not work so well, even though random sampling is guaranteed to find approximations that are asymptotically worst-case optimal.

2. Computing the approximation error

Let P be a set of n points and Q be a set of m points in the plane, with $m \leq n$. In this section we show how to compute the approximation error of Q with respect to P for three different families of ranges: unit squares, arbitrarily sized squares, and arbitrarily sized rectangles. By $\delta := n/m$ we denote the dot value of the points in Q .

2.1. Unit squares as ranges

Let \mathcal{R} be the family of all possible unit squares. When we want to compute the approximation error of Q with respect to P for unit squares, it can make a difference whether we consider open or closed squares. In the description of the algorithm, we will consider the squares to be closed; it is easy to adapt the algorithm to the case of open squares.

Recall that we use the absolute value of the error in the definition of approximation error. It is convenient to compute separately the maximum positive error and the maximum negative error. Below we describe how to compute the maximum positive error; computing the maximum negative error can be done in a similar way.

A unit square contains a point if and only if the center of the unit square is contained in the unit square centered at the point. Hence, instead of considering the point sets P and Q and the family of all unit squares as ranges, we can use the sets S_P and S_Q of unit squares centered at the points in P and Q , and all points in the plane as ranges. Call the squares in S_P the *red squares*, and the squares in S_Q the *blue squares*. The (positive) approximation error of a point x in the plane is now

$$(\# \text{ of red squares containing } x) - \delta \cdot (\# \text{ of blue squares containing } x).$$

The approximation error of S_Q with respect to S_P is the maximum approximation error over all points in the plane. From the discussion above it follows that this is the same as the approximation error of Q with respect to P for the family \mathcal{R} of unit squares as ranges.

The arrangement formed by the squares S_Q and S_P partitions the plane into faces where the approximation error of any point in a face of the arrangement is the same. Therefore, finding the maximum approximation error amounts to finding the face with maximum approximation error. We compute the approximation of S_Q with respect to S_P with a plane-sweep algorithm. In this algorithm, we sweep a vertical line ℓ from left to right over the arrangement. As we sweep the arrangement, we maintain the maximum approximation error over the faces of the arrangement intersected by ℓ . Since the arrangement is formed by squares, the only events are when the sweep lines reaches a left or right edge of a square. At each event we compute the maximum error of all points on ℓ and of all points slightly to the right of ℓ (but to the left of the previous event). The maximum error found in all the events will be the maximum error of S_Q with respect to S_P . We now describe the information we maintain during the sweep—the status structure—and how to handle the events.

2.1.1. A dynamic 1-dimensional structure

The status structure is a dynamic data structure for solving the following 1-dimensional version of the problem. We are given a set I_R of red segments and a set I_B of blue segments on the real line, and a parameter δ . The (positive) approximation error of a point $x \in \mathbb{R}$ is defined as

$$(\# \text{ of red segments containing } x) - \delta \cdot (\# \text{ of blue segments containing } x).$$

We want to maintain the maximum error over all points in \mathbb{R} under insertions and deletions of segments.

The structure we use is essentially the structure described in [4] in the context of grid placement problems. A similar structure is also presented in [9]. The structure maintains a function $f: \mathbb{R} \rightarrow \mathbb{R}$. Initially, it is assumed that $f(x) = 0$, for all $x \in \mathbb{R}$. The following update and query operations are allowed on the structure:

- (1) Insert($[a : b], d$): Increase the value of $f(x)$ by d over the interval $[a : b]$.
- (2) Delete($[a : b], d$): Decrease the value of $f(x)$ by d over the interval $[a : b]$.
- (3) Max(): Return $\max\{f(x): x \in \mathbb{R}\}$.

The first two operations can be performed in $O(\log n)$ time where n is the number of intervals currently inserted and the third operation takes $O(1)$ time. Essentially, the data structure is a balanced binary tree (similar to a segment tree [3]) whose leaves represent the *elementary intervals* (of the inserted intervals) ordered from left to right. An internal node of the tree represents the interval that is the union of the elementary intervals of the leaves in its subtree. The nodes have been augmented with additional information in order to answer the queries. The structure has $O(n)$ size where n is the number of intervals currently in the structure. For more details on the structure, the reader is referred to the paper of Bose et al. [4].

With this structure, the 1-dimensional problem is easily solved. When inserting (respectively deleting) a red segment, we increase (respectively decrease) the value of $f(x)$ by 1 over this segment. Similarly, when inserting (respectively deleting) a blue segment, we decrease (respectively increase) the value of $f(x)$ by δ over this segment. Max() allows one to recover the maximum approximation error over the currently inserted segments.

This leads to the following lemma.

Lemma 2.1. *The maximum approximation error of a set of red and blue segments on a line can be maintained with a structure of $O(n)$ space that takes $O(\log n)$ time per insertion and deletion, where n is the number of red and blue segments.*

We now return to the 2-dimensional problem, where we want to compute the approximation error of a set of blue squares with respect to a set of red squares, with the points in the plane as ranges. Recall that our approach is a plane-sweep algorithm. The algorithm maintains the maximum error along the sweep line ℓ using the structure \mathcal{T} just described above. Whenever the left edge of a square is encountered, we insert its y -interval into the structure along with the appropriate value (that is, 1 if it is red and $-\delta$ otherwise), and whenever the right edge of a square is encountered, we delete its y -interval. If events happen simultaneously—multiple edges have the same x -coordinate—then we process the events in the following order. First we handle all left boundaries. After this, Max() tells us the maximum error on ℓ . Next, we handle all the right boundaries, and get the maximum error slightly to the right of ℓ . Hence, every event takes $O(\log n)$ time to process and the initialization takes $O(n \log n)$ time. Since there are $O(n)$ events to process, we get the following theorem.

Theorem 2.2. *Let P be a set of n points in the plane, and let Q be a set of m points in the plane, with $m \leq n$. The approximation error of Q with respect to P for the family of all unit squares can be computed in $O(n \log n)$ time.*

2.2. Arbitrarily sized squares as ranges

The case of squares of arbitrary size as ranges is probably the most interesting in our application. Note that, unlike in the case of unit squares, the approximation error does not depend on whether we consider open or closed squares: for any open (closed) square, there is a slightly smaller closed (larger open) square that contains exactly the same points. We start by showing a fairly simple algorithm that runs in $O(m^2n + n \log n)$ time.

We first prove a lemma which restricts the number of candidate squares. Let B be the bounding box of $P \cup Q$.

Lemma 2.3. *There is an open square with maximum positive error such that two opposite sides of the square each either contain a point from Q or are contained in the boundary of B . Similarly, there is a closed square with maximum negative error such that two opposite sides of the square each contain a point from Q .*

Proof. Let s be an open square of maximum positive error, that is, a square that maximizes $|r \cap P| - \delta \cdot |r \cap Q|$. Suppose the top and right edge do not contain a point from Q or a part of the boundary of B . Fix the bottom left corner of s and grow the square until either a point from Q or the boundary of B hits the top or right edge of s . No point of P can enter s during this process since otherwise s was not a maximum. Next, fix the top right corner of s and grow the square until either a point from Q or the boundary of B hits the bottom or left edge of s . Again, no point of P can enter s during this process. At this point, the conditions of the lemma are met or two adjacent edges of s contain a point from Q or the boundary of B . If the latter holds, then assume, without loss of generality, that the top and left edges of s contain a point of Q . Then fix the top left corner and grow the square until the condition of the lemma is met.

Now let s be a closed square of maximum negative error, and suppose the top and right edge do not contain a point from Q or a part of the boundary of B . We can transform s into a square with the same error that satisfies the conditions of the lemma using the same procedure as above, except this time we shrink s instead of growing s . Because of the shrinking, we do not have the case where the boundary of s hits the boundary of B , since we can assume that initially s lies completely inside B . \square

2.2.1. A simple algorithm

Next we describe a simple algorithm, based on Lemma 2.3, to compute the maximum positive approximation error; the maximum negative error can be computed in a similar way.

By Lemma 2.3, the square of maximum discrepancy must have a blue point (i.e., a point from Q) on two opposite sides. Given two blue points, if the absolute value of the difference in their x -coordinate is larger than the absolute value of the y -coordinate difference, then the two points can only lie on the left and right sides of a square. Similarly, if the y -coordinate difference is larger, than the two points can only lie on the top and bottom edges of the square. Finally, if the differences are the same, then there is a unique square with the points at the opposite corners. This implies that a given pair of blue points determines the size of the square and the direction of search. Since there are m blue points, there are $\binom{m}{2}$ candidate pairs. Select one such pair, q_i, q_j , and assume without loss of generality that the y -coordinate difference is larger. The case where the x -coordinate difference is larger is symmetric, and the other case is trivial.

Given q_i and q_j , let h_i and h_j be the horizontal lines through the respective points. We have to find the maximum error over all squares whose top and bottom edges are contained in those lines. In order to find this maximum, we will sweep (i.e., move) the square from left to right through the strip.

Consider the points in $P \cup Q$ that lie within this strip. Sort these points by their x -coordinates, and let S represent this set in sorted order. Start with the left boundary of the square on the left boundary of B . Compute the discrepancy of this square by finding the points of S in this square. Now, sweep the square from left to right until the right boundary reaches the right boundary of B and maintain the maximum at each step. The events in this sweep are that either a point leaves the square or a point enters the square. The order in which the points enter as well as the order in which the points leave is the sorted order. Processing an event amounts to adding or subtracting the appropriate amount to the current discrepancy, depending on which point enters or leaves. (Events that occur simultaneously should be handled together; the details of how this should be done are easy to fill in.) Note that we do not need to sweep the whole strip but only the portion of the strip where q_i and q_j are on the top and bottom edges of the square. However, this optimization does not make a difference asymptotically. Since each event can be processed in $O(1)$ time given the sorted order, we can compute in $O(n)$ time the maximum discrepancy given a candidate pair of points provided the points in the strip are sorted. If we pre-sort the points of $P \cup Q$ in $O(n \log n)$ time then $O(n)$ time the sorted order of points of $P \cup Q$ within a strip can be obtained. Since there are $O(m^2)$ possible candidates and each candidate can be verified in $O(n)$ time, the total time for the algorithm is $O(m^2n + n \log n)$.

Theorem 2.4. *Let P be a set of n points in the plane, and let Q be a set of m points in the plane, with $m \leq n$. The approximation error of P with respect to Q for squares can be computed in $O(m^2n + n \log n)$ time.*

2.2.2. A subcubic algorithm

The algorithm of Theorem 2.4 runs in cubic time if $m = \Theta(n)$. Next we describe a subcubic algorithm. Recall that in the case of unit squares as ranges, we replaced every point in $P \cup Q$ with a unit square and we looked at the maximum error of points in the plane with respect to the resulting sets S_P and S_Q of squares. This approach does not apply here since the size of the square is not fixed. However, we can do something similar: replace every point in $P \cup Q$ with a square of size ρ , let ρ grow from zero to infinity, and maintain the maximum error of points in the plane with respect to the resulting sets S_P and S_Q of squares over the whole growing process. Notice that the maximum error only changes when two squares start overlapping. This is precisely when new faces in the arrangement of squares appear and other faces disappear. This happens $O(n^2)$ times. The approach is to maintain the maximum over the whole growing process. In the remainder we develop a structure that allows us to compute in $O(\sqrt{n} \log n)$ time the new maximum discrepancy when such an event takes place, leading to an algorithm with overall running time of $O(n^2 \sqrt{n} \log n)$.

A dynamic 2-dimensional structure. We develop a dynamic structure for the following 2-dimensional problem. We are given sets R_R and R_B of red and blue rectangles, respectively, and a parameter δ . The error of a point x in the plane is defined as

$$(\# \text{ of red rectangles containing } x) - \delta \cdot (\# \text{ of blue rectangles containing } x).$$

The approximation error of R_B with respect to R_R is the maximum approximation error over all points in the plane. Our goal is to maintain the approximation error of R_B with respect to R_R under insertions into and deletions from R_R and R_B . We use n to denote the total number of rectangles in the current sets.

Our structure uses a partitioning of the plane similar to the one used by Overmars and Yap [14]. More precisely, we partition the plane into vertical *slabs* by drawing $O(\sqrt{n})$ vertical lines such that in between any two consecutive lines there are at most \sqrt{n} vertices of rectangles (in the current set). A rectangle is said to *belong* to a slab if any vertex of the rectangle is contained within the slab. A rectangle is said to *cross* a slab if the intersection of the slab and the rectangle is not empty but the rectangle does not belong to the slab. Each slab σ is further subdivided by drawing horizontal segments connecting its two bounding lines through every vertex of a rectangle inside σ . This way we obtain a subdivision of the plane into cells with the following properties.

- There are $O(\sqrt{n})$ slabs and $O(n)$ cells.
- No cell contains a vertex of a rectangle from $R_R \cup R_B$ in its interior.
- A cell is crossed by at most $O(\sqrt{n})$ vertical edges.

Let \mathcal{A} represent the arrangement of $R_B \cup R_R$. Essentially, we need to maintain the face of this arrangement with maximum error under insertions and deletions of red or blue rectangles. The main idea is to maintain the maximum for each slab σ . In order to do this, we maintain for each cell in a given slab, the maximum in that cell with respect to the rectangles belonging to the slab. We first describe how to maintain the maximum in a given cell and then we show how to use this in order to maintain the maximum in a given slab.

As we are concentrating on a particular cell C in a slab σ , let $R(\sigma)$ represent the rectangles of $R_R \cup R_B$ that belong to σ clipped to within σ , and let $\mathcal{A}(\sigma)$ be the arrangement of rectangles $R(\sigma)$. Since C has no vertices in its interior and all rectangles in $R(\sigma)$ belong to σ , the part of $\mathcal{A}(\sigma)$ within C is formed by $O(\sqrt{n})$ vertical edges crossing C . Therefore, maintaining the maximum in C is a 1-dimensional problem and we can use the tree structure \mathcal{T}_C described in Lemma 2.1 to maintain the maximum. For each rectangle r of $R(\sigma)$ in C , insert an interval $[r_s, r_e]$ in \mathcal{T}_C where r_s and r_e are the x -coordinates of the left and right edges of r . The value associated with this interval is 1 if r is red and $-\delta$ otherwise. If r_s is to the left of the left side of the slab or r_e is to the right of the right side of the slab, we truncate r_s or r_e to the slab boundary since we only concentrate on what is within C .

We now turn our attention on how to maintain the maximum within a slab σ . There are two types of rectangles that contribute to the error of the faces of \mathcal{A} within σ : those that belong to σ and those that cross σ . The error of the rectangles that belong to σ are taken care of within each cell, so to maintain the maximum for the slab, we need to incorporate the information pertaining to the rectangles crossing the slab. Let $X(\sigma)$ represent all of the rectangles that cross σ . For a rectangle $r \in X(\sigma)$, consider the cells of σ that are contained in r . For each such cell, the maximum error in that cell only changes by a constant (depending on the color of r) because the whole cell is contained in r . So again, we are able to transform the problem to a 1-dimensional problem. Let Y_σ be the y -intervals of the cells of σ and let $Y_{X(\sigma)}$ be the y -intervals of the rectangles in $X(\sigma)$. To maintain the maximum in σ , we construct a tree \mathcal{T}_σ . For each interval y_i in Y_σ , we insert y_i in \mathcal{T}_σ and the associated value is the maximum error in the cell. For each y -interval y_j in $Y_{X(\sigma)}$, we insert y_j in \mathcal{T}_σ and the associated value is either 1 or $-\delta$ depending on the color of the rectangle.

Our structure consists of one tree for each slab and one tree for each cell within a slab. Therefore, the structure consists of $O(\sqrt{n})$ slab trees and $O(n)$ cell trees.

Updates. To insert a rectangle r into the structure, we proceed as follows. First, we find in $O(\sqrt{n})$ time all $O(\sqrt{n})$ slabs that are completely crossed by r . For each such slab σ , we insert the y -interval of r and the value of r into \mathcal{T}_σ . This takes at most $O(\sqrt{n} \log n)$ time.

Next we deal with the at most two slabs that contain a vertex of r . For each such slab σ , we find in $O(\sqrt{n})$ time all $O(\sqrt{n})$ cells intersected by r . There are two types of cells: the ones that contain a vertex of r and the ones that are crossed from top to bottom by r . For cell C of the latter type, we insert the error of r and the x -interval of r (clipped to within σ) into \mathcal{T}_C . Cells of the former type—there are at most two of them—have to be split using horizontal segments through the vertices of r . Since the number of intervals stored with a cell is $O(\sqrt{n})$, splitting a cell and rebuilding its structure can be done in $O(\sqrt{n})$ time. For each affected cell C in σ we now know its new error, so we update the slab tree \mathcal{T}_σ by deleting and re-inserting the affected cells. Overall, we spend $O(\sqrt{n} \log n)$ time to handle the at most two slabs containing an endpoint of r .

After this, we go over all slabs to recompute the new maximum error.

Deleting a rectangle is done using a similar procedure, so we omit the details. Finally, during the course of insertions and deletions, we may have to split a slab into two or merge two neighboring slabs in order to maintain the partition into $O(\sqrt{n})$ slabs each containing $O(\sqrt{n})$ cells. Merging or splitting can be done in $O(n \log n)$ time by simply reconstructing the trees for the slabs and cells. If we split a slab whenever its size is more than $2\sqrt{n}$ or merge two slabs when both their sizes are less than $\sqrt{n}/2$ then a standard amortization argument shows that given an update sequence containing n insertions and deletions that merges and splits can be achieved in amortized $O(\sqrt{n} \log n)$ time.

Lemma 2.5. *The maximum approximation error of a set of red and blue rectangles in the plane can be maintained in $O(\sqrt{n} \log n)$ amortized time per update, where n is the number of rectangles in the set.*

We now return to the original problem, of computing the approximation error of a set Q of m points in the plane with respect to a set P of n points, where the set of ranges is the family of all possible squares. Let $s_\rho(p)$ denote the square of size ρ centered at a point p . Furthermore, let $S_P(\rho) = \{s_\rho(p) : p \in P\}$ and let $S_Q(\rho) = \{s_\rho(q) : q \in Q\}$. Define the maximum error of $S_P(\rho)$ with respect to $S_Q(\rho)$ for point ranges as before. Then the maximum error we want to compute is given by $\max_{\rho>0} \{\text{error of } S_P(\rho) \text{ wrt } S_Q(\rho)\}$. When we let ρ increase, the error of $S_P(\rho)$ with respect to $S_Q(\rho)$ can only change when two edges of squares meet. Hence, we proceed as follows. We normalize the problem by replacing the x -coordinates of the vertical edges by their rank, and by replacing the y -coordinates of the horizontal edges by their rank. Now every square becomes a rectangle with coordinates from a universe of size $U = 2(n + m)$. We store these rectangles in a dynamic structure as described above. Whenever two vertical (or horizontal) edges swap ranks, we delete the rectangles from our structure, and re-insert the rectangles with their new normalized coordinates. This takes $O(\sqrt{n} \log n)$ time, and gives us the new error.

We start the process with a value of ρ that is small enough so that all rectangles are disjoint (and the error is δ or $\delta - 1$, depending on whether $Q \subset P$). The number of swaps we have to process is $O(n^2)$. We get the following result.

Theorem 2.6. *Let P be a set of n points in the plane, and let Q be a set of m points in the plane, with $m \leq n$. The approximation error of Q with respect to P for the family of all squares can be computed in $O(n^2 \sqrt{n} \log n)$ time.*

2.3. Rectangles as ranges

Let \mathcal{R} be the set of all possible rectangles in the plane. Dobkin et al. [9] present an algorithm that computes the approximation error of Q with respect to P in $O(n^2 \log n)$ time. Their algorithm is not sensitive to the size of the set Q . We present an algorithm that is sensitive to the relative sizes of the two point sets.

Like in the case of arbitrarily sized squares, the approximation error does not depend on whether we consider open or closed rectangles: for any open (closed) rectangle, there is a slightly smaller closed (larger open) rectangle that contains exactly the same points.

We start with a simple lemma limiting the number of rectangles to consider. Let B be the bounding box of $P \cup Q$.

Lemma 2.7. *There is an open rectangle with maximum positive error such that each side either contains a point from Q or is contained in the boundary of B . Similarly, there is a closed rectangle with maximum negative error such that each side contains a point from Q .*

Proof. Let r be an open rectangle of maximum positive error, that is, a rectangle that maximizes $|r \cap P| - \delta \cdot |r \cap Q|$. While there is a side of r that does not contain a point from Q on its boundary, move that side away from the center of r until it does contain a point of Q on its boundary or until the side is contained in the boundary of B . Since r is open, $|r \cap Q|$ remains the same as before the growing operation. Clearly, $|r \cap P|$ has not decreased, so the new rectangle still gives the maximum error.

Similarly, we can transform a closed rectangle of maximum negative error to one satisfying the conditions of the lemma by moving the sides towards the center (thus shrinking the rectangle) until each side contains a point of Q on its boundary. \square

Herein lies the main difference in approach between our algorithms and the algorithm of Dobkin et al. [9]: they verify all pairs of points rather than exploiting the above lemma. Lemma 2.7 immediately implies a fairly simple algorithm with $O(m^2 n + n \log n)$ running time, very similar to the first algorithm we presented for squares. However, we can do better, by using the following divide-and-conquer approach. As before, we show how to compute the positive approximation error; the negative error can be computed in a similar way.

Split the plane into two half-planes using a vertical line, and recursively compute the maximum error over all rectangles lying completely to the left of ℓ , and the maximum error over all rectangles lying completely to the right of ℓ . What remains is to compute the maximum error over all rectangles crossing ℓ , i.e. the merging step, which we describe below. The maximum of the three values is the global maximum.

In the merging step we have to find the rectangle r^* giving the largest error over all rectangles crossing a given line $\ell : x = \ell_x$. Our algorithm is based on the following observation. Let ℓ^- be the closed half-plane to the left of ℓ , and let ℓ^+ be the open half-plane to the right of ℓ . (We make one of the half-planes open to ensure that points on ℓ are counted only once.)

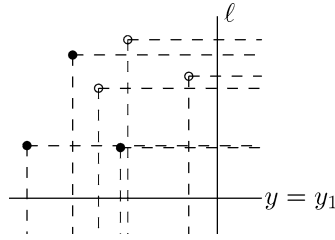


Fig. 2. Replacing points by quadrants in the merging step for rectangles.

Observation 2.8. The rectangle $r^* \cap \ell^-$ (respectively $r^* \cap \ell^+$) has the largest error of all rectangles whose right (respectively left) side lies on ℓ and whose top and bottom sides have the same y -coordinates as the top and bottom sides of r^* .

Let Y be the set of all y -coordinates of points in Q together with the y -coordinates of the top and bottom edge of the bounding box B . Next we show how to compute, for each y -interval $(y_1 : y_2)$ defined by y -coordinates in Y , the rectangle of maximum error over all rectangles with this y -interval whose right edge lies on ℓ .

Fix some y_1 . We can now restrict our attention to the quadrant to the left of ℓ and above the line $y = y_1$. Let $P(y_1)$ and $Q(y_1)$ be the subsets of P and Q , respectively, inside this quadrant. The rectangles we are interested in all have (ℓ_x, y_1) as bottom right corner, so if we restrict our attention to $P(y_1) \cup Q(y_1)$, we can regard the rectangles as being quadrants that are unbounded to the right and bottom. Hence, we can apply the same algorithm as we used for unit squares: First, we replace every point in $P(y_1) \cup Q(y_1)$ by a quadrant unbounded to the top and left. See Fig. 2. Next we sweep the arrangement of quadrants from bottom to top. Events are the y -coordinates of Y larger than y_1 —these include those of the points in $Q(y_1)$ —and the y -coordinates of the points in $P(y_1)$. We maintain a tree \mathcal{T} that maintains the maximum error of the intersections of the quadrants with the sweep line, as described in Lemma 2.1, and at each $y \in Y$ we report the maximum error. This gives us for a fixed y_1 in $O((m + n) \log(n + m))$ time the maximum error for each interval (y_1, y_2) .

We perform this procedure with each $y_1 \in Y$, taking $O(m(m + n) \log(n + m))$ in total. Hence, the merging step can be performed in this amount of time. To get a nicely balanced recursion tree, we choose ℓ at odd levels of the recursion tree such that at most half the points from P lie on either side of ℓ , and we choose it at even levels such that at most half the points from Q lie on either side of ℓ . This way we get the following recurrence for the running time, $T(n, m)$:

$$T(n, m) = O(m(m + n) \log(n + m)) + \sum_{i=1}^4 T(n_i, m_i),$$

$$\text{with } \sum_{i=1}^4 n_i = n \text{ and } n_i \leq n/2 \text{ for } i = 1, \dots, 4,$$

$$\text{and } \sum_{i=1}^4 m_i = m, \text{ and } m_i \leq m/2 \text{ for } i = 1, \dots, 4.$$

This gives $T(n, m) = O(m(n + m) \log(n + m))$.

Theorem 2.9. Let P be a set of n points in the plane, and let Q be a set of m points in the plane, with $m \leq n$. The approximation error of P with respect to Q for the family of all rectangles can be computed in $O(nm \log n)$ time.

Remark 2.10. If $m = o(n^{1/3})$, then the following simple approach is more efficient than the one above: preprocess the points in P for range counting, and query with each rectangle defined by four points from Q . Using $O(n \log n)$ preprocessing, range counting queries can be answered in $O(\log n)$ time [1], so this approach leads to a total time of $O((n + m^4) \log n)$.

3. Finding approximations with small error

We now turn our attention to finding good approximations of a specified size m for a given set P of n points. We will concentrate on square ranges, as this seems most natural in our application.

3.1. Data sets

Our input sets P consist of n points in the unit square, for various values of n . We use three types of distributions: uniform, clustered and real-world data. The clustered data sets were generated as follows. We randomly choose 20 cluster centers, draw a circle around each center, and generate points randomly within that circle according to a distribution that generates more points close to the center. Which fraction of the points goes to which cluster is also determined randomly. Fig. 3 shows an example of a clustered

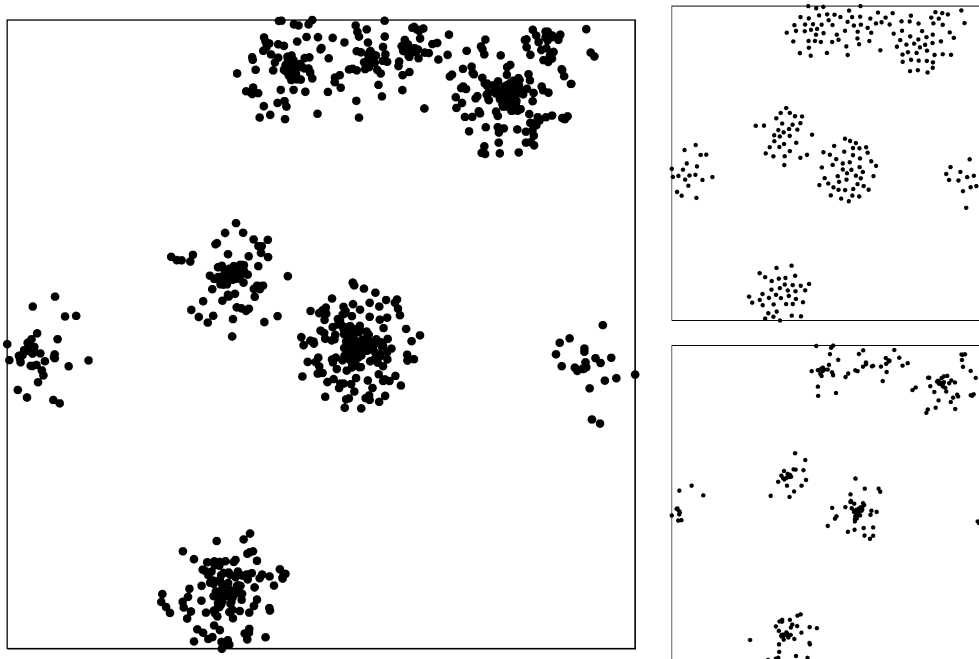


Fig. 3. Clustered 1000-point set P (on the left), and two 250-point approximations for P .

data set generated in this manner. The real-world data set represents the acres of harvested cropland in the USA in 1992 [15]—see Fig. 1.

3.2. Computing the error

Our heuristics call a subroutine to compute the error for given P and Q many times. We have implemented the $O(m^2n + n \log n)$ algorithm for computing the error for square ranges. For large n and m , this is rather slow. To speed up the heuristics we therefore want to replace the subroutine by a faster one. We do this by computing the error for squares of a fixed size, for several different sizes; for a fixed size we used the $O(n \log n)$ algorithm of Theorem 2.2. The hope is that if the number of sizes is large enough, the error we find is close enough to the real error, so that it will not harm our heuristics. Our first experiment is to test whether this hope is justified: we compare the real error, computed with the $O(m^2n + n \log n)$ algorithm, to the error computed by looking at squares of k different sizes only, for various values of k .

The results are summarized in Table 1.

For each distribution we have generated between 5 and 10 different sets P , and for each P between 8 and 20 different sets Q . Half of the choices for Q were taken as random samples from P , the other half was generated using another distribution. The table shows the average difference between the error for arbitrary squares and the error for k different fixed sizes, where the sizes were equally spaced. (We also tried sizes on a logarithmic scale, but obtained poorer results.) The numbers between brackets in the table give the maximum difference found in the experiments. If we take $k = 60$, then the average difference between the real error and the estimated error is always close to (and often smaller than) the dot value, and the maximum difference is close to twice the dot value. We conclude from this that it is safe to use the estimated error in our heuristics.

We use this estimate out of necessity. Computing the error exactly takes hours for the US data set, while computing the estimated error takes only a few seconds. Since our algorithms for finding good approximations have to repeatedly compute the error of an approximation, computing the exact error is not a reasonable option.

Table 1
Estimating the square error by k fixed sizes

n	Uniform						Clustered						Real	
	1000 (20 samples)			5000 (5 samples)			1000 (5 samples)			5000 (5 samples)			500	1000
m	50	100	250	100	250	500	50	100	250	100	250	500	500	1000
δ	20	10	4	50	20	10	20	10	4	50	20	10	22	11
$k = 5$	30	22	18	117	68	95	27	22	18	160	88	82	68	10
	(104)	(80)	(80)	(296)	(183)	(401)	(121)	(74)	(92)	(517)	(230)	(307)	(68)	(10)
$k = 10$	17	14	9	78	47	53	17	12	9	102	58	54	68	10
	(50)	(60)	(36)	(177)	(138)	(230)	(71)	(38)	(40)	(220)	(228)	(192)	(68)	(10)
$k = 30$	9	7	4	41	27	17	8	5	4	45	29	20	19	9
	(30)	(22)	(15)	(86)	(65)	(42)	(28)	(18)	(22)	(92)	(119)	(126)	(19)	(9)
$k = 60$	6	4	3	27	17	12	4	3	3	28	19	13	19	9
	(18)	(13)	(9)	(68)	(44)	(24)	(27)	(12)	(12)	(70)	(90)	(86)	(19)	(9)

3.3. The heuristics

Next, we experimented with several heuristics for generating an approximation Q of a desired size m for a given set P of n points. These heuristics fall into two classes:

- (1) Iterative algorithms that start with a random solution and then apply some iteration rule to try and improve upon it. These algorithms include traditional optimization algorithms such as simulated annealing and taking the best of k random samples.
- (2) Clustering algorithms that partition the set S into m groups and then choose one representative point for each group. These algorithms may partition the point set S directly (see Dobkin-Tal below) or may partition the plane thereby inducing a partition of S .

3.3.1. Iterative algorithms

The first class of heuristics that we consider are *iterative* algorithms. For this class, we consider any algorithm that works by testing many different solutions, i.e., subsets Q , and taking the best one. The differences between various iterative algorithms come from how the subsets Q are selected.

Heuristic 1. *Best of k random samples.* Here we take k random samples Q_1, \dots, Q_k of P , compute the approximation error for each of them, and return the best sample. Here k is a parameter. The larger the value of k , the better the approximation.

Heuristic 2. *Simulated annealing.* Simulated annealing (cf. [13]) is a general search technique that starts with an initial random solution (i.e. a random sample) and then tries to converge to an optimal solution by introducing random changes. A random change is kept if (1) it improves the current solution, or (2) some *annealing condition* is met.

In our implementation of simulated annealing, a random change involves choosing a random point of Q and replacing it with a random point from $P \setminus Q$. The annealing criterion is the following: During the i th round of annealing, we replace the solution Q by the solution Q' with probability $\exp((\Delta(Q, P) - \Delta(Q', P))/T_i)$. Here, T_i is a temperature parameter whose value is $T_i = (k - i)/k$, where k is the total number of rounds the algorithm runs for.

Heuristic 3. *Swapping.* We first obtain an approximation Q by taking a random sample of size m , and then try to improve it as follows. We compute a range r_{pos} with the largest positive error and a range r_{neg} with the largest negative error. We remove a random point in $Q \cap r_{\text{neg}}$ from Q , and add a random point in $(P \setminus Q) \cap r_{\text{pos}}$ to Q .

Initial experimental results with the swapping heuristic were encouraging. However, this heuristic is highly dependent on having a good starting configuration. When this doesn't happen, the algorithm can get stuck in a local minimum, after which no further improvement can be made. To overcome this problem, we implemented two variants of the swapping heuristic.

Heuristic 4. *Swapping with restart.* This is a version of the swapping heuristic that starts with a new random sample Q if 10 consecutive rounds of swapping fail to improve the solution.

Heuristic 5. *Swapping with 10% perturbation.* This is a version of the swapper that, after 10 consecutive rounds of swapping fail to improve the solution, removes $\lceil m/10 \rceil$ points of Q at random and replaces them with $\lceil m/10 \rceil$ points of $P \setminus Q$ selected at random.

3.4. Experimental results

Fig. 1 shows the progress of iterative algorithms for 1000 rounds on a set P of $n = 5000$ points chosen uniformly at random from the unit square. The algorithms are attempting to find a good approximation Q of size $m = 100$. The x -axis of the figure represents time (number of rounds) and the y -axis represents the approximation error. Note that, although the figure looks as if the various heuristics began with different starting configurations, this is caused by a lack of resolution, and is not actually the case. All experiments began with the same initial configuration.

The worst of the heuristics is clearly simulated annealing (Heuristic 2), which makes some quick improvements in the first few rounds and then gets trapped in a local minimum. Part of the problem with simulated annealing is that it completely ignores the problem and tries to make progress by introducing small random changes. It is very easy for the simulated annealing strategy to get stuck in a local minimum and never improve. Although it may be possible to improve the performance of the simulated annealing heuristic by tweaking the parameters, we were unable to do significantly better than the results presented here.

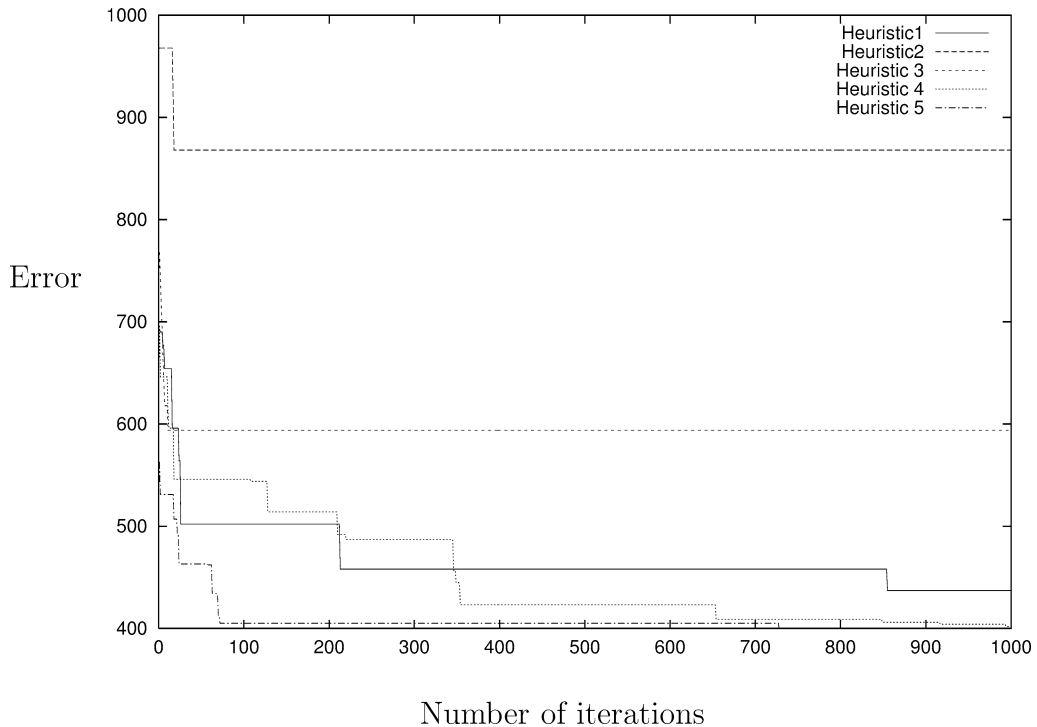


Fig. 4. The progress of Heuristics 1–5 over 1000 rounds.

The second-worst heuristic is the swapping heuristic (Heuristic 3). The swapper performs better than simulated annealing because it introduces a carefully-chosen change that is more likely to improve the current solution. However, it still only changes the current solution by one point and therefore quickly gets caught in a local minimum.

The two best heuristics are modifications of the swapping heuristic. Swapping with restart (Heuristic 4) and swapping with 10% perturbation (Heuristic 5) both achieve comparable results after 1000 rounds. However, swapping with 10% perturbation converges more quickly to a good solution. This seems to be due to the fact that, when it gets stuck in a local minimum, it restarts with a new solution that is still much better than a random sample.

Choosing the best of k random samples (Heuristic 1), a technique that is often mentioned in the literature, does not perform as well as the modified swapping heuristics. It reliably finds good approximations, but these are not quite as good as those found by the two modified swapping heuristics.

3.5. Clustering algorithms

We also considered algorithms that can be loosely termed “clustering” algorithms. These are algorithms that (implicitly or explicitly) partition the point set S into m groups and then select a representative point or points from each group.

Heuristic 6. Rows and columns. This heuristic produces a subset Q with $m = r \times s$ points by first sorting the points by x -coordinate and grouping the points into r columns, i.e., vertical strips, each containing n/r points. Next, the points within each column are sorted and grouped into $s = m/r$ rows, i.e. horizontal strips, of size n/m . Thus we obtain a partition of the plane into m rectangular cells each containing exactly n/m points.

For our set Q , we take a sample from each cell. Several strategies for choosing the best sample in each square were implemented. The one that worked best was to try $k = 50$ random samples and choose the sample with smallest error constrained to that cell.

Note that, in these experiments, the value of m is given, so we must factor m into r and s . We did this by taking $r = \lfloor \sqrt{m} \rfloor$ and then taking s to be the largest integer so that $r \times s \leq m$. This gives us an approximation that uses at most m points. When computing the quality of the resulting approximation we adjust the dot value δ accordingly.

Heuristic 7. Quadtrees. This heuristic is based on the well-known quadtree data structure. Let S be some axis-aligned square containing the point set P . We recursively partition S into squares as follows. If S contains fewer than 4δ points of P then we do nothing. Otherwise, we partition S into 4 equal squares and recursively partition each square.

Once this partition is computed, we choose a sample from each square of the partition. If a square contains k points of P then we choose a sample of size $\lfloor k/\delta + 1/2 \rfloor$ from that particular square. The sampling strategy is the same as used for Heuristic 6. As before, this does not always yield a solution with exactly n/δ points so, when computing the error we adjust the dot value δ accordingly.

Heuristic 8. Dobkin–Tal. The algorithm proposed by Dobkin and Tal [10] produces an approximation that is not a subset of P , by repeatedly finding closest pairs and replacing them by their midpoint.

Dobkin and Tal were originally interested in the dual setting of our problem: given a set of lines, find a smaller set of lines whose arrangement approximates the original arrangement. They solve the problem using dualization, so they arrive exactly at our problem.

Although their approach seems more suited to minimizing the Hausdorff distance between P and Q —indeed, they prove bounds on the minimal Hausdorff distance they achieve—they also use their algorithm in an application that is closely related to ours. Namely, they want to approximate the area half-plane discrepancy [5] of P with the discrepancy of Q . (The area half-plane discrepancy of a set of points in the unit square is defined as the maximum, over all half-planes, of the absolute difference between the fraction of points in the half-plane and the fraction of the unit square covered by the half-plane.) Now if we considered half-planes as regions, then the approximation error of Q with respect to P is an upper bound on the difference between the area discrepancies of P and Q . Dobkin and Tal claim that for some distributions of P the area discrepancy of P can be estimated better by a set Q computed with their algorithm than by a random sample. For this reason we also consider their algorithm in our experiments.

3.6. Experimental results

Table 2 shows the results for Heuristic 5, the best of the iterative heuristics, after 50 rounds and all the clustering algorithms. The tests were performed on 6 data sets of size $n = 5000$ and one real world data set. The data sets U5K{a,b,c} each consist of 5000 points uniformly distributed in the unit square. The data sets C5K{a,b,c} each consist of 5000 points drawn from the “city” distribution described earlier. The US1 data set is the data set shown in Fig. 1 and consists of 82516 points. For each data set, we used the algorithms to compute approximations with dot values $\delta = 100, 50, 20$ and 10.

These results suggest that the “rows and columns” heuristic seems to be the best choice of the clustering heuristics. For large values of δ , it is competitive with the quadtree heuristic and much better

Table 2
Experimental results for clustering algorithms

Heuristic	δ	US1	U5Ka	U5Kb	U5Kc	C5Ka	C5Kb	C5Kc
	100	3206	570	674	655	570	730	680
Heuristic 5	50	3066	556	507	427	515	407	614
(Swap w. 10%)	20	4396	359	268	310	364	368	343
	10	789	309	236	243	227	231	223
	100	1924	650	682	657	491	487	524
Heuristic 6	50	1036	705	359	748	405	410	367
(Rows & Cols.)	20	885	315	256	294	188	273	203
	10	780	153	195	156	119	128	132
	100	2289	582	604	739	520	400	524
Heuristic 7	50	1612	500	508	356	421	321	353
(Quadtree)	20	4285	261	230	276	225	217	207
	10	6728	439	467	398	398	396	343
	100	–	824	742	691	1445	1806	1335
Heuristic 8	50	–	575	519	704	1479	1752	1133
(Dobkin–Tal)	20	–	414	392	399	1378	1631	1028
	10	–	280	302	309	1302	1586	1011

than Dobkin–Tal. For small values of δ , the “rows and columns” heuristic is definitely the method of choice and outperforms the quadtree heuristic by a significant margin. This seems to be because the quadtree heuristic has trouble controlling the number of points in each cell, while the “rows and columns” heuristic has exactly δ points per cell.

Surprisingly, the simple “rows and columns” heuristic also seems to perform better than Heuristic 5, even though we allow Heuristic 5 to run for 50 rounds. This makes the “rows and columns” heuristic a very fast method of obtaining good quality solutions. In terms of computation time, the entire running time of the rows and columns heuristic is roughly the same as one or two rounds of an iterative heuristic.

Finally, the Dobkin–Tal heuristic does reasonably well for uniformly distributed points, but does very poorly with clustered point sets. This seems to be an artifact of the averaging effect obtained by repeatedly taking the midpoints of the pairs of points.

4. Concluding remarks

In some applications, it may be desirable to give outliers in P a bigger chance to be present in Q . This can be done by giving these points a higher weight. For instance, we can let the weight of each point be dependent on the number of points within a fixed distance from that point. By giving more and more weight to isolated points, the approximation is likely to become more and more uniform. The definition of approximation error and our algorithms can easily be extended to the weighted case, and it would be interesting to experiment with this.

In our application it seems most reasonable to look at the approximation error for families of squares or discs. We studied the case of squares, but it would be interesting to see if our algorithm to compute the approximation error in this case can be improved. We did not study discs at all in this paper. It is easy to compute the approximation error for discs in (close to) cubic time, but it remains open whether this can be done faster.

Finally, we suspect that computing the best approximation of a given size with respect to a given set P is NP-hard, but we have not been able to prove this.

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