# Algorithms for Stable Matching and Clustering in a Grid

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**Abstract.** We study a discrete version of a geometric stable marriage problem originally proposed in a continuous setting by Hoffman, Holroyd, and Peres, in which points in the plane are stably matched to cluster centers, as prioritized by their distances, so that each cluster center is apportioned a set of points of equal area. We show that, for a discretization of the problem to an  $n \times n$  grid of pixels with k centers, the problem can be solved in time  $O(n^2 \log^5 n)$ , and we experiment with two slower but more practical algorithms and a hybrid method that switches from one of these algorithms to the other to gain greater efficiency than either algorithm alone. We also show how to combine geometric stable matchings with a k-means clustering algorithm, so as to provide a geometric political-districting algorithm that views distance in economic terms, and we experiment with weighted versions of stable k-means in order to improve the connectivity of the resulting clusters.

#### 1 Introduction

A long line of research considers algorithms on objects embedded in  $n \times n$  grids, including problems in computational geometry (e.g., see [1,2,8,17,19,26,28,29]), graph drawing (e.g., see [5,10,14,30]), geographic information systems (e.g., see [13]), and geometric image processing (e.g., see [9,11,15,20]). Continuing this line, we consider in this paper the problem of matching grid points (which we view as pixels) to k center points in the grid. Pixels have a preference for centers closer to them, and centers prefer closer pixels as well. The goal is to match every center to an equal number of pixels and for the matching to be stable, meaning that no two elements prefer each other to their specified matches. For example, the centers could be facilities, such as polling places, fire stations, or post offices, that have assigned jurisdictions and equal operational capacities (in terms of how many pixels they can serve). Rather than optimizing some computationally challenging global quality criterion based on distance or area, we seek an assignment of pixels to centers that is locally stable. Figure 1 illustrates a solution to this stable grid matching problem for a  $900 \times 900$  grid and 100random centers. Note that some centers are matched to disconnected regions.

Stable grid matching is a special case of the classic *stable matching prob-*lem [18], which was originally described in terms of arranging marriages between N heterosexual men and women in a closed community. In this case, stability

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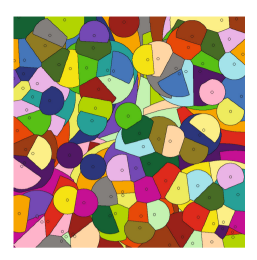


Fig. 1. An example solution to the *stable grid matching* problem for a  $900 \times 900$  grid and 100 centers distributed randomly. Pixels of the same color are assigned to the same center. (Color figure online)

means that no man-woman pair prefers each other to their assigned mates, which is necessary (and more important than, e.g., total utility) to prevent extramarital affairs. The Gale-Shapley algorithm [18] finds a stable matching for arbitrary preferences in  $O(N^2)$  time. For stable grid matching in an  $n \times n$  grid this would give a running time of  $O(n^4)$ , since each "man" would correspond to a pixel and each "woman" would correspond to one of  $\lceil n^2/k \rceil$  copies of a center. As we show, the geometric structure of the stable grid matching problem allows for significantly more efficient solutions.

We also study the effect of integrating a stable matching with a k-means clustering method, which alternates between assigning points to cluster centers and moving cluster centers to better represent their assigned points. Using stable matching for the assignment stage of this method allows us to fix the size of the clusters (for instance, to be all equally sized), which might be advantageous in some applications.

Prior Related Work. As mentioned above, there is considerable prior research on algorithms involving objects embedded in an  $n \times n$  grid. The stable grid matching problem that we study can be viewed as a grid-restricted version of the classic "post office" problem of Knuth [27], where one wishes to identify each point in the plane with its closest of k post offices, with the added restriction that the region assigned to each post office must have the same area. The continuous version of the stable grid matching problem, which deals with points in  $\mathbb{R}^2$  instead of discrete pixels, was studied by Hoffman et al. [21]. They showed that there is a unique solution, and there is a simple numerical method to find it: Start growing a circle from each center at the same time, all growing at the same speed. When a yet-unmatched point is reached by a circle, it is assigned to the corresponding

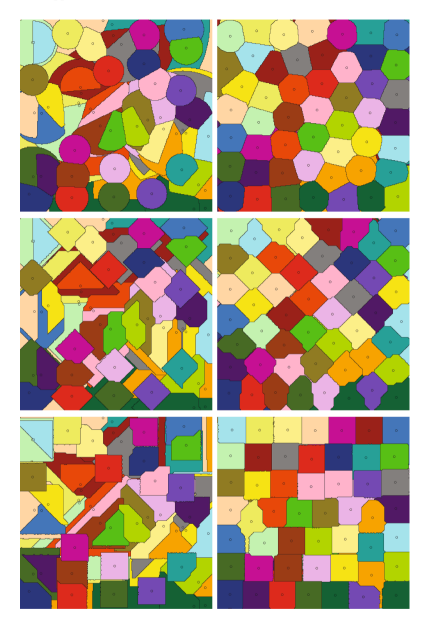
center. When a center reaches its quota (its region covers 1/k of the area of the square), its circle halts. (Note that if the halting condition is removed, we obtain the Voronoi diagram of the centers instead, as in the well-known solution to Knuth's post office problem, e.g., see [3].) Due to its continuous, numerical nature, Hoffman et al. did not analyze the running time of their method; hence, there is motivation to study the grid-based version of this problem.

With respect to the related problem of k-means clustering, we are interested in a grid-based version of this problem as well, which has been studied extensively in non-grid discrete contexts (e.g., see [22,24]). In the continuous version of this problem, one is interested in partitioning a geometric region into subregions that all have the same area (e.g., see [6]). One of the motivations for such partitions is in *political districting*, for which there is additional related prior work (e.g., see [32]). The goal of political districting is to partition a territory into regions (districts) which all have roughly the same population size and are "compact", which informally means that their shape should be connected and resemble a circle rather than an octopus [32]. Ricca et al. [31] adapted the concept of Voronoi regions to the discrete setting in order to use them for political districting. Voronoi regions ensured good compactness but poor population balance, however. Thus, there is motivation for a clustering algorithm based on the use of stable matchings, since such partitions enforce the property that all regions have the same size (at the possible cost of connectivity). Finding a scheme that guarantees both size equality and compactness is an open problem of interest.

Problem Definition. In the stable grid matching problem, we are given a square  $n \times n$  grid and k points called centers within the grid. The lattice points are called pixels or sites. Sites implicitly rank the centers in increasing order of distance, and centers similarly implicitly rank pixels in increasing order by distance. A matching is a mapping from sites to centers. The goal is to find a matching with the following two properties (see Fig. 2, left column):

- 1. The region of each center (the set of sites assigned to it) must have the same size up to roundoff errors. The quota of a center is the number of sites that must be in its region. If  $n^2$  is a multiple of k, then all the quotas are  $n^2/k$ . Otherwise, some centers are allowed one extra site.
- 2. The matching must be *stable*. A matching is not stable when a pair of sites  $(p_1, p_2)$  is assigned to centers  $c_1$  and  $c_2$  such that  $p_1$  prefers (i.e., according to some metric is closer to)  $c_2$  over  $c_1$  and  $c_2$  prefers  $p_1$  over  $p_2$ . This is unstable because  $p_1$  and  $c_2$  prefer each other to their current matches.

Combining k-means with Stable Assignment. The k-means clustering method is to partition a data set (which, in our case, is an  $n \times n$  grid) into k regions, based on a simple iterative refinement algorithm (which is called the k-means algorithm or Lloyd's algorithm, e.g., see [24]): We begin by choosing k points, called cluster centers, randomly in the space. Then, we iteratively repeat the following two phases: (1) assignment step: each object is assigned to its closest center, and (2) update step: each center is moved to the centroid of the objects assigned to it.



**Fig. 2.** Left: stable matching in a  $300 \times 300$  grid with the same 50 random centers for the Euclidean (top), Manhattan (center), and Chebyshev (bottom) metrics. Right: result of the stable k-means algorithm with unweighted centroids for each metric.

Lloyd's algorithm converges to a (locally optimal) partition that minimizes the sum of the squared distances from each object to its assigned center [24]. In this paper, we propose a variation, which we call  $stable\ k$ -means, where the

assignment step is replaced by a stable matching between objects and centers, so as to achieve the additional property that the regions all have equal area (to within roundoff errors). Intuitively, the goal is to implement Lloyd's algorithm with stable grid matching so as to improve the compactness of the regions while preserving equal-sized clusters.

We have found through experimentation that, although the stable k-means method succeeds in improving compactness, centers can sometimes stop moving while we are executing Lloyd's algorithm before their regions became completely connected (e.g., see Fig. 2). Thus, we introduce in this paper an additional heuristic, where we use weighted centroids, which are more sensitive to the outlying parts of their region. The usual centroid of a set of points S is defined as  $(\sum_{q \in S} q)/|S|$ , where the points are regarded as two-dimensional vectors so that the sum makes sense. Instead, we can compute a weighted centroid as  $(\sum_{q \in S} w_q q)/(\sum_{q \in S} w_q)$ . A natural choice to use for the weight  $w_q$  of a point q assigned to the region of the center c is the distance from q to c raised to some exponent p that we can choose,  $d(q,c)^p$ . The larger p is, the more sensitive the weighted centroids are to outliers. When p = 0, we get the usual centroid. When  $p \to +\infty$ , we get the circumcenter of the region, and when  $p \to -\infty$  we get the current center.

#### Contributions. In this paper, we provide the following results:

- The stable grid matching problem, for a grid of  $n \times n$  pixels with k centers, can be solved by a randomized algorithm with expected running time  $O(n^2 \log^5 n)$ . Since an  $n \times n$  grid has  $\Theta(n^2)$  pixels, this quasilinear bound improves the  $O(n^4)$  time of the Gale-Shapley algorithm. However, this algorithm uses intricate data structures that make it challenging to implement in practice.
- Given the pragmatic challenges of the above-mentioned quasilinear-time algorithm, we provide two alternative algorithms, a "circle-growing algorithm" and a "distance-sorting" method, both of which are simple to implement and have running times of  $O(n^2k)$ .
- We provide an experimental analysis of these two practical algorithms, where we observe that the circle-growing algorithm is more efficient at finding low-distance matched pairs, while the distance-sorting based method is more efficient when pairs are farther apart. Therefore, we show that it is advantageous to switch from one algorithm to the other partway through the matching process, potentially achieving running times with a sublinear dependence on k. We experiment with the optimal cutoff for switching between these two algorithms.
- We also provide the results of experiments to test the connectivity of the clusters obtained by our stable k-means algorithm, with weighted variants for finding centroids. Our experiments support the conclusion that no choice of a weight exponent p will always result in total connectivity. Nevertheless, our experiments provide evidence that the best results come from the range  $-0.8 \le p \le 0.4$ . Empirically, more highly negative values of p tend to make

the algorithm converge slowly or fail to converge, while more highly positive values of p lead to oscillations in the center placement. See the full version of the paper for additional figures of these cases.

## 2 Algorithms

Our stable grid matching algorithms start with an empty matching and add center—site pairs to it. Given a partial matching, we say a site is *available* if it has not been matched yet, and a center is *available* if the size of its region is smaller than its quota. A center—site pair is available if both the center and site are available, and it is a *closest available pair* if it is available and the distance from the center to the site is minimum among all available pairs. It is simple to prove that if an algorithm starts with an empty matching and only adds closest available pairs to it until it is complete, the resulting matching is stable.

#### 2.1 Circle-Growing Algorithm

In this section we describe our main practical algorithm, the circle-growing algorithm, which mimics the continuous construction from [21]. First, we obtain the list of all the lattice points with coordinates ranging from -n to n sorted by distance to the origin. The resulting list P emulates a circle growing from the origin. When initializing P, we can gain a factor of eight savings in space by sorting and storing only the points in the triangle  $\triangle(0,0)(0,n)(n,n)$ . The remaining points can be obtained by symmetry: if p=(x,y) is a point in the triangle, the eight points with coordinates of the form  $(\pm x, \pm y)$  and  $(\pm y, \pm x)$  are at the same distance from the origin as p. Moreover, in applications where we find multiple stable grid matchings, such as in the stable k-means method, we need only initialize P once. The way we use P depends on the type of centers we consider.

Integer Centers. In this case we can use the fact that if we relocate the points in P relative to a center, then they are in the order in which a circle growing from that center would reach them. To respect that all the circles grow at the same rate, we iterate through the points in P in order. For each point p, we relocate it relative to each center c to form the site p+c (the order of the centers does not matter). We add to the matching any available center—site pair (c, p+c). We iterate through P until the matching is complete.

We require  $O(n^2)$  space and  $O(n^2 \log n)$  time to sort the points in P. For the Euclidean metric instead of using distances to sort P we can use squared distances, which take integer values between 0 and  $2n^2$ . Then, we can use an integer sorting algorithm such as counting sort to sort in  $O(n^2)$  time [12, Chap. 8.2]. Since each point in P results in up to O(k) center—site pairs, we need  $O(n^2k)$  time to iterate through P.

Real Centers (Algorithm 1). If centers have real coordinates, we cannot translate the points in P relative to the centers, because p+c is not necessarily a lattice point. The workaround is to associate each center c to its closest lattice point  $p_c$ . Let  $\delta$  be the maximum distance  $d(c, p_c)$  among all centers. Then, the centersite pairs "generated" by each point p in P have the form  $(c, p+p_c)$  and their distances can vary between  $d(p, O) - \delta$  and  $d(p, O) + \delta$  (where O denotes the origin, (0,0)). Consequently, the distances of pairs generated by points  $p_i, p_j$  in P with i < j may intertwine, but only if  $d(p_j, O) - \delta \le d(p_i, O) + \delta$ . The points in P after  $p_i$  whose pairs might intertwine with those of  $p_i$  form an annulus centered at O with small radius  $d(p_i, O)$  and big radius  $d(p_i, O) + 2\delta$ .

Since  $\delta$  is a constant (for the Euclidean metric,  $\delta \leq \sqrt{2}/4$ ), it can be derived from the Gauss circle problem that such an annulus contains  $O(d(p_i, O)) = O(n)$  points.

#### **Algorithm 1.** Circle growing algorithm for k real centers on an $n \times n$ grid.

```
Set all sites as unmatched.
Set the quota of the first n^2 \mod k centers to \lceil n^2/k \rceil.
Set the quota of the remaining centers to \lfloor n^2/k \rfloor.
Let P = \text{list of points } (x, y) \text{ such that } -n < x, y < n.
Sort P by nondecreasing distance to (0,0).
For each center c, let p_c = (\text{round}(c_x), \text{round}(c_y)).
Let \delta = \max\{\operatorname{dist}(c, p_c)\} among all centers.
j \leftarrow 1
while the matching is not complete do
     L \leftarrow \text{ empty list}
     i \leftarrow \min(j + n, |P|)
     for all p \in P_j, \ldots, P_i do
                                           \triangleright Add to L pairs generated by points in the next chunk
          for all centers c with quota > 0 do
               s \leftarrow p + p_c
               if 0 \le s_x, s_y < n and s is still available then
                    Add (c, s) to L.
     Let d = \max\{\operatorname{dist}(c, s)\} among all pairs (c, s) \in L.
     for all p \in P_{i+1}, \ldots, P_{|P|} do
                                                     \triangleright Add to L pairs closer than pairs already in L
          if dist(p, O) > dist(P_i, O) + 2\delta then
               break
          for all centers c with quota > 0 do
               s \leftarrow p + p_c
               if 0 \le s_x, s_y < n and s is still available and dist(c, s) \le d then
                    Add (c, s) to L.
     Sort L by nondecreasing center–site distance.
     for all (c,s) \in L do
          if c and s are available then
               Match s and c.
               Reduce the quota of c by 1.
     j \leftarrow i + 1
```

The algorithm processes the points in P in chunks of n at a time, adding available center—site pairs generated by points in the chunk (or points after it, as we will see) to the matching in order by distance. The invariant is that after a chunk is processed, its points do not generate any more available pairs, and we can move on to the next one until the matching is complete. To do this, for each chunk we construct the list L of all the pairs generated by its points. Let d be the maximum distance among these pairs. If  $p_i$  is the last point in the chunk, the points in P from  $p_{i+1}$  up to the last point at distance to the origin at most  $d(p_i, O) + 2\delta$  can generate pairs with distance less than d. We add any such pair to L. We have to check O(n) additional points, so L still has size O(kn). We sort all these pairs and consider them in order, adding any available pair to the matching. Since each chunk has size n, there will be O(n) chunks. Each one requires sorting a list of O(kn) pairs, which requires  $O(kn \log n)$  time (since  $k \leq n^2$ ) and O(kn) space. In total, we need  $O(n^2k \log n)$  time and  $O(n^2 + nk)$  space.

#### 2.2 Distance-Sorting Methods

Unless the centers are clustered together, the circle-growing algorithm finds many available pairs in the early iterations. However, it reaches a point in which most circles overlap. Even if the centers are randomly distributed, in the typical case a large fraction of centers have "far outliers", sites which belong to their region but are arbitrarily far because all the area in between is claimed by other centers. Consequently, many centers have to scan a large fraction of the square. At some point, thus, it is convenient to switch to a different algorithm that can find the closest available pairs quickly. In this section, let m and  $k \leq m$  denote, respectively, the number of available sites and centers after a matching has been partially completed.

Pair Sort Algorithm. This algorithm simply sorts all the center–site pairs by distance and considers them in order, adding any available pair to the matching until it is complete. This algorithm is convenient when we can use integer sorting techniques, as in the case of the Euclidean metric and integer centers. Then, it requires O(mk) time and space.

While the pair sort algorithm has a big memory requirement to be used starting with an empty matching, used after the circle-growing algorithm has matched a large fraction of sites results in improved performance.

Pair Heap Algorithm. When centers have real coordinates, sorting all the pairs takes  $O(mk \log m)$  time, but we can do better. We find for each site s its closest center  $c_s$ , and build a min-heap with all the center—site pairs of the form  $(c_s, s)$  using  $d(c_s, s)$  as key. Clearly, the top of the heap is a closest available pair. We can iteratively extract and match the top of the heap until one of the centers becomes unavailable. When a center c becomes unavailable, all the pairs in the heap containing c become unavailable. At this point, there are two possibilities:

Eager update. We find the new closest available center of all the sites that had c as closest center and rebuild the heap from scratch so that it again contains one pair for each available site and its closest available center.

Lazy update. We proceed as usual until we actually extract a pair  $(c_s, s)$  with an unavailable center. Then, we find the new closest available center only for s, and reinsert the new pair in the heap.

In both cases, we repeat the process until the matching is complete.

We have not addressed yet how to find the closest center to a site. For this, we can use a nearest neighbor (NN) data structure that supports deletions. Such a data structure maintains a set of points and is able to answer *nearest neighbor queries*, which provide a query point q and ask for the point in the set closest to q. For the pair heap algorithm, we initialize the NN data structure with the set of centers and delete them as they become unavailable.

Since we need deletions we can use a *dynamic* NN data structure, i.e., with support for insertions as well as deletions. The simplest NN algorithm is a linear search, and a dynamic data structure based on it has O(k) time per query and O(1) time per update. The best known complexity of a dynamic NN data structure is  $O(\log^5 k)$  amortized time per operation [7,25].

Given that we know all the query points for our NN data structure ahead of time (the sites), we can build for each site s an array  $A_s$  with all the centers sorted by distance to s. Then, the closest center to a site s is  $A_s[i_s]$ , where  $i_s$  is the index of the first available center in  $A_s$ . When a center is deleted we simply mark it. When we get a query for the closest center to a site s, we search  $A_s$  until we find an unmarked center. We can start the search from the index of the center returned in the last query for s. This data structure requires O(mk) space and has a  $O(mk \log k)$  initialization cost to sort all the arrays. The interesting property is that if we do O(k) queries for a given site s, we require O(k) time for all of them, as in total we traverse  $A_s$  only once. We call this data structure presort, although it is not strictly a NN data structure because it knows the query points ahead of time.

In the pair heap algorithm, we can combine eager and lazy updates with any NN data structure. In any case, the running time is influenced by  $\alpha$ , the sum among all centers c of the number of sites that had c as closest center when c became unavailable. In the worst case  $\alpha = O(km)$ , but assuming that each center is equally likely to be the closest center to each site, the expected value of  $\alpha$  is O(m). In the full version of the paper we test the value of  $\alpha$  empirically in several different settings, and in every case we find  $\alpha < 10 \, m$ .

With eager updates in total we have to initialize the NN data structure, perform m extract-min operations,  $O(m+\alpha)$  NN queries, k NN deletions, and rebuild the heap k times. Thus, the running time is  $O(P(k,m) + m \log m + (m+\alpha)Q(k) + kD(k) + km)$ , where P(k,m) is the cost of initializing the NN data structure of choice with k points (and m query points, in the case of the presort data structure), and Q(k) and D(k) are the costs of queries and deletions, respectively. With lazy updates, instead of rebuilding the heap we have  $O(\alpha)$  extra insert and extract-min heap operations, which requires  $O(\alpha \log m)$  time.

For real centers, the best worst-case bound is with eager deletions and the presort NN data structure. In that case, we have that the NN queries take O(km) for any  $\alpha$ , so the total running time is  $O(mk \log k + m \log m)$ . If we assume that  $\alpha = O(m)$ , then the best time is with lazy deletions and the NN data structure from [7,25]. The running time with this heuristic assumption is  $O(m \log^5 k + m \log m)$ .

#### 2.3 Bichromatic Closest Pairs and Nearest Neighbor Chains

We now describe a less-practical solution based on bichromatic closest pairs which achieves the best theoretical running time that we have been able to prove. A bichromatic closest pair (BCP) data structure maintains a set of points, each colored red or blue, and is able to answer queries asking for the closest pair of different color.

The stable grid matching problem can be solved with a BCP data structure that supports deletions, either on its own or after the circle-growing algorithm. We first initialize the data structure with the available sites and centers as blue and red points, respectively. Then, we repeatedly find and match the closest pair, remove the site, and remove the center if it becomes unavailable. The running time is O(P(m) + mQ(m) + mD(m)), where P(m), Q(m), and D(m) are the initialization, query, and deletion costs, respectively, for the BCP data structure of choice containing m blue points and  $k \leq m$  red points.

Eppstein [16] proposed a fully dynamic BCP data structure that uses an auxiliary dynamic NN data structure. Using it, the sequence of operations required to solve the stable grid matching problem takes  $O(mT(m)\log^2 m)$  time, where T(m) is the cost per operation of the NN data structure. In particular, combining this with the dynamic nearest neighbor data structure of Chan [7] and Kaplan et al. [25] gives a total time bound of  $O(n^2 \log^7 n)$  for this problem.

To improve this, we observe that (with a suitable tie-breaking rule to ensure that no two distances are equal) it is not necessary to find the bichromatic closest pair in each step: it suffices, instead, to find a mutual nearest neighbor pair: a pixel and a center that are closer to each other than to any other pixel or center. The reason is twofold. First, in the algorithm that repeatedly finds and removes closest pairs, every pair (c,p) of mutual nearest neighbors eventually becomes a closest pair, because until they do, nothing else that the algorithm does can change the fact that they are mutual nearest neighbors. So (c,p) will eventually become matched by the algorithm. Second, if we find a pair (c,p) that will eventually become matched (such as a mutual nearest neighbor pair), it is safe to match them early; doing so cannot affect the correctness of the rest of the algorithm.

To find these, we may adapt the nearest-neighbor chain algorithm from the theory of hierarchical clustering [4,23] which uses a stack to repeatedly find pairs of mutual nearest neighbors at a cost of O(1) nearest neighbor queries per pair. In more detail, the algorithm is as follows.

1. Initialize two dynamic nearest neighbor structures for the pixels and centers, and an empty stack S.

- 2. Repeat the following steps until all pixels have been matched:
  - (a) If S is empty, push an arbitrary point (either a pixel or a center) onto S.
  - (b) Let p be the point at the top of S, and use the nearest neighbor data structure to find the nearest point q of the opposite color to p.
  - (c) If q is not already on S, push it onto S. Otherwise, q must be the second-from-top point on S, and is a mutual nearest neighbor with p. Pop p and q, match them to each other, and remove one or both of p or q from the nearest neighbor data structure (always remove the pixel, and remove the center if it becomes unavailable).

Note that in step 2. (c) q must be second-from-top because we have a cycle of (non-mutual) nearest neighbors starting with  $p \to q$  and then up the stack back to p. At each step along this cycle, the distance decreases or stays equal. But it cannot decrease, because there would be no way to increase back again, and nothing but  $q \to p$  can be equal to  $p \to q$ , because we are using a tie-breaking rule. So the cycle has length two and q is second-from-top.

Each step that pushes a new point onto S can be charged against a later pop operation and its associated matched pixel, so the number of repetitions is  $O(n^2)$ . This algorithm gives us the following theorem.

**Theorem 1.** The stable grid matching problem can be solved in  $O(n^2)$  operations of a dynamic nearest neighbor data structure. In particular, with the structure of Chan [7] and Kaplan et al. [25], the time is  $O(n^2 \log^5 n)$ .

## 3 Experiments

Datasets. Table 1 summarizes the parameters used in the different experiments. We use the following labels for the algorithms: CG the circle-growing algorithm alone, and PS and PH for the combination of CG and the pair sort and pair heap algorithms, respectively. Moreover, for the pair heap algorithm we consider the following variations: eager/presort  $(PH_{E,P})$ , eager/linear search  $(PH_{E,L})$ , lazy/presort  $(PH_{L,P})$ , and lazy/linear search  $(PH_{L,L})$ .

We focus on the Euclidean metric, but in the full version of the paper we also consider the Manhattan and Chebyshev metrics. The parameter n is the length of the side of the square grid, and k is the number of centers. In all the experiments, the centers are chosen uniformly and independently at random. Moreover, every data point is the average of 10 runs, each starting with different centers.

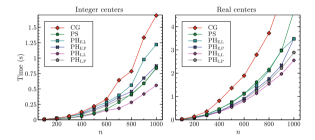
The cutoff is the parameter used to determine when to switch from the circle-growing algorithm to a different one. We define it as a ratio between the number of available pairs and the number of pairs already considered by the circle-growing algorithm.

The algorithms were implemented in C++ (gcc version 4.8.2) and the interface in Qt. The experiments were executed by a Intel(R) Core(TM) CPU i7-3537U 2.00 GHz with 4 GB of RAM, on Windows 10.

Experiment	Algorithms	Metric	n	k	Cutoff
Exec. time (Fig. 3)	All	$L_2$	varies	10n	0.15
Cutoff (Fig. 4)	$CG, PH_{L,L}$	$L_2$	1000	varies	varies

Table 1. Summary of parameters used in the experiments section.

Algorithm Comparison. Figure 3 contains a comparison of all the algorithms. Pair heap is generally better than pair sort, even for integer distances where it has a higher theoretical complexity. Among pair heap variations, lazy/linear is the best for both types of centers. In general lazy updates perform better, but eager/presort is also a strong combination because they synergize: eager updates require more NN queries in exchange for less extract-min heap operations, and the presort data structure has fast NN queries.



**Fig. 3.** Execution time of the various algorithms for integer (left) and real (right) centers. For all the methods but CG, the cutoff is 0.15. Each data point is the average of 10 runs with 10n randomly distributed centers and the  $L_2$  metric.

Optimal Cutoff. When combining the circle-growing algorithm with another algorithm, the efficiency of the combination depends on the cutoff used to switch between both. If we switch too soon, we don't exploit the good behavior of the circle-growing algorithm when circles are still mostly disjoint. If we switch too late, the circle-growing algorithm slows down as it grows the circles in every direction just to reach some outlying region.

Figure 4 illustrates the role of the cutoff. It shows that most of the execution time of the circle-growing algorithm is spent with the very few last available pairs, so even a really small cutoff prompts a substantial improvement. After that, the additional time spent in the pair heap algorithm slightly beats the savings in the circle-growing algorithm, resulting in a steady increase of the total running time.

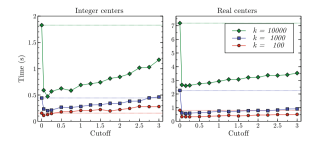


Fig. 4. Execution time of the circle-growing algorithm for integer (left) and real (right) centers, combined with the pair heap algorithm with lazy updates and a linear search NN data structure. The dotted lines denote the running time of the circle-growing algorithm alone, i.e., with cutoff 0. Each data point is the average of 10 runs with randomly distributed centers, n = 1000, and the  $L_2$  metric.

#### 4 Discussion

We have defined the stable grid matching problem, developed efficient theoretical algorithms and practical implementations of slower but simpler algorithms for this problem, and used our implementation to test different strategies for center placement in k-means like stable clustering algorithms. However, this work leaves several open questions:

- For which n and k does the stable grid matching problem have a placement of centers for which all clusters are connected, and how can such centers be found?
- Can the worst-case running time of our theoretical  $O(n^2 \log^5 n)$ -time algorithm be improved? Is it possible to achieve similar runtimes without going through fully-dynamic bichromatic closest pair data structures?
- Can we obtain practical algorithms whose runtime has lower worst-case dependence on k than our  $O(n^2k)$ -time circle-growing and distance-sorting methods?
- Our bichromatic closest pair and distance-sorting algorithms can be made to work for arbitrary point sets (not just pixels) but the circle-growing method assumes that the points form a grid, and its time analysis depends on the fact that the grid is a fat polygon (so that the area of each circle is proportional to the number of grid points that it covers) and that testing whether a point belongs to the grid is trivial. Can this method be extended to pixelated versions of more complicated polygons?
- How efficiently can we perform similar distance-based stable matching problems for graph shortest path distances instead of geometric distances? Can additional structure (such as the structures found in real-world road networks) help speed up this computation?

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