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Efficient trajectory queries under the Fréchet distance (GIS Cup)

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ABSTRACT
Consider a set \( \mathcal{P} \) of trajectories (polygonal lines in \( \mathbb{R}^2 \)), and a query given by a trajectory \( Q \) and a threshold \( \epsilon > 0 \). To answer the query, we wish to find all trajectories \( P \in \mathcal{P} \) such that \( \delta_\mathcal{P}(P, Q) \leq \epsilon \), where \( \delta_\mathcal{P} \) denotes the Fréchet distance. We present an approach to efficiently answer a large number of queries for the same set \( \mathcal{P} \). Key ingredients are (a) precomputing a spatial hash that allows us to quickly find trajectories that have endpoints near \( Q \); (b) precomputing simplifications on all trajectories in \( \mathcal{P} \); (c) using the simplifications and optimizations of the decision algorithm to efficiently decide \( \delta_\mathcal{P}(P, Q) \leq \epsilon \) for most \( P \in \mathcal{P} \).

CSC CONCEPTS
- Information systems → Geographic information systems;
- Theory of computation → Computational geometry;

KEYWORDS
Fréchet distance, trajectories, range searching

ACM Reference Format:

1 INTRODUCTION
With GPS sensors becoming low-cost and near-ubiquitous, trajectory data has seen a rapid increase in volume. A central problem, underlying many trajectory analyses in one way or another, is the need to quantify trajectory similarity and to find trajectories similar to a given one. A common way to quantify trajectory similarity is the Fréchet distance [3], which has been used to, e.g., find similar subtrajectories [4] or detect interaction between moving objects [5].

To define this metric, consider two trajectories (polygonal lines), \( P = \langle p_0, \ldots, p_m \rangle \) and \( Q = \langle q_0, \ldots, q_n \rangle \), defined by their vertices. We interpret these as functions \( P: [0, m] \rightarrow \mathbb{R}^2 \) using \( P(i + \lambda) = (1 - \lambda)p_i + \lambda p_{i+1} \), and likewise for \( Q \). Now, consider the set \( \Psi \) of all orientation-preserving homeomorphisms between the two parameter spaces, that is, of continuous functions \( \psi: [0, m] \rightarrow [0, n] \) with a continuous inverse such that \( \psi(0) = 0 \) and \( \psi(m) = n \). The Fréchet distance is now defined as follows:

\[ \delta_\Psi(P, Q) = \inf_{\psi \in \Psi} \max_{\epsilon \in [0, m]} \| P(\psi) - Q(\psi(\epsilon)) \| \]

The challenge. The GIS Cup 2017 sets out the following challenge. Consider a database \( \mathcal{P} \) of trajectories, together with a set \( \mathcal{Q} \) of range searching queries [1], where each query is a pair \( (Q, \epsilon) \) of a trajectory \( Q \) and a threshold \( \epsilon > 0 \). For each query \( (Q, \epsilon) \), we are tasked to find all trajectories \( P \in \mathcal{P} \) with \( \delta_\mathcal{P}(P, Q) \leq \epsilon \). In particular, \( \mathcal{P} \) and \( \mathcal{Q} \) are sufficiently large for preprocessing to pay off.

Contributions. We provide an efficient approach to solve the given challenge, by combining and optimizing a number of techniques. The key ingredients are spatial hashing, simplification, and optimized exact decision procedures. We first provide a high-level overview of our algorithm, followed by a detailed explanation of each component. The implementation is available online\(^1\).

2 ALGORITHM OVERVIEW
The algorithm solves the GIS Cup problem in two stages: a preprocessing stage, followed by a query stage. The preprocessing stage consists of building two main data structures: a spatial hash and four levels of simplifications for each trajectory in \( \mathcal{P} \). The query stage uses these two data structures determine for each \( P \in \mathcal{P} \) whether \( \delta_\mathcal{P}(P, Q) \leq \epsilon \), for each query \( (Q, \epsilon) \). Each query is processed in four steps, as summarized below.

1. We first use the spatial hash to obtain a candidate set \( C \subseteq \mathcal{P} \), which contains all trajectories whose start and endpoints are within distance \( \epsilon \) of the start and endpoints of \( Q \).
2. For each trajectory \( P \in C \), we now do the following steps, stopping as soon as we know how \( \delta_\mathcal{P}(P, Q) \) compares to \( \epsilon \).
   a. Starting with the most coarse level, we compare the simplifications of \( P \) and \( Q \) to try to decide whether \( \delta_\mathcal{P}(P, Q) \leq \epsilon \).
   b. An imprecise but fast greedy algorithm is run to check if \( \delta_\mathcal{P}(P, Q) \leq \epsilon \) for cases in which a simple linear homeomorphism suffices (equal-time distance).
   c. We run our exact continuous decision Fréchet algorithm to establish whether \( \delta_\mathcal{P}(P, Q) \leq \epsilon \). This is always conclusive.

The key idea of the above is to spend as little time as possible to establish whether \( P \in \mathcal{P} \) is within Fréchet distance \( \epsilon \) of a trajectory \( Q \). Each consecutive step takes more time, but is more precise. The

\(^{1}\)https://github.com/TWTDIG/GISCUP17TUE
earlier steps are powerful enough to filter out a majority of the trajectories. In particular, only very few trajectories require Step 2c, which could in theory take quadratic time, whereas the other steps are linear or use significantly simplified trajectories. We use parallelism to speed up computation where possible, using a number of parallel tasks up to what the system can handle.

Figure 1 shows the pruning percentages on our test data set\(^2\). We constructed this data set, using a database \(\mathcal{P}\) consists 3200 trajectories randomly sampled from the challenge sample trajectory set. The corresponding query set \(\mathcal{Q}\) consists of 6400 trajectories randomly selected from the challenge sample trajectory set, with ranges \(\epsilon\) generated such that the average query returns 2\% of \(\mathcal{P}\), while rare queries return no trajectories or up to 30\% of \(\mathcal{P}\). This makes the queries fairly realistic while also containing exceptional cases. In our tests, the algorithm solves the problem in 7 seconds, which is 30 times faster compared to using no pruning steps (i.e. only using Step 2c). Total processing time increases from 7 to 94 seconds when using fewer to no simplification levels, while reducing the preprocessing time by only 200 milliseconds per omitted level. Despite the observed effectiveness of the algorithm, we chose not to add a fifth level: it affects the memory footprint, and performance gain decreases for additional levels.

In the next sections, we provide details for each step. Throughout, we assume that a database trajectory \(P \in \mathcal{P}\) has \(m + 1\) vertices, \(p_0, \ldots, p_m\); a query trajectory \(Q\) has \(n + 1\) vertices, \(q_0, \ldots, q_n\).

3 A SPATIAL HASH

During the preprocessing stage, we create a spatial hash. It is implemented as a regular grid, dividing the bounding box (minimal axis-aligned rectangle enclosing all trajectories in \(\mathcal{P}\)) in 500 equally-sized rows and columns, yielding 2500000 cells in total. Each cell \(c\) contains a list of vertices that have coordinates within \(c\). We place \(p_0\) and \(p_m\) into the spatial hash for each trajectory \(P \in \mathcal{P}\). The insertions take only \(O(1)\) time per trajectory and end up as a relatively insignificant amount of time with respect to the total execution time. Though parallelization could be considered, the need for synchronization in accessing the list of a cell makes for only a very small potential gain and we hence did not use concurrency here.

During the query stage, we need the spatial hash only to establish the set \(C\) of candidates with close-enough endpoints to \(Q\) in Step 1. We retrieve the trajectories \(P \in \mathcal{P}\) with \(|p_0 - q_0| \leq \epsilon\) or with \(|p_m - q_n| \leq \epsilon\), by using the vertices stored with cells in the hash that are close enough to \(q_0\) and \(q_n\), respectively. We then intersect these lists to obtain \(C\). Note that any trajectory with \(|p_0 - q_0| > \epsilon\) or \(|p_m - q_n| > \epsilon\) cannot have \(\delta_F(P, Q) \leq \epsilon\), since the endpoints must map to each other in any homeomorphism; thus any trajectory not in \(C\) is indeed not part of the answer for query \((Q, \epsilon)\).

4 SIMPLIFICATIONS

In the preprocessing stage, we simplify each trajectory \(P \in \mathcal{P}\). For this we use the algorithm by Agarwal et al. \(^2\) which computes a simplification \(T'\) of some trajectory \(T\) with \(\delta_F(T, T') \leq \epsilon\), given some threshold \(\delta > 0\). There is a direct trade off between \(\delta\) and the number of vertices of \(T'\); that is, between the power and computation time of subsequent comparisons in the query stage. We therefore construct four simplifications, with increasing level of detail. These simplifications are thus increasingly time-consuming to compute with, but also yield more more accurate results. This way, easily decidable trajectory comparisons can be discarded with little computation, while trajectories more difficult to decide still benefit from a detailed comparison.

In particular, for each trajectory \(P \in \mathcal{P}\) of size \(m\), we run four binary searches to find threshold that achieve simplifications with 0.07m, 0.19m, 0.24m, and 0.32m vertices. These values were determined experimentally by observing the algorithm behavior on our constructed test data set (see also Section 2). We denote the obtained simplifications by \(P^0, \ldots, P^3\), where \(P^0\) is the coarsest simplification, and \(P^3\) the finest. These simplifications are computed in parallel per trajectory.

We can use the simplifications in the query stage, by exploiting that the Fréchet distance is a proper metric and thus the triangle inequality can be used. First, we similarly compute simplifications \(Q^0, \ldots, Q^3\) for query trajectory \(Q\). Instead of searching for exact thresholds, we estimate good thresholds using the thresholds used to simplify \(P\). The estimation is computed by averaging, for each trajectory, the relation between the computed threshold and the diagonal of the trajectory bounding box. This reduces computation time per query, and implicitly assuming that the trajectories from \(\mathcal{P}\) and \(\mathcal{Q}\) have similar characteristics and thus can be similarly simplified. For each \(i = 0, \ldots, 3\), we do the following:

(I) Compute the equal-time distance \(d(P_i)\) between \(P_i\) and \(Q_i\) (Section 5). If \(d < \epsilon - \delta_i - \gamma_i\), then by triangle inequality we may conclude that \(\delta_F(P, Q) \leq \epsilon\).

(II) We now use our exact decision algorithm (Section 6). If \(\delta_F(P_i, Q_i) \leq \epsilon - \delta_i - \gamma_i\), we conclude that \(\delta_F(P, Q) \leq \epsilon\).

(III) Finally, we use the exact decision algorithm a second time. If \(\delta_F(P_i, Q_i) > \epsilon + \delta_i + \gamma_i\), we can conclude \(\delta_F(P, Q) > \epsilon\).

If we reach a conclusion in any of the above steps, we stop processing \(P\) and report it for the query’s answer if we conclude \(\delta_F(P, Q) \leq \epsilon\) (I or II). If we reach no conclusion, we proceed with the next more detailed level of simplification.

\(^2\)Available online at https://github.com/TWTDIG/GISCUP17TUE
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5 USING THE EQUAL-TIME DISTANCE

In Step 2a and 2b of the algorithm, we use the equal-time distance to determine an upperbound on the Fréchet distance. That is, we use a linear homeomorphism \( \psi : [0, m] \rightarrow [0, n] \) with \( \psi(t) = t \cdot n/m \), to map points on \( P \) to \( Q \). It is straightforward to compute \( d = \max_{\psi(0), \psi(n]} ||P(t) - Q(\psi(t))|| \) in linear time; and since \( \psi \in \Psi \), this readily implies that \( \delta_\Psi (P, Q) \leq d \). Thus, if \( d \leq \varepsilon \), we can immediately conclude that \( \delta_\Psi (P, Q) \leq \varepsilon \) by transitivity.

6 THE FRÉCHET DECISION PROBLEM

We need an algorithm to decide whether \( \delta_\Psi (P, Q) \leq \varepsilon \) in Step 2c and similarly for Step 2a. The decision algorithm by Alt and Godau [3] runs in \( \Theta(mn) \) time. This would be problematic for efficient processing of complex trajectories, thus especially for Step 2c—comparing the full trajectories. Below, we present an improved variant of this algorithm, used in both Step 2a and Step 2c.

The Alt and Godau algorithm conceptually works on a parameter-space representation of the problem. This is called the free-space diagram, a \([0, m] \times [0, n] \) diagram where a point \((s, t)\) represents the pair of points \( P(s) \) and \( Q(t) \). We call a point \((s, t)\) of the diagram free if \( ||P(s) - Q(t)|| \leq \varepsilon \). The free space is then the union of all free points. Alt and Godau showed that a bimonotone path from \((0, 0)\) (bottomleft of the diagram) to \((m, n)\) (topright) through the free space corresponds to a homeomorphism which acts as a witness for \( \delta_\Psi (P, Q) \leq \varepsilon \) [3]. Points \((s, t)\) that are free and reachable with such a bimonotone path are called reachable; the union of reachable points is called the reachable space. The decision problem is reduced to deciding whether \((m, n)\) is reachable.

The vertices of \( P \) and \( Q \) partition the diagram into \( m \cdot n \) cells. Alt and Godau provide a dynamic program that processes each cell to find out which points along the cell boundaries are reachable. This algorithm uses the interval of reachable points on the left and bottom side of the cell, to compute those for the top and right side.

We define the complexity of the reachable space as the number of reachable vertex pairs \( p_i, q_j \) plus the number of reachable points \((s, t)\) for which \( ||P(s) - Q(t)|| = \varepsilon \) and either \( P(s) \) or \( Q(t) \) is a vertex. To improve upon the quadratic run time, we design our method such that the run time is \( O(mn) \), that is, proportional to the complexity of the reachable space rather than the \( \Theta(mn) \) number of cells.

We first present a basic version of our algorithm, which uses queues to efficiently find the reachable space. We then describe how we re-use information from the simplification steps, to further speed up our algorithm.

6.1 Queue-based approach

Algorithm 1 provides an overview of our basic algorithm and is further explained below. For this, we assume that \( m \leq n \), and number the rows \( I \) though \( m \) (corresponding to edges of \( P \)) and the columns \( J \) through \( n \) (the edges of \( Q \)). Note that if \( m > n \), we simply swap the roles of \( P \) and \( Q \). We denote the reachable intervals on the left, right, top and bottom side of a cell \( i, j \) by \( L_{ij}, R_{ij}, T_{ij} \) and \( B_{ij} \) respectively. Note that \( R_{ij} = L_{i(j+1)} \) and \( T_{ij} = B_{i(j+1)} \).

The main idea is to extend the column-by-column dynamic program, to maintain a list of row indices with reachable intervals that we could reach in the previous column to jump over unreachable parts. That is, we use a queue \( \Lambda_j \) for each vertex \( q_j \) of \( Q \), that stores the indices of edges \( p_i, q_j \) of \( P \) that have a nonempty reachable interval with \( q_j \), in sorted order. We set \( L_0 \) to \((1)\), as we know that \((0, 0)\) is reachable. As we process a column \( j = 1, \ldots, n \), we use queue \( \Lambda_{j-1} \) and build queue \( \Lambda_j \) (see also Figure 2(a)).

We process a column as follows (Line 4 to 16). We immediately jump to the first index in \( \Lambda_{j-1} \), knowing that any earlier cells are not reachable. We then essentially start Alt and Godau’s procedure from there, going up in the column, each time processing cell \( i, j \) by computing \( R_{ij} \) and \( T_{ij} \) using \( L_{ij} \) and \( B_{ij} \) (Line 7 to 16). When we find that \( i \) matches the index stored in at the head of the queue \( \Lambda_{j-1} \), we remove it and obtain the corresponding nonempty \( L_{ij} \); otherwise, we know that \( L_{ij} \) is empty. If \( R_{ij} \neq 0 \), we add the row index \( i \) with corresponding \( R_{ij} \) to \( \Lambda_j \). We stop this process after we find that \( T_{ij} = 0 \), that is, after we conclude that we cannot reach the next cell from below. Then, if there are still elements in \( \Lambda_{j-1} \), we jump to the next index there; otherwise, we stop processing the column and move to the next, as no cell in the remainder of the current column can be reachable.

To implement the above algorithm, we observe that we only ever need \( \Lambda_{j-1} \) and \( \Lambda_j \). We therefore implement this using only two queues, using one as \( \Lambda_{j-1} \) and the other as \( \Lambda_j \); the queues swap roles when moving to the next column, to avoid copying queues.

We further improve this method by compressing the queues. Instead of remembering each reachable row index, we save space in the queue by instead storing maximal vertical intervals of reachable space, characterized by a start and end point (which may be part of different rows). At this point this optimization saves only space and queue operations; we still need to process all rows in between to check whether a long reachable interval splits into multiple. However, it in fact also allows us to do one more optimization, where we leverage information from our previous simplifications; this is described in the upcoming section.

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Algorithm 1 Decide\((P, Q, \varepsilon)\)

Given: \( P \) and \( Q \) are trajectories with \( m \) and \( n \) edges, and \( \varepsilon > 0 \)

Decide: whether \( \delta_\Psi (P, Q) \leq \varepsilon \)

1. Set \( \Lambda_0 \) to \((1)\)
2. Set \( j \) to 1
3. While \( \Lambda_{j-1} \) is not empty and \( j \leq n \)
   4. While \( \Lambda_{j-1} \) is not empty
      5. Set \( i \) to index at the head of \( \Lambda_{j-1} \)
      6. Set \( B \) to an empty interval
      7. Repeat
         8. If the head of \( \Lambda_{j-1} \) stores index \( i \) then
            9. Set \( L \) to the reachable interval at the head of \( \Lambda_{j-1} \)
            10. Remove the head of \( \Lambda_{j-1} \)
            11. Else
            12. Set \( L \) to an empty interval
            13. Use \( L \) and \( B \) to compute reachable intervals \( R \) and \( T \)
            14. If \( R \neq \emptyset \), add \((i, R)\) to queue \( \Lambda_j \)
            15. Set \( B \) to \( T \) and increase \( j \) by one
            16. Until \( B = \emptyset \)
            17. Increase \( j \) by one
   18. Return true if \( m \) is contained in \( \Lambda_n \)
We use information from our simplification steps to define free-space jumps. These jumps allow us to find sequences of connected reachable rows in $\Lambda_j$ without processing each row individually, given certain preconditions. In our tests on the sample data sets, the application of free-space jumps reduced the number of cells processed, and so the number of steps in our algorithm by a factor of five, while requiring little additional storage or computation.

During trajectory simplification, we have calculated the Fréchet distance of many subtrajectories to the source trajectory in order to find a simplification matching the given threshold. We can use the (normally discarded) Fréchet distance of these subtrajectories to skip computations in the decision Fréchet problem. Given start and end indices $(s, e)$ of a subtrajectory and an accompanying Fréchet distance $d$, we can skip the calculation of the rows between $s$ and $e$ in any decision Fréchet problem if the following conditions hold:

1. Interval $(s, e)$ is contained in a single interval in queue $\Lambda_{j-1}$.
2. $\max(\delta(s, j), \delta(j, e)) + d \leq \epsilon$

The first condition ensures that skipping range $(s, e)$ does not result in a loss of information. That is, by having $(j - 1, e)$ reachable, we do not need to propagate the reachability information up through all these cells: the top of each of the intermediate cells is fully reachable by this condition. The second condition ensures that the jump is valid: the range $(s, e)$ on column $j$ is part of the free-space, and so can be skipped. This is illustrated in Figure 2(b). Skipping these calculations as described constitutes a free-space jump. As only jumps along a column are possible, only one of the trajectories needs to retain the subtrajectory Fréchet computation results.

6.3 Performance

Though a theoretic difference between $O(mn)$ and $O(mn)$ may seem minor, it has a major impact on our running time. We explain this as follows. When the need arises to compare two trajectories in Step 2c, we can safely assume that the trajectories are very similar: otherwise, a previous step would have likely already been able to give us a definite answer for the comparison. With the two trajectories being very similar, the reachable-space complexity is likely much lower than quadratic; this makes our optimized version efficient, even when executed with two full trajectories. Note that this is not a hard guarantee: trajectories with a high reachable-space complexity may remain, even after all pruning steps.

7 FURTHER CONSIDERATIONS

Our method uses various parameters (e.g. spatial hash size, number of simplifications and their thresholds). We set these based on experiments with several example data sets. However, more experiments could be set up to further fine-tune these parameters and investigate how characteristics of the data sets may inform these parameter settings; the latter would allow us to move towards an approach that adapts its parameters to fit the actual data set.

We use equal-time distance as a simple greedy approach, but various alternatives could be considered. On the sample data set, it decided only 25% of the cases that made it to Step 2b. Other heuristic methods could be considered, if they have significantly better pruning percentage and/or average running time.

Our approach to the decision Fréchet problem requires little storage in practice compared to other column-based approaches. This results in efficient CPU processing, but also makes it a good candidate for a GPU-based implementation, where IO is often a bottleneck and memory is expensive.

Finally, the free-space jumps can be used to convert the decision Fréchet problem into a graph problem, where the edges are pairs of jumps and reachability is computed by traversing this graph. We expect that other algorithms may be developed in this new problem space, especially greedy approximation algorithms.

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