Towards Efficient Indexing of Spatiotemporal Trajectories on the GPU for Distance Threshold Similarity Searches

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Abstract

Applications in many domains require processing moving object trajectories. In this work, we focus on a trajectory similarity search that finds all trajectories within a given distance of a query trajectory over a time interval, which we call the distance threshold similarity search. We develop three indexing strategies with spatial, temporal and spatiotemporal selectivity for the GPU that differ significantly from indexes suitable for the CPU, and show the conditions under which each index achieves good performance. Furthermore, we show that the GPU implementations outperform multithreaded CPU implementations in a range of experimental scenarios, making the GPU an attractive technology for processing moving object trajectories. We test our implementations on two synthetic and one real-world dataset of a galaxy merger.

1 Introduction

Trajectory data is generated in a wide range of application domains, such as the motions of people or objects captured by global positioning systems (GPS), the movement of objects in scientific applications, such as stars in astrophysical simulations, vehicles in traffic studies, animals in zoological studies and a range of applications of geographical information systems (GIS). We study historical continuous trajectories [6], where a database of trajectories is given as input and is searched to gain domain-specific insight. In particular, we study the distance threshold search: Find all trajectories within a distance \( d \) of a given query trajectory over a time interval \([t_{start}, t_{end}]\). An example of this search would be to find all prey within 200 m of all predators over the period of a month.

The challenges associated with moving object trajectories in comparison to stationary objects has prompted a literature on efficient trajectory indexing and processing strategies. Many of the methods developed by the spatial and spatiotemporal database communities focus on sequential implementations, where a fraction of the data is stored in memory, and the rest is stored on disk. Thus, reduction of disk accesses is the main optimization goal in these works. Alternatively, with relatively large memories available in modern workstations, sizable in-memory databases have become feasible. Furthermore, with the proliferation of multicore and manycore architectures, parallel in-memory implementations can provide
significant performance improvements over sequential out-of-core implementations. In instances where memory capacity on a single node is insufficient, historical continuous trajectory datasets can be partitioned and queried in-memory across multiple compute nodes in parallel.

To this end, we study the efficient processing of distance threshold searches on trajectory databases using General Purpose Computing on Graphics Processing Units (GPGPU). We focus on developing and comparing the performance of GPU-friendly indexing strategies, and make the following contributions:

- We develop three indexing techniques that are suitable for distances threshold searches on the GPU.
- For each of the indexes, we develop an associated GPU kernel that minimizes branch instructions to achieve good parallel efficiency.
- We compare our GPU implementation to a previously developed CPU-only implementation that uses an in-memory R-tree index, and show that using the GPU can afford significant speedup.
- We find that when using large datasets, in contrast to smaller datasets previously used in the literature, efficient trajectory splitting strategies for an R-tree index, at least for the in-memory case, provides limited or no performance improvements.
- We evaluate our algorithms and kernel implementations with 4-D datasets (3 spatial dimensions and 1 temporal dimension), including a real-world astrophysics dataset (of a galaxy merger) and two synthetic datasets.

The paper is outlined as follows: Section 2 outlines a motivating example and discusses related work. Section 3 formally defines the problem. Section 4 describes our three indexing techniques and search algorithms. Section 5 presents our experimental results. Finally, Section 6 concludes with a summary of our findings and a discussion of future research directions.

2 Background and Motivating Example

2.1 Motivating Example

One motivating application for this work is in the area of astrophysics/astrobiology [10]. Astrobiology is the study the evolution, distribution and future of life in the universe. Biologists study the habitability of the Earth and find that life can exist in a multitude of environments (including extreme environments, such as temperature, pressure, salinity, radiation exposure, and others). The past decade of exoplanet searches implies that the Milky Way, and hence the universe, hosts many rocky, low mass planets that may be capable of supporting complex life (land-based animal life). Given that there are many planets in the Milky Way and given the broad range of conditions in which life is found to thrive on Earth, the notion of the Galactic Habitable Zone has emerged, i.e., the region(s) of the Galaxy that
may favor the development of complex life. With regards to long-term habitability, some regions of the Milky Way may be inhospitable due to transient radiation events, such as supernovae explosions or close encounters with flyby stars that can gravitationally perturb planetary systems. Studying habitability thus entails solving the following two types of distance threshold searches on the trajectories of (possibly billions of) stars orbiting the Milky Way: (i) Find all stars within a distance $d$ of a supernova explosion (or gamma ray burst), i.e., a non-moving point over a time interval; and (ii) Find the stars, and corresponding time periods, that host a habitable planet and are within a distance $d$ of all other stellar trajectories.

2.2 Background and Related Work

A key question in database research is the efficient retrieval of data. In the most general context, database management systems provide information about database content and support arbitrary queries. However, in specific domains it is possible to achieve more efficient retrieval if there are structures and constraints on the data stored in the database and/or if particular types of queries are expected. Such a domain is that of spatial and spatiotemporal databases that store the trajectories of moving objects. A trajectory is a collection of points associated with the positions of an object over time, where the points are connected by polylines (line segments). Such data, which arises in many scientific domains but is also pervasive in modern society (GPS data, GIS applications), presents both opportunities and challenges that are studied in the spatiotemporal database community. The main goal of these databases is to perform trajectory similarity searches, i.e., finding trajectories within a database that exhibit similarity in terms of spatial and/or temporal proximity, or exhibit similarity in terms of spatial and/or temporal features so that trajectories can be classified as belonging to a certain group. Similarity searches have been studied in various domains, such as convoys [18], flocks [30], and swarms [22]. A predominant trajectory similarity search that is used in many application areas is the kNN (k Nearest Neighbors) search [7, 5, 8, 14].

The typical approach in previous spatiotemporal database works proceeds in two phases: (i) search an index to obtain a preliminary result set; (ii) use refinement to produce the final result set. The search phase focuses on pruning, i.e., avoiding parts of the index based on the selecting criteria of the query. To this end, several index-trees have been proposed inspired by the success of the popular R-tree [15], such as TB-trees [25], STR-trees [25], 3DR-trees [28], SETI [3], and implemented in systems such as TrajStore [4] and SECONDO [14]. More specifically, these works map nodes in an index-tree to pages stored on disk. Performance is a function of the number of index-tree nodes that are accessed, aiming to keep this number low so as to avoid avoiding costly data transfers between memory and disk. Index-trees have been used extensively for kNN searches.

In this work we study distance threshold searches, which can be viewed as kNN searches with an unknown value of $k$ and thus unknown result set size. As a result, several of the aforementioned index-trees, while efficient for kNN searches, are not efficient for distance threshold searches. This is because, as $k$ is unbounded, standard index pruning methods cannot be used. Distance-threshold searches, although relevant to several application domains, have not received a lot of attention in the literature. Our previous work in [12] studies in-memory sequential distance threshold searches, using an R-tree to index trajectories inside
hyperrectangular minimum bounding boxes (MBBs). The main contribution therein is an indexing method that achieves a desirable trade-off between the index overlap, the number of entries in the index, and the overhead of processing candidate trajectory segments. The work in [1] solves a similar problem, i.e., finding trajectories in a database that are within a query distance \( d \) of a search trajectory and the authors propose four query processing strategies. A key difference with the work in [12] is that part of the database resides on disk. Other trajectory similarity searches rely on metrics of similarity at coarse grained resolutions [9]. Instead, the similarity search we study in this work necessitates precise comparisons between individual polylines, to find the exact time intervals when trajectories are within the threshold distance. The large number of such comparisons is a motivation for using the GPU.

In the context of in-memory moving object trajectory databases several authors have explored the use of multicore and manycore architectures. Spatial and spatiotemporal indexing methods have been advanced for use on the GPU [33, 32, 31, 23]. Given the single instruction multiple data (SIMD) nature of the GPU, proposed indexes for this architecture tend to be less sophisticated than the index-trees used in the context of out-of-core databases. This is in part because branches in the instruction flow cause thread serialization and thus loss of parallel efficiency [17]. The \( k \)NN query (not on trajectories) has been studied in the context of the GPU [24, 20] and on hybrid CPU-GPU environments [21]. In this work we focus on indexing techniques for distance threshold similarity searches on trajectories for the GPU, which to our knowledge has only been explored in our previous work [11]. That previous work focuses on a scenario in which the query set cannot fit entirely on the GPU due to memory constraints, thereby requiring back-and-forth communication between the host and GPU, and thus a particular indexing scheme. Instead, in this work, the query set fits on the GPU, which makes it possible to explore a range of indexing schemes (while still having to consider memory constraints).

3 Problem Statement

3.1 Problem Definition

Let \( D \) be a spatiotemporal database that contains \( n \) 4-dimensional (3 spatial and 1 temporal dimensions) entry line segments. A line segment \( l_i, i = 1, \ldots, |D| \), is defined by a spatiotemporal start point \((x_{i}^{\text{start}}, y_{i}^{\text{start}}, z_{i}^{\text{start}}, t_{i}^{\text{start}})\), an end point \((x_{i}^{\text{end}}, y_{i}^{\text{end}}, z_{i}^{\text{end}}, t_{i}^{\text{end}})\), a segment id and a trajectory id. Segments belonging to the same trajectory have the same trajectory id and are ordered temporally by their segment ids. We call \( t_{i}^{\text{end}} - t_{i}^{\text{start}} \) the temporal extent of \( l_i \).

The distance threshold search searches for entry segments within a distance \( d \) of a query set \( Q \), where \( Q \) is a set of line segments that belong to a series of moving object trajectories. We call the line segments in \( Q \) query segments and denote them by \( q_k, k = 1, \ldots, |Q| \). The search is continuous, such that an entry segment may be within the distance threshold \( d \) of particular query segment for only a subinterval of that segment’s temporal extent. We call a comparison between an entry segment and a query segment an interaction. The result set thus contains a set of query and entry segment pairs, and for each pair the time interval
during which the two segments are within a distance $d$ of each other. For example, a search may return $(q_1,l_1,[0.1,0.3])$ and $(q_1,l_2,[0.5,0.95])$, for a query segment $q_1$ with temporal extent $[0,1]$.

We consider a platform that consists of a host, with RAM and CPUs, and a GPU with its own memory and Streaming Multi-Processors (SMPs) connected to the CPU via a (PCI Express) bus. We consider an in-memory database, meaning that $D$ is stored once and for all in global memory on the GPU, i.e., the database is stored once and queried multiple times. The objective is to minimize the response time for processing the queries in $Q$. This is the typical objective considered in other spatiotemporal database works such as the ones reviewed in Section 2. We consider the case in which both $D$ and $Q$ can fit in GPU memory. This means that GPU memory is large enough and not shared with other users. Our intended scenario is that of a distributed memory environment in which a number of GPU-equipped compute nodes are reserved by a user.

### 3.2 Memory Management on the GPU

Previous works on indexing trajectories for the purpose of distance threshold similarity searches have targeted multi-core CPU implementations [12, 13, 1] and GPU implementations [11]. CPU implementations rely on (in-memory) index trees that have been used traditionally for out-of-core implementations, such as the R-tree [15]. Each thread traverses the tree and creates a candidate segment set to be further processed to create the final result set. Although many candidate segments are not within distance $d$ of the query segments due to the “wasted space” in the index (an unavoidable consequence of using MBBs) [12], memory must still be allocated to store these candidate segments. Furthermore, the size of the final result set is non-deterministic as it depends on the spatiotemporal nature of the data. Consequently, memory allocation for the result set must be conservative and overestimate the memory required (this overestimation grows linearly with $|Q|$). On the CPU these memory management issues are typically not problematic in practice since the number of threads is limited (e.g., set to the number of physical cores) and the memory is large.

On the GPU, even though we assume that both $D$ and $Q$ fit in memory, the same memory management issues are problematic. This is because we have a large number of threads that each need memory to store candidate segments, in addition to the memory needed to store the final result set. To address this issue of non-deterministic storage requirements, on the GPU one must define a fixed size for a statically allocated memory buffer for each thread. If the memory requirements exceed this buffer then it is necessary to perform a series of kernel invocations so as to “batch” the generation of the candidate sets and the final result set.

### 4 Indexing Trajectory Data

In this section we outline three trajectory indexing techniques for the GPU. For each we discuss shortcomings and possible solutions regarding the memory management issues discussed in Section 3.2. Although our GPU implementations use OpenCL, in what follows we use the more common CUDA terminology to describe our algorithms (GPU as opposed to device, kernel as opposed to program, thread as opposed to work-item, etc.).
4.1 Spatial Indexing: Flatly Structured Grids

Previous work has proposed the use of grid files, or “flatly structured grids” (FSG), to index trajectory data on the GPU spatially [32]. In that work the authors focus on 2-D spatial data (and Hausdorff distance) while our context is 3-D spatiotemporal data (and Euclidian distance). An interesting question is whether spatial indexing with FSGs is effective even when the data has a temporal dimension. In what follows we describe an FSG indexing scheme and accompanying search algorithm for the GPU. We call this approach GPUSpatial.

4.1.1 Trajectory Indexing

We define a FSG as a 3-D rectangular box partitioned into cells with \( \text{grid}_x, \text{grid}_y, \text{grid}_z \) cells in the \( x, y, \) and \( z \) spatial dimensions, respectively, for a total of \( \text{grid}_x \times \text{grid}_y \times \text{grid}_z \) cells. Each line segment \( l_i \) in \( D \) is contained in a spatial MBB defined by two points \( \text{MBB}_{\text{min}}^i \) and \( \text{MBB}_{\text{max}}^i \), where

\[
\text{MBB}_{\text{min}}^i(x) = \min(x_{\text{start}}^i, x_{\text{end}}^i), \quad \text{MBB}_{\text{min}}^i(y) = \min(y_{\text{start}}^i, y_{\text{end}}^i), \quad \text{MBB}_{\text{min}}^i(z) = \min(z_{\text{start}}^i, z_{\text{end}}^i)
\]

and

\[
\text{MBB}_{\text{max}}^i(x) = \max(x_{\text{start}}^i, x_{\text{end}}^i), \quad \text{MBB}_{\text{max}}^i(y) = \max(y_{\text{start}}^i, y_{\text{end}}^i), \quad \text{MBB}_{\text{max}}^i(z) = \max(z_{\text{start}}^i, z_{\text{end}}^i)
\]

Each line segment is assigned to the FSG by rasterizing its MBB to grid cells. Figure 1 shows a 2-D example for two line segments and a 5 \( \times \) 5 FSG. Each line segment may occupy more than one grid cell, and some grid cells can remain empty. We store the FSG as an array of non-empty cells, \( G \). Each cell is defined by \( h \), and by an index range \([A_{\text{min}}^h, A_{\text{max}}^h]\) in an additional integer “lookup” array, \( A \). \( A[A_{\text{min}}^h : A_{\text{max}}^h] \) contains the indices of the line segments whose MBBs overlap cell \( C_h \) (the notation \( X[a:b] \) is used to denote the “slice” of array \( X \) from index \( a \) to index \( b \), inclusive). In other terms, if \( l_i \)'s MBB overlaps \( C_h \), then \( i \in A[A_{\text{min}}^h : A_{\text{max}}^h] \).

Since the MBB of line segment \( l_i \) can overlap multiple grid cells, \( i \) can occur multiple times in array \( A \). Figure 2 shows an example to highlight the relationship between \( G, A, \) and \( D \). This example is discussed in the next section.

One of the objectives of the above design is to reduce the memory footprint of the index. This is why we only index non-empty grid cells, and why for each cell \( C_h \) we do not store its spatial coordinates but instead compute \( h \) whenever needed (thereby trading off memory space for computation time). Furthermore, the use of lookup array \( A \) makes it possible for array \( G \) to consist of same-size elements (even though some cells contain more line segments than others). Without this extra indirection through lookup array \( A \), it would have been necessary to store entry segment ids directly into the elements of \( G \). However, it would have been necessary to pick an element size large enough to accommodate the cell with the largest
Figure 2: Example relationship between the grid \((G)\), the lookup array \((A)\) and the database of entry line segments \((D)\) in the GPUSpatial approach.

number of entry segments, thereby wasting memory space. \(D\), \(A\), and \(G\) are stored in GPU memory before query processing begins.

4.1.2 Search Algorithm

The trajectory segments in \(Q\) are not sorted by any spatial or temporal dimension. This is because sorting segments temporally would not be effective when using a spatial index. Regarding spatial sorting, it is not clear by which dimension the segments should be sorted. However, segments that are part of the same query trajectory are stored contiguously, thus providing a natural advantageous ordering of data elements. Each query segment \(q_k\) is assigned to a GPU thread. The kernel first calculates the MBB for \(q_k\) and the FSG cells that overlap this MBB. Given the \(x\), \(y\), \(z\) coordinates of each such cell in the FSG, the kernel computes its linearized coordinate \((h)\) using a row-major order. A binary search is used to find whether cell \(C_h\) occurs in array \(G\), in \(O(\log |Q|)\) time. In this manner the kernel creates a list of non-empty cells that overlap \(q_k\)’s MBB. For each cell \(C_h\) in this list, the indices of the entry segments it contains are computed as \(A[A^\text{min}_h : A^\text{max}_h]\). These indices are appended to a buffer \(U_k\). A key point here is that with a spatial indexing scheme there is no good approach for storing index entry segments in a contiguous manner (since one would have to arbitrarily pick one of the spatial dimensions). This is why we must resort to using buffer \(U_k\) as opposed to, for instance, a 2-integer index range in a contiguous array of entry segments. Each entry in \(U_k\) is then compared to the query segment \(q_k\) to see if it is within the threshold distance; however, note that while the segments are expected to be relatively nearby each other spatially (given their FSG overlap), they may not overlap temporally.

Consider the example in Figure 2 which shows partial contents of arrays \(G\), \(A\), and \(D\). Consider a query \(q_1\) (not shown in the figure), which overlaps grid cells \(C_0\), \(C_1\), and \(C_7\). Cell \(C_1\) is not in \(G\), meaning that it contains no entry segments. Therefore, the only two cells to consider are \(C_0\) and \(C_7\), which have \([A^\text{min}_h : A^\text{max}_h]\) values of \([0,2]\) and \([25,90]\), respectively. In lookup array \(A\), we find that \([0,2]\) corresponds to entries 2, 100, and 22, while \([25,90]\) corresponds to entry indices 100, 867, \ldots, 400. These indices are copied from \(A\) into buffer \(U_k\). Note that in this step the search algorithm does not remove duplicate indices (such as entry index 100 in this example) and thus may perform some redundant entry segment processing. Removing duplicates would amount to sorting buffer \(U_k\), as done for instance in
which thus comes at an additional computational cost that may offset the benefits of removing redundant segment processing.

Since the number of entry segments that overlap $q_k$’s MBB can be arbitrary large (it depends on the spatial features of $D$ and $Q$, and on the query distance $d$), the use of buffer $U_k$ creates memory pressure, especially since both $D$ (along with $G$ and $A$) and $Q$ are stored on the GPU. This same issue has been encountered in previous work, e.g., when using a parallel R-tree index on the GPU [23]. We define an overall buffer size, $s$, that is split equally among all queries ($|U_k| = s/|Q|$). If the processing of query $q_k$ exceeds the capacity of $U_k$, then the thread terminates, and stores the query id into an array that is sent back to the host. Once the kernel execution finishes, the host re-attempts the execution of those queries that could not complete due to memory pressure. In this re-attempt, memory pressure is lower because fewer queries are executed (i.e., $|U_k|$ is larger). This method implicitly has the effect that threads with similar (large) amounts of work to execute together, resulting in improved load-balancing.

Algorithm 1 GPUSpatial kernel.

```plaintext
1: procedure SearchSpatial(G, A, D, Q, queryIDs, U, d, redo, resultSet)
2:  gid ← getGlobalId()
3:  if queryIDs = ∅ and gid≥|Q| return
4:  if queryIDs ≠ ∅ and gid≥|queryIDs| return
5:  if queryIDs = ∅ then
6:      queryID ← gid
7:  else
8:      queryID ← queryIDs[gid]
9:  end if
10: (overflow , candidateSet) ← getCandidates(G, A, D, Q[queryID], U, d)
11:  if overflow then
12:      atomic: redo ← redo ∪ { queryID }
13:      return
14:  end if
15:  for all entryID ∈ candidateSet do
16:      result ← compare(D[entryID], Q[queryID])
17:      if result ≠ ∅ then
18:          atomic: resultSet ← resultSet ∪ result
19:      end if
20:  end for
21:  return
22: end procedure
```

The pseudo-code of the search algorithm is shown in Algorithm 1. It takes the following arguments: (i) the FSG array ($G$); (ii) the lookup array ($A$); (iii) the database ($D$); (iv) the set of queries ($Q$); (v) an array that contains the ids of the queries to be reprocessed ($queryIDs$), which is empty for the first kernel invocation; (vi) buffer space ($U$); (vii) the query distance ($d$); (viii) an output array in which the kernel stores the ids of the queries that must be reprocessed ($redo$); and (ix) the memory space to store the result set ($resultSet$). Arguments that lead to array transfers between the host and the GPU, either as input or
output, are shown in boldface. Other arguments are either pointers to pre-allocated zones of (global) GPU memory or integers. The algorithm begins by checking the global thread id and aborts if it is greater than \( Q \) or \(|\text{queryIDs}|\), depending on whether this is a first invocation or a re-invocation (lines 3|4). The id of the query assigned to the GPU thread is then acquired from \( Q \) or using an indirection via \( \text{queryIDs} \) (lines 8|8). Function \text{getCandidates} searches the FSG and returns a boolean that indicates whether buffer space was exceeded and the (possibly empty) set of candidate entry segment ids (line 10). If buffer space was exceeded, then the query id is atomically added to the \text{redo} array and the thread terminates (line 11|13). The algorithm then loops over all candidate entry segment ids (line 15), compares each entry segment spatially and temporally to the query (line 16) and atomically adds a query result, if any, to the result set (line 18). Once all GPU threads have completed, \text{resultSet} and \text{redo} are transferred back to the host. If \|\text{redo}\| \) is non-zero, then the kernel is re-invoked, passing \text{redo} as \( \text{queryIDs} \). Duplicates in the result set are filtered out on the host.

### 4.2 Temporal Indexing

In this section we propose a purely temporal partitioning strategy, which we call \text{GPUTEMPORAL}.

#### 4.2.1 Trajectory Indexing

We begin by sorting the entries in \( D \) by ascending \( t_{\text{start}} \) values, re-numbering the entry segments in this order, i.e., \( t_{i,\text{start}} \leq t_{i+1,\text{start}} \). The full temporal extent of \( D \) is \([t_{\text{min}}, t_{\text{max}}]\) where \( t_{\text{min}} = \min_{i\in D} t_{i,\text{start}} \) and \( t_{\text{max}} = \max_{i\in D} t_{i,\text{end}} \). We divide this full temporal extent so as to create \( m \) logical bins of fixed length \( b = (t_{\text{max}} - t_{\text{min}})/m \). We assign each entry segment, \( l_i \), \( i = 1, \ldots, |D| \), to a bin, where \( l_i \) belongs to bin \( B_j \), \( j = 1, \ldots, m \), if \( \lfloor t_{i,\text{start}}/b \rfloor = j \). There can be temporal overlap between the line segments in adjacent bins. For each bin \( B_j \) we defined its start times as \( B_{j,\text{start}} = j \times b \) and its end time as \( B_{j,\text{end}} = \max((j + 1) \times b, \max_{i\in B_j} t_{i,\text{end}}) \). \( B_{j,\text{start}} \) does not depend on the line segments in bin \( B_j \), but \( B_{j,\text{end}} \) does. The temporal extent of bin \( B_j \) is defined as \([B_{j,\text{start}}, B_{j,\text{end}}]\). Given the definitions of \( B_{j,\text{start}} \) and \( B_{j,\text{end}} \), the union of the temporal extents of the bins is equal to the full temporal extent of \( D \). We define \( B_{j,\text{first}} = \arg\min_{i\in B_j} t_{i,\text{start}} \) and \( B_{j,\text{last}} = \arg\max_{i\in B_j} t_{i,\text{start}} \), i.e., the ids of the first and last entry segments in bin \( B_j \), respectively. \([B_{j,\text{first}}, B_{j,\text{last}}]\) forms the index range of the entry segments in \( B_j \). Bin \( B_j \) is thus fully described as \((B_{j,\text{start}}, B_{j,\text{end}}, B_{j,\text{first}}, B_{j,\text{last}})\). The set of bins forms the temporal database index.

Figure 3 shows an example of how line segments may be assigned to a set of temporal bins. In this example, 15 entry segments are assigned to 4 temporal bins over a database temporal extent of 12 time units (spatial dimensions are ignored, and thus line segments are simply represented as horizontal lines in the figures). The \( B_{\text{start}}, B_{\text{end}}, B_{\text{first}} \) and \( B_{\text{last}} \) values are shown for each bin. For instance, three entry segments are assigned to bin \( B_2 \): \( l_9, l_{10} \), and \( l_{11} \). Thus, \( B_{2,\text{first}} = 9 \) and \( B_{2,\text{last}} = 11 \). \( B_{2,\text{start}} = 2 \times (12/4) = 6 \) and \( B_{2,\text{end}} = t_{11} \).
4.2.2 Search Algorithm

Before performing the actual search, the following pre-processing steps must be performed. First, query segments in $Q$ are sorted by non-decreasing $t_{start}$ values, in $O(|Q| \log |Q|)$ time. For each query segment $q_k$, we calculate the index range of the contiguous bins that it overlaps temporally. A naïve algorithm for computing this overlap would be to scan all bins in $O(m)$ time. A binary search could be used to obtain a logarithmic time complexity. In practice, however, there are many temporally contiguous query segments and each overlaps only a few bins. Since segments in $Q$ are sorted by non-decreasing $t_{start}$, the search can be done efficiently by using the first temporal bin that overlaps the previous query segment as the starting point for the scan for the temporal bins that overlap the next query segment. The search thus typically takes near-constant time. Let $B_k$ denote the set of contiguous bins that temporally overlap query segment $q_k$, as identified by the above search. In constant time we can now compute the index range of the entry line segments that may overlap and must be compared with $q_k$: $E_k = [\min_{B \in B} B_{first}, \max_{B \in B} B_{last}]$. We term the mapping between $q_k$ and $E_k$ the schedule, $S$. Each GPU thread compares a single query to the line segments in $D$ whose indices are in the $E_k$ range. Assuming that $|Q|$ is moderately large, one is then insured that all GPU cores can be utilized.

In our implementation, all preprocessing described in the previous paragraph is performed on the CPU. Some of this preprocessing could be performed on the GPU (e.g., sorting the query segments). In an initial implementation, we performed the calculation of $E_k$ on the GPU; however, this did not result in any performance improvement. As explained earlier, on the host the search for temporally overlapping bins can be drastically improved by relying on the same search for the previous query segment. However, this cannot be implemented on the GPU as it would require thread synchronization and communication, which cannot be performed across thread blocks. In all of our experiments, the time to compute $S$ on the CPU is a negligible portion of the overall query response time.

The pseudo-code of the search algorithm is shown in Algorithm 2. It takes the following arguments: (i) the database ($D$); (ii) the query set ($Q$); (iii) the schedule ($S$); (iv) the query distance ($d$); and (v) the memory space to store the result set ($resultSet$). As in Algorithm 1, arguments that lead to array transfers between the host and the GPU are shown in boldface.
Algorithm 2 GPUTemporal kernel.

1: procedure SearchTemporal(D, Q, S, d, resultSet)
2:     gid ← getGlobalId()
3:     if gid ≥ |Q| return
4:     queryID ← gid
5:     entryMin ← S[gid].EntryMin
6:     entryMax ← S[gid].EntryMax
7:     for all entryID ∈ {entryMin,...,entryMax} do
8:         result ← compare(D[entryID], Q[queryID])
9:         if result ≠ ∅ then
10:             atomic: resultSet ← resultSet ∪ result
11:         end if
12:     end for
13:     return
14: end procedure

The algorithm first checks the global thread id and aborts if it is greater than |Q| (line 3). The query assigned to the thread is then acquired from Q (line 4). Next, the algorithm retrieves the minimum and maximum entry segment indices from the schedule (lines 5-6). From line 7 to 13 the algorithm then operates as Algorithm 1.

4.3 Spatiotemporal Indexing

In the two previous sections we have proposed a purely spatial and a purely temporal indexing scheme. The spatial scheme leads to segments in Q and D being compared that are spatially relevant but may be temporal misses (no temporal overlap). Likewise, the temporal indexing scheme compares temporally relevant segments in Q and D, but these segments may be spatial misses (no spatial overlap). Therefore, either approach can outperform the other depending on the spatiotemporal characteristics of Q and D. Assuming for the sake of discussion that these characteristics do not give any such particular advantage to either one of the two indexing approaches, we can reason about their relative performance. First, the spatial indexing approach requires buffer space to store the spatially overlapping trajectory segments. In contrast, because the temporal indexing scheme is indexed in a single dimension, the temporally overlapping entry segments can be defined by an index range in D, which represents significant memory space savings. The same method could possibly be used with a spatial indexing scheme if considering only one of the spatial dimensions, making the index no longer a multi-dimensional grid, but instead a linear array. This approach would however drastically decrease the spatial selectivity of the search, leading to large increases in wasted computational effort (i.e., comparisons of segments that have no overlap in one or two of the spatial dimensions). Second, to minimize the memory footprint on the GPU, the spatial scheme requires two additional arrays (G and A), thus leading to two indirections in global GPU memory. In contrast, the temporal scheme requires a single indirection. Moreover, the entry segments are stored contiguously in the temporal scheme, while this is not the case in the spatial scheme.

Given the features of both the spatial and the temporal indexing scheme, we attempt
to find an alternate spatiotemporal index that retains the benefit of both schemes without some of the drawbacks mentioned above. We term this approach GPUSpatioTemporal.

4.3.1 Trajectory Indexing

GPUSpatioTemporal adopts a temporal index so as to avoid the buffering and multiple indirection issues of spatial indexing, but subdivides each temporal bin into spatial subbins to achieve spatial selectivity. Entry segments in $D$ are assigned to $m$ temporal bins exactly as for GPUTemporal. We then compute the spatial extent of $D$ in each dimension. For instance, in the $x$ dimension the extent of $D$ is:

$$[x_{\text{min}}, x_{\text{max}}] = \left[ \min_{l \in D} \min(x^l_{\text{start}}, x^l_{\text{end}}), \max_{l \in D} \max(x^l_{\text{start}}, x^l_{\text{end}}) \right].$$

Spatial extents in the $y$ and $z$ dimensions are computed similarly. We then compute the maximum spatial extent in each dimension of the entry segments, which for the $x$ dimensions is $\max_{l \in D} |x^l_{\text{start}} - x^l_{\text{end}}|$. Maximum spatial extents are computed similarly for the $y$ and $z$ dimension. For each of the temporal bins, we create $v$ spatial subbins along each dimension, with the constraints that these subbins are larger than the maximum spatial extent of the entry segments. For instance, in the $x$ dimension, this constraint is expressed as $v \leq (x_{\text{max}} - x_{\text{min}})/\max_{l \in D} |x^l_{\text{start}} - x^l_{\text{end}}|$. We place this constraint for two reasons, which will be clarified when we describe the search algorithm: (i) to eliminate duplicates in the result set, and (ii) to reduce the amount of redundant information in the index. In total we have $m \times v$ subbins and we denote each subbin as $\hat{B}_{i,j}$, with $i = 1, \ldots, m$ and $j = 1, \ldots, v$.

The part of Figure 4 above the dashed line shows an example of how entry line segments are logically assigned to bins and subbins. The very top of the figure shows $m = 3$ temporal bins, $B_0$ to $B_2$. Each temporal bin contains contains the segments with ids in the range $[B^1_{j,\text{first}}, B^1_{j,\text{last}}]$. For instance, $B^1_{1,\text{first}} = 4$ and $B^1_{1,\text{last}} = 7$. Each entry segment is described by an id and 2 spatial ($x, y, z$) extremities. For instance, segment $l_6$ is in temporal bin $B_1$ and its spatial extremities are (8, 9, 10) and (10, 9, 8). Temporal dimensions are omitted in the figure. Below the temporal bins, we depict 9 temporal spatial subbins, $\hat{B}_{0,j}$ to $\hat{B}_{2,j}$ ($v = 3$ subbins per temporal bin). For each subbin, we indicate its spatial range in the $x$, $y$, and $z$ dimension. Each subbin spans 4 spatial units in the $x$ and $y$ dimensions, and 5 spatial units in the $z$ dimension. Given segment lengths in the database these subbin dimensions meet the constraints described in the previous paragraph. For each subbin and each dimension, we show the overlapping entry segment ids. For instance, consider subbin $\hat{B}_{0,1}$. It is overlapped in the $x$ dimension by $l_0$, $l_2$ and $l_3$, in the $y$ dimension by $l_3$, and in the $z$ dimension by $l_1$ and $l_3$.

The part of Figure 4 below the dashed line shows how the logical assignment of segments to spatial subbins is implemented physically in memory. We create three integer arrays, $X$, $Y$, and $Z$, depicted at the bottom of the figure. Each array stores the ids of the line segments that overlap the subbins in one spatial dimension. The ids for a subbin are stored contiguously, for the subbins $\hat{B}_{i,j}$’s sorted by $(j, i)$ lexicographical order. This is illustrated using colors in the figure and amounts to storing contiguously all ids in the first subbins of the temporal bins, then all ids in the second subbins of the temporal bins, etc. For instance, for the $y$ dimension, the $Y$ array in our example consists of $v = 3$ chunks.
first chunk corresponds to the ids in subbins \( \hat{B}_{0,0} (l_0, l_1, l_3), \hat{B}_{1,0} (l_4, l_5, l_8), \) and \( \hat{B}_{2,0} (l_9), \) the second chunk corresponds to the ids in subbins \( \hat{B}_{0,1} (l_3), \hat{B}_{1,1} (l_5, l_7), \) and \( \hat{B}_{2,1} (l_8), \) and the third chunk corresponds to the ids in subbins \( \hat{B}_{0,2} (\text{none}), \hat{B}_{1,2} (l_6), \) and \( \hat{B}_{2,2} (l_8). \) The reason for storing the ids in this manner is as follows. Consider a query segment with some spatial and temporal extent. This query may overlap several contiguous temporal bins (as shown in Section 4.2). However, because of the way in which we choose the sizes of the spatial subbins, most queries will not overlap multiple subbins in all three dimensions. Identifying potential overlapping entry segments then amounts to examining the \( i \)-th subbin of contiguous temporal bins, for some \( 0 \leq i \leq v. \) In other words, based on the example in Figure 4, this amounts to examining sequences of same-color subbins.

Given the \( X, Y, \) and \( Z \) array, each spatial subbin is then described with the index range of the entries in those arrays, i.e., 6 integers. For instance, consider subbin \( \hat{B}_{0,1} \) in our example. Its description is index range \( 5 - 7 \) in the \( x \) dimension (i.e., it overlaps with segments \( l_X[5] \) to \( l_X[7] \) in the \( x \) dimension), index range \( 7 - 7 \) in the \( y \) dimension (i.e., it overlaps with segment \( l_X[7] \) in the \( y \) dimension), and index range \( 4 - 5 \) in the \( z \) dimension (i.e., it overlaps with segment \( l_X[4] l_X[5] \) in the \( z \) dimension). Using this indirection, each spatial subbin is of fixed size. When compared to the purely temporal index, this spatiotemporal indexing scheme requires only additional space in GPU memory for the \( X, Y, \) and \( Z \) integer arrays, which corresponds to \( \geq 3|D| \times 4 \text{ bytes}. \)
4.3.2 Search Algorithm

On the host, as in the GPUTEMPORAL approach, we first sort $Q$ and for each query segment calculate the temporally overlapping entries from the temporal bins. We also compute the set of spatially overlapping subbins in each dimension. This computation also takes place on the host, where the description of the bins and subbins are stored. Arrays $X$, $Y$, and $Z$ are stored on the GPU. One option would be to compute the intersection of entry segments that belong to these subbins so as to select only spatially relevant entry segments. This turns out to be inefficient because we would then have to send a list of entry segment indices to the GPU, which has high overhead. Instead, we seek a solution in which we send a fixed and small number of indices to the GPU. As a result, we opt for a poorer but easier to encode selection of the candidate entry segments. Among the three spatial dimensions we pick the one in which the number of entry segments that overlap the query segment is the smallest. We then simply send an index range, 2 integers, in the $X$, $Y$, or $Z$ array, depending on the dimension that was picked. This approach may lead to wasteful computation on the GPU (i.e., evaluation of entry segments that do not overlap with the query segment in one of the other two spatial dimensions), but the overhead of these wasteful computations is offset by the gain from the reduced amount of data that is sent to the GPU. Let us demonstrate how this approach exploits the way in which the $X$, $Y$, and $Z$ arrays are constructed in the previous section. For the example in Figure 4 consider a query segment that overlaps temporal bins 0 and 1, and overlaps spatially with subbins $\hat{B}_{0,0}$ and $\hat{B}_{1,0}$ in the $x$ dimension (entries 1,3,4,5), with subbins $\hat{B}_{0,1}$ and $\hat{B}_{1,2}$ in the $y$ dimension (entries 3,5,7), and with subbins $\hat{B}_{0,0}$ and $\hat{B}_{1,0}$ in the $z$ dimension (entries 0,1,4,5). Because the smallest number of entries in the overlapping subbins is along the $y$ dimension, we opt to compare the query with entries 3,5, and 7. In array $Y$, these entries are stored contiguously at indices 7, 8, and 9. So we simply compare the query to the entry segments stored in array $Y$ from index 7 to index 9, which is encoded as one dimension specification and two integers, i.e., a constant size w.r.t. to the number of entry segments. We perform a comparison of the query segment with entry segment 7, even though entry 7 does not overlap the query along the $x$ and $z$ dimension. This comparison will thus lead to wasteful computation due to our non-perfect spatial selectivity of entry segments.

On the host, we generate a schedule $S$, which contains for each query segment $q_k$ a specification of which lookup array to use (0 for $X$, 1 for $Y$, or 2 for $Z$) and an index range into that array, which we encode using 4 integers (which preserves alignment). GPUSPATIOTEMPORAL requires only 1 extra indirection in comparison to GPUTEMPORAL, and avoids storing the overlapping entry indices in a buffer like in GPUSPATIAL. We then sort $S$ based on the lookup array specification so as to avoid thread serialization due to branching as much as possible. As for GPUTEMPORAL, Section 4.2 calculating $S$ on the host takes negligible time.

As explained in the previous section, we enforce a minimum size for the spatial subbins. Ensuring that subbins are not too small is necessary for two reasons. First, with small subbins each entry segment could overlap many subbins with high probability. As a result, the query id would occur many times in arrays $X,Y$, and/or $Z$, thereby wasting memory space on the GPU and causing redundant calculations. Second, given our indexing scheme and search algorithm described hereafter, a query that overlaps multiple subbins along all
three spatial dimensions may lead to duplicates in the result set. These duplicates would then need to be filtered out (either on the GPU or the CPU). To avoid duplicates, we simply default to the purely temporal scheme whenever duplicates would occur. While this behavior wastes computation (i.e., we lose spatial filtering capabilities), the constraint on subbin size described in the previous section ensure that it occurs with low probability.

The pseudo-code of the search algorithm is shown in Algorithm 3. It takes the following arguments: (i) the $X$, $Y$, and $Z$ arrays; (ii) the database ($D$); (iii) the query set ($Q$); (iv) the schedule ($S$); (v) the query distance ($d$); and (vi) the memory space to store the result set ($resultSet$). As in Algorithm 2, arguments that lead to array transfers between the host and the GPU are shown in boldface. The algorithm begins by checking the global thread id and aborts if it is greater than $|Q|$ (line 3). The query assigned to the thread is acquired from $Q$ (line 4). A helper array is constructed that holds pointers to the $X$, $Y$, and $Z$ arrays (line 5). If schedule $S$ does not give a specification for one of the $X$, $Y$, or $Z$ arrays ($S[gid].arrayXYZ = -1$) then the algorithm defaults to the temporal scheme (line 17). Otherwise, the algorithm retrieves the pointer to the correct $X$, $Y$, or $Z$ array (line 7) and determines the index range for the entry segments (lines 8-9). It then processes the entry segments (line 10) as Algorithm 2.

Algorithm 3 GPUSpatioTemporal kernel.

1: procedure SearchSpatiotemporal($X,Y,Z,D,Q,S,d$, resultSet)
2: gid ← getGlobalId()
3: if $gid \geq |Q|$ return
4: queryID ← gid
5: arraySelector ← \{X, Y, Z\}
6: if $S[gid].arrayXYZ \neq -1$ then
7: arrayXYZ ← arraySelector[$S[gid].arrayXYZ$]
8: entryMin ← $S[gid].entryMin$
9: entryMax ← $S[gid].entryMax$
10: for all $i \in \{\text{entryMin}, \ldots, \text{entryMax}\}$ do
11: entryID = arrayXYZ[$i$]
12: result ← compare($D[\text{entryID}],Q[\text{queryID}]$)
13: if result $\neq \emptyset$ then
14: atomic: resultSet $\leftarrow$ resultSet $\cup$ result
15: end if
16: end for
17: else
18: Lines 5-12 in Algorithm 2
19: end if
20: return
21: end procedure
5 Experimental Evaluation

5.1 Datasets

To evaluate the performance of our various indexing methods we use 3 datasets (1 real world and 2 synthetic) of 4-dimensional trajectories (3 spatial + 1 temporal). In previous work \cite{13} we have evaluated a purely temporal indexing scheme that shares the general principles of the scheme described in Section 4.2 (but assuming that $Q$ cannot fit in GPU memory). In that work we evaluated the performance of distance threshold search for datasets with varying statistical temporal properties, and found the index to perform equally well across these datasets. In this work, based on our previous experience and because we consider spatial and spatiotemporal indexing schemes, we use trajectory datasets that vary in terms of their sizes and spatial properties (e.g., spatial trajectory density):

- **Random-1M**: a small, sparse synthetic dataset;
- **Merger**: a large, real-world astronomy dataset;
- **Random-dense**: a high density synthetic dataset that is motivated by astronomy applications.

The **Random-1M** dataset consists of 2,500 trajectories generated via random walks over 400 timesteps, for a total of 997,500 entry segments. Trajectory start times are sampled from a uniform distribution over the $[0,100]$ interval.

The **Merger** dataset is from the field of astronomy and consists of particle trajectories that simulate the merger of the disks of two galaxies. It contains the positions of 131,072 particles over 193 timesteps for a total of 25,165,824 entry segments. Figure 5 depicts particles positions projected onto the $x-y$ plane at different times, showing the merger evolution.

The **Random-dense** dataset is generated as follows. Consider the stellar number density of the solar neighborhood, i.e., at galactocentric radius $R_\odot = 8$ kpc (kiloparsecs), of Reid et al. \cite{27}, $n_\odot = 0.112$ stars/pc$^3$. We develop a dataset with the same number of particles of one disk in the **Merger** dataset (65,536), and 193 timesteps. To match the density of \cite{27}, we require a volume of $65536/0.112 = 585142$ pc$^3$. This yields a cube with length, width and height dimensions of 83.64 pc. Note that we could have made the dataset more spatially dense by picking a region close to the galactic center, since the stellar density decreases as a function of $R$. We generate actual trajectories as random walks as in the **Random-1M** dataset, where all of the particles are initially populated within the aforementioned cube. We allow the trajectories to move a variable distance in each of the x,y,z dimensions at each timestep (between 0.001 and 0.005 kpc), and if a particle moves outside of the cube by 20% of the length of the cube in any dimension, the particle is forced back towards the cube. The particles, on average, cannot travel too far from the cube such that we maintain a fairly consistent trajectory density at each timestep. This dataset aims to represent a density consistent within the range of possible densities within the Milky Way that a single node might process. The characteristics of each dataset are summarized in Table 1.

\footnote{This dataset was obtained from Josh Barnes \cite{2}.}
**Table 1: Characteristics of Datasets**

| Dataset      | Trajec. | Entries       |
|--------------|---------|---------------|
| Random-1M    | 2,500   | 997,500       |
| Merger       | 131,072 | 25,165,824    |
| Random-dense | 65,536  | 12,582,912    |

Figure 5: Sample particle positions in the Merger dataset at times 0 Gyr (a), 1.5 Gyr (b) and 3 Gyr (c).

### 5.2 Experimental Methodology

For all our distance threshold search implementations the GPU-side is developed in OpenCL and the host-side is developed in C++. The host-side implementation is executed on one of the 6 cores of a dedicated 3.46 GHz Intel Xeon W3690 processor with 12 MiB L3 cache. The GPU-side implementation runs on an Nvidia Tesla C2075 card with 6GiB of RAM and 448 cores. In all experiments we measure query response time as an average over 3 trials (standard deviation over the trials is negligible). We allocate a buffer to hold the result set of the search on the GPU that can hold $5.0 \times 10^7$ items. In the description of the results we indicate when this buffer is overcome, thus requiring incremental processing of the query. The response time does not include the time to build the index or the time to store $D$ and the index in GPU memory. These operations can be performed off-line before query processing begins.

We consider three experimental scenarios, each for one of our datasets:

- **S1**: The Random-1M dataset and a query with 100 trajectories each with 400 timesteps for a total of 39,900 query segments.
- **S2**: The Merger dataset and a query set with 265 trajectories each with 193 timesteps for a total of 50,880 query segments.
- **S3**: The Random-dense dataset and a query set with 265 trajectories each with 193 timesteps for a total of 50,880 query segments.
For each scenario we use ranges of query distances (in units of kpc for S2 and S3).

In addition to our GPU implementations, we also evaluate a CPU-only implementation. This implementation relies on an in-memory R-tree index, and is multithreaded using OpenMP. Threads traverse the R-tree in parallel, each for a different query segment, and return candidate entry segments. This implementation was developed in our previous work [12, 13]. In that work we investigated “trajectory splitting,” i.e., the impact of the number of segments stored in each MBB in the R-tree index, \( r \). There is a trade-off between the time to search the index (which decreases as \( r \) increases due to lower tree depth) and the time to process the candidate (which increases as \( r \) increases due to higher index overlap). All executions of the CPU implementation use 6 threads on our 6-core CPU. Results in [11] show that this implementation achieves high parallel efficiency. Like for the GPU implementation, our response time measurements do not include the time to build the index tree.

Although the experimental results in the following sections are constrained by the specifics of our platform, the results of the CPU implementation are used to demonstrate that the GPU can be used efficiently for distance threshold searches. A fundamental difference between the CPU implementation and the GPU implementation is that the former relies on index-tree traversal while the latter relies on flat indexing schemes. This is because tree traversals on the GPU are problematic, e.g., due to thread divergence slowdowns.

### 5.3 Results for the Random-1M Dataset

In this section, we present results for the Random-1M dataset, first giving results for individual implementations and then combining results that make it possible to compare the implementations. The Random-1M dataset is representative of small and sparse datasets in which few or no entry segments are expected to lie within distance \( d \) of a query segment, i.e., with a low number of interactions.

Figure 6 shows response time vs. the number of entry segments per MBB \( (r) \) for the CPU implementation for a range of query distances. Using a single entry segment per MBB...
Figure 7: Response time vs. $d$ for GPUSpatial in scenario S1. Different curves are shown for different numbers of spatial cells in the $x$, $y$, and $z$ dimensions (i.e., “Cells=10” means a $10 \times 10 \times 10$ grid).

$(r = 1)$ does not lead to the best response time. For this experimental scenario using, e.g., $r = 10$ leads to good response times across all query distances.

Figure 7 plots response time vs. $d$ for GPUSpatial. Results are shown for a range of grid sizes. We use a total buffer size, $|U|$, of 2GiB to store overlapping entry segments. This is larger than the space necessary to store $D$. This is thus an optimistic configuration for the FSG index. Using too few grid cells leads, e.g., 10 per dimension, to poor performance due to poor spatial selectivity. With poor spatial selectivity (i) a large candidate set must be processed and (ii) many GPU threads overflow their entry buffers ($U_k$) thus requiring multiple query processing attempts. Likewise, using too many grid cells also leads to poor performance because entry segments overlap multiple cells. As a result there is duplication of index entries, and thus in the result set. Although filtering out these duplicates takes negligible time, transferring them from the GPU back to the host incurs non-negligible overhead. In these experiments, and among the FSG configurations we have attempted, using 50 cells per dimension leads to the best result.

Regardless of FSG configuration, we see rapid growth in response time as $d$ increases. The disposition of the FSG index to prefer small $d$ values has also been alluded to in [32]. This suggests that FSGs may not be particularly useful for spatiotemporal trajectory searches due to the large spatial extent of the data and absence of temporal discernment, unless query distances are small. However, a FSG index is likely to perform well with fewer requirements, such as indexing data with no temporal dimension, and/or focusing on point searches (instead of line segments), which will not cause data duplication when a large number of grid cells is used.

Figure 8 is shows response times vs. $d$ for GPUTemporal. Results are shown for a range of number of temporal bins. Unlike for GPUSpatial, this method is insensitive to the query distance. With too few temporal bins there is not enough temporal discrimination leading to large numbers of interactions. But as the number of bins increases the response time reaches a minimum (increasing beyond 5,000 bins does not differentiate entries as a function of temporal extent in the Random-1M dataset).
Figure 8: Response time vs. $d$ for GPUTEMPORAL in scenario S1. Different curves are shown for different numbers of temporal bins (100, 1000, 5000, 10000).

Figure 9: Response time vs. the number of subbins ($v$) for GPUSpatioTEMPORAL in scenario S1. The number of temporal bins is set to 10,000. Different curves are shown for different query distances ($d = 5, 10, ..., 50$).

Figure 9 shows response time vs. the number of subbins for GPUSpatioTEMPORAL, where 10,000 temporal bins are used. Curves are shown for a range of $d$ values. For low $d$ values a greater number of spatial subbins is desirable. This is because it is unlikely that a query will overlap multiple subbins, which would cause our algorithm to revert to the purely temporal method, which has no spatial selectivity. As $d$ increases, queries overlap multiple spatial subbins with higher probability. As a result, better performance is achieved with fewer subbins. Recall that we require that a query fall within a single subbin so as to avoid duplication in the result set. Without this requirement, an increasing number of subbins would suggest an increase in the duplication of entries in the index, thereby increasing the number of candidates that need to be processed (the same trade-off discussed for GPUSpatial). There is thus a trade-off between having too few or too many subbins, even when duplicates in the result set are permitted.

We note that using 1 subbin in the spatiotemporal index is equivalent to using a purely
temporal index with no spatial selectivity. Comparing results between GPUSpatialTemo-
ral with 1 subbin and GPUTemporal shows the effect of the additional indirection in the
spatiotemporal index. At $d = 50$ (yielding the greatest number of indirections in S1), with
1 subbin in the spatiotemporal index, the response time is 1.36 s, whereas the response time
is 1.21 s when using the temporal index without any indirection. This is a 12.4% increase in
response time due to the indirection.

Figure 10 shows response time vs. $d$ for our four implementations. Each implementa-
tion is configured with best or good parameter values based on previous results in this sections
(see the caption of Figure 10 for details). The CPU implementation is best across all query
distances. Comparing the GPU implementations, we see that GPUSpatial performs better
than GPUTemporal and GPUSpatialTemporal when $d < 20$, but that it does not scale
good for larger $d$ values. One may wonder whether this lack of scalability comes from the
overhead of re-launching the kernel due to buffer overflows. Figure 10 plots an “optimistic”
curve that discounts this overhead. We see that the same trend, if not as extreme, remains.
The temporal and spatiotemporal indexing methods have consistent response times across
query distances. Note that we could have selected the best number of subbins for each
value of $d$ from Figure 9 which would have improved results. Comparing GPUTemporal
and GPUSpatialTemporal, we see that having spatial selectivity in addition to temporal
indexing provides performance gains, even on this small and sparse dataset. We conclude
that an in-memory R-tree is a good approach when indexing small and sparse trajectory
datasets that lead to few interactions. For such a dataset, the overhead of using the GPU is
simply too great.
Figure 11: Response time vs. number of entry segments per MBB \((r)\) for the CPU implementation in scenario S2 with \(d = 0.01, 0.1, 0.05\).

### 5.4 Results for the Merger Dataset

In this section, we present results for our largest dataset, Merger, which contains over 25 million entry segments. From Section 5.3, we find that the purely spatial FSG method leads to extremely high response times for this larger dataset and as a result, we do not consider it. In GPU executions, for some values of \(d\), we have to process \(Q\) incrementally due to insufficient space for storing the full result set on the GPU. This is reflected in the measured response times.

Figure 11 shows response time vs. \(r\) for the CPU implementation for 3 query distances. With this large dataset, unlike with Random-1M, storing more than \(r = 1\) segments per MBB leads to higher response time. A higher \(r\) value decreases the time to search the R-tree index, but this benefit is offset by the increase in candidate set size. This is an important result. There is a literature devoted to assigning trajectory segments to MBBs for improving response time [16, 26, 12]. These works, however, do not consider large datasets. For these datasets, an intriguing future research direction is to take the opposite approach as that advocated in the literature: splice individual polylines to increase the size of the dataset (which can be thought of as setting \(r < 1\)).

We do not show results for GPUTEMPORAL as they are similar to those for the Random-1M dataset. Using 1,000 temporal bins leads to the lowest response time, which is consistent across all query distances.

Figure 12 shows response time vs. number of subbins for GPUSpatialTemporal, where 1,000 temporal bins are used. Curves are shown for a range of \(d\) values. A good number of subbins is \(v = 16\) across all query distances, and this value is in fact best for most query distances. This implies that picking a good \(v\) value can likely be done for a dataset regardless of the queries. Figure 9 shows a dependency between \(v\) and \(d\) for the Random-1M dataset. This dependency vanishes for a large dataset with many interactions.

Figure 13 compares the performance of the CPU implementation and GPUTEMPORAL and GPUSpatialTemporal (GPUSpatial is omitted). Each method is configured with best or good parameter values based on results in Figures 11 and 12. GPUSpatialTemporal outperforms GPUTEMPORAL across the board, with response times at least 23.6%
faster. At low query distances the CPU implementation yields the lowest response times. It is overtaken by GPUSpatioTemporal at $d \sim 1.5$. At $d = 0.001$ the response time for the CPU implementation is 9.70 s vs. 41.75 s for GPUSpatioTemporal (the GPU implementation is 330.4% slower). At $d = 5$ these response times become 184.4 s, and 116.09 s, respectively (the GPU implementation is 58.8% faster). We conclude that the GPU implementation outperforms the CPU implementation when using large datasets or when large query distances are considered.

5.5 Results for the Random-dense Dataset

We now present results for the Random-dense dataset, which is smaller than Merger and representative of scenarios in which many trajectories are located in a small spatial region, as motivated by the stellar number density at the solar neighborhood. Note that increasing the density by even $> 4 \times$ would still be consistent with that resembling the disk in the inner Galaxy.

Figure 14 shows response time vs. query distance for the CPU implementation for $r = 1, 2, 4, 8$. Unlike for Merger, which has $2 \times$ the number of entries as Random-dense, storing multiple segments/MBB improves response time. We find that $r = 4$ yields low response time values across all query distances.

As in the previous section, we do not show results for GPUTemporal as they are similar to those for the Random-1M dataset. Using 1,000 temporal bins leads to the lowest response time, which is consistent across all query distances.

Figure 15 (a) shows response time vs. the number of subbins ($v$) for scenario S3 for GPUSpatioTemporal. With this dataset, the use of subbins for reducing response time is only possible for small query distances ($d = 0.001, 0.01, 0.03$). This is because the dataset is smaller than Merger and because with larger values of $d$, the queries are more likely to fall within multiple subbins (in which case the search algorithm degenerates into a purely temporal scheme). Figure 15 (b) shows the fraction of queries that utilized the entries
Figure 13: Response time vs. $d$ for our implementations for scenario S2. For the CPU implementation we use $r = 1$ segments/MBB; for GPUTEMPORAL, we use 1,000 bins; for GPUSPATIOTEMPORAL, we use 1,000 temporal bins and $v = 16$ spatial subbins. We indicate three distance thresholds that would be interesting for the study of the habitability of the Milky Way based on such datasets. Red: close encounters between stars and planetary systems \cite{19}; Blue: supernova events on habitable planetary systems \cite{10}, and Magenta: studying the effects of gamma ray bursts on habitable planets \cite{29}. Both the Red and Blue lines are close to the vertical axis.

provided by the subbins for $d = 0.001, 0.01, 0.03$. Only the smallest query distance, $d = 0.001$, permits usage of the spatiotemporal index across a sizable fraction of the number of subbins. For instance for $d = 0.03$ and $v = 2$, just over 60% of the queries use the spatiotemporal index over the pure temporal index, and when $v = 4$, the entries provided by the spatiotemporal index are not used. This explains why in Figure 15 (a), there is no performance improvement for $d > 0.03$ when $v$ increases.

Given the density of the dataset, for larger values of $d$, only a fraction of the queries can be solved per kernel invocation as there is insufficient memory space for the result set. Since Random-dense has half as many entries as Merger, we can increase the size of the buffer on the GPU for the result set (from $5 \times 10^7$ elements for Merger to $9.2 \times 10^7$ elements for Random-dense). Figure 16 shows the response time vs. $d$ for GPUTEMPORAL and GPUSPATIOTEMPORAL with two buffer sizes. Increasing the buffer size by 84% (thus requiring fewer kernel invocations) leads to decreases in response time due to fewer host-GPU communications. For instance, at $d = 0.09$ (which requires the greatest number of kernel invocations), the spatiotemporal index, with $v = 2$, using an increased buffer size for the result set has a response time that is 65.76% lower than with the initial buffer size. Although we could not run experiments with a larger buffer size for scenario S2 (due to the large size of the Merger dataset), we expect similar performance gains. Since current trends point to improvements in host-to-GPU bandwidth, in the future, our indexing methods should provide even better performance improvements compared to CPU implementations.

Figure 17 shows response time vs. $d$ for the CPU implementation and GPUTEMPORAL and GPUSPATIOTEMPORAL with the larger buffer sizes. The query distance range spans a wide range of result set sizes. When $d = 0.001 \approx 0\%$ of the entries are within the query
distance, and when \( d = 0.09 \), 73.9% of the entries are within the query distance. For very small query distances \( d \lesssim 0.02 \), the CPU implementation yields the lowest response time, and is outperformed by the GPU implementations for larger \( d \). For \( d > 0.03 \), GPUSpatioTemporal performs slightly worse than GPUTemporal. This suggests that for dense datasets, when moderate to large query distances are required, the pure temporal indexing method performs the best. At \( d = 0.05 \), GPUTemporal is 223% faster than the CPU implementation (with \( r = 4 \)).

Comparing Figures 13 (Merger dataset) and 17 (Random-dense dataset), we see that the range of query distances for which the GPU method is preferable to the CPU method is much larger for the Random-dense dataset (considering the query distances that correspond to relevant application scenarios – the red, blue, and magenta vertical lines). In the astronomy application domain, datasets denser than the Random-dense dataset are relevant (i.e., to study the galactic regions at \( R < 8 \) kpc). For these datasets a GPU approach will provide even more performance improvement over a CPU implementation.

To summarize our results, Figure 18 shows the ratio of the response times of the GPU to CPU implementations for the 3 datasets for a few representative query distances. Data points below the \( y = 1 \) line correspond to instances in which the GPU implementation outperforms the CPU implementation. The main findings are that although the CPU is preferable for small and sparse datasets (Figure 18 (a)), the GPU leads to significant improvements for large and/or dense datasets (Figure 18 (b)) unless query distances are very small.

6 Conclusions

In this paper, we have proposed indexing methods and accompanying algorithms for efficient distance threshold similarity searches on spatiotemporal trajectory datasets. Our main result is that GPU-friendly indexing methods can outperform a multicore CPU implementation that uses an in-memory R-tree index. This is the case when the datasets are large and/or dense and the query distances are relatively large. Such scenarios are routine in some applications, and in particular in our driving application domain (astronomy). Overall, we
Figure 15: (a) Response time vs. number of subbins \(v\) for \texttt{GPUSpatioTemporal} for scenario S3 for a range of query distances. The number of temporal bins is set to 1,000. (b) The fraction of queries that use the entries provided by subbins vs. the number of subbins \(v\).

find that spatiotemporal indexing methods, which achieves selectivity both in time and space but without the use of an index tree, is effective on the GPU. The trends and future plans for GPU technology point to key improvements (faster host-to-GPU transfers, increased memory, etc.) that will give a further advantage to GPU implementation of spatiotemporal similarity searches.

Our results show that for the in-memory R-tree CPU implementation, the well-studied question of how to split a trajectory and store it in multiple MBBs is not pertinent for large datasets. For these datasets, storing a single segment by MBB is appropriate, and in fact it is likely appropriate to splice segments and increase dataset size so as to trade-off higher index-tree search time for small candidate sets to process. This result should apply to other similarity searches, such as \(k\)NN searches.

The main future work direction is to apply our indexing techniques to other spatial/spatiotemporal trajectory searches and investigating hybrid implementations of the distance threshold search that uses both the CPU and GPU for query processing.

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Figure 16: Response time vs. \( d \) for \text{GPUTemporal} and \text{GPUSpatioTemporal} for scenario S3. Results are shown for the original buffer size \((5 \times 10^7)\) and for a larger buffer size \((9.2 \times 10^7)\).

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Figure 17: Response time (left vertical axis) and fraction of entries with distance $d$ of the query (right vertical axis) vs. $d$ for the CPU implementation, GPUTEMPORAL, and GPUSPATIOTEMPORAL for scenario S3. For the CPU we show results for $r = 1$ and $r = 4$. 1,000 temporal bins are used for both the temporal and spatiotemporal indexing methods. $v = 2$ spatial subbins are used for the spatiotemporal indexing method.

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Figure 18: Ratio of GPU to CPU response times across all datasets for (a) S1 and (b) S2 and S3. Values below the $y = 1$ line indicate improvements over the CPU implementation.

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