

# Dynamic Graphs on the GPU

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**Abstract**—We present a fast dynamic graph data structure for the GPU. Our dynamic graph structure uses one hash table per vertex to store adjacency lists and achieves 3.4–14.8x faster insertion rates over the state of the art across a diverse set of large datasets, as well as deletion speedups up to 7.8x. The data structure supports queries and dynamic updates through both edge and vertex insertion and deletion. In addition, we define a comprehensive evaluation strategy based on operations, workloads, and applications that we believe better characterize and evaluate dynamic graph data structures.

**Index Terms**—dynamic, graph, data structures, GPU

## I. INTRODUCTION

While interest in graph analytics on GPUs has exploded in recent years, the vast majority of this work focuses on static graphs that never change during the graph computation. Today’s GPU graph analytic frameworks generally lack a GPU-managed dynamic graph data structure that supports changes (insertions and deletions) to the graph as well as queries into this data structure. The key challenge is representing the neighbors of each vertex (the “adjacency list”), where that data structure must be flexible enough to support a wide range of sizes and efficiently allow both queries into and changes to this data structure.

Previous efforts to support GPU-based dynamic graph data structures represent an adjacency list with a list-based data structure [1, 2] or an array-based data structure that maintains sort order [3]. Its implementation as a list/array presents a dilemma for its designer:

- Adjacencies can be stored as an unsorted list, which is easy to maintain. However, unsorted lists are unacceptably slow for important operations on the data structure, e.g., edge-existence queries (“does  $v$  have  $u$  as a neighbor”) or insertions that do not result in duplicates, both of which require traversing the entire list. Consequently, with an unsorted list, many operations are  $O(n)$  in the size of the list. This cost is prohibitive for vertices with many neighbors, as is common in scale-free graphs.
- To eliminate this  $O(n)$  cost, the adjacencies can instead be stored as a sorted list. These expensive operations become  $O(\log n)$  in the size of the list, but now the data structure must maintain the list in sorted order, which incurs a significant cost.

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From a performance standpoint, neither alternative is acceptable. Now, some graph operations can be implemented with high performance on an unsorted list, and thus for a subset of graph workloads, a list-based data structure may deliver acceptable performance. But many graph operations cannot be implemented without paying the high cost of a full search of an unordered list or the maintenance cost of preserving sorted order.

We believe that existing dynamic GPU graph data structures like faimGraph [2] and Hornet [1] are suboptimal when considering real-world dynamic graph scenarios. Truly dynamic data structures need to support continuous modifications not only from running the algorithm (e.g., edge deletion in k-truss), but also from a flowing stream of edge and vertex insertions and deletions. While a reasonable first step, the experiments presented in the faimGraph and Hornet works lack the true dynamism we expect in real world scenarios. Their chosen approaches both rely heavily on potentially expensive sorting operations necessary for vertex and edge deduplication. In general, duplicate entries lead to incorrect graph analytic results for the many graph primitives where idempotence does not apply (e.g., triangle counting or betweenness centrality).

We show that using a more sophisticated data structure (e.g., a hash table), we achieve better performance compared to list-based techniques provided by alternative data structures (e.g., faimGraph or Hornet). A major reason for our superior performance is the fast query rates that hash tables offer and their ability to ensure uniqueness while performing updates. In contrast, list-based data structures require explicit sorting for deduplication to maintain uniqueness. To evaluate our data structure, we integrate it into the Gunrock GPU graph analytics framework [4] and compare it against other dynamic graph data structures. The resulting overall performance of our data structure on a range of workloads and applications, particularly on insertions, is superior to existing alternatives and also allows reasonable tradeoffs dependent on the selected workload.

Our contributions in this work are:

- 1) A high-performance hash table based dynamic graph data structure that supports extremely high rates of insertions and deletions (sections III and IV);
- 2) An evaluation strategy that defines a set of workloads to benchmark a dynamic graph data structure (section V); and

- 3) Exploring the use of a dynamic graph data structure in applications while maintaining an updated graph (section VI).

## II. BACKGROUND AND PREVIOUS WORK

### A. Background

Consider a directed weighted graph  $\mathcal{G} = (\mathcal{V}, \mathcal{E}, \mathcal{W})^1$ , where  $\mathcal{V}$ ,  $\mathcal{E}$ , and  $\mathcal{W}$  represent the vertex set, edge set, and edge weight set respectively. For each arbitrary vertex  $u \in \mathcal{V}$ , we represent its outgoing neighbors (adjacency list) with  $\mathcal{A}_u$ .  $e = \langle u, v, w \rangle \in \mathcal{E}$  represents an edge from vertex  $u \in \mathcal{V}$  to  $v \in \mathcal{V}$  with weight  $w$ .

It is common to use an adjacency matrix with  $|\mathcal{V}|^2$  elements to represent dense graphs. Updating an adjacency matrix is trivial. However, if  $\mathcal{G}$  is a sparse graph (i.e.,  $|\mathcal{E}| \ll |\mathcal{V}|^2$ ), then the adjacency matrix is largely empty and, in practice, requires far too much memory for large graphs. Several alternative sparse graph representations exist. For example, *Compressed Sparse Row* (CSR), *Compressed Sparse Column* (CSC), and *Coordinate list* (COO) all offer better memory efficiency than the equivalent sparse adjacency matrix.

In CSR, for instance,  $\mathcal{A}_u$  is stored as an array of values (weights) and an array of destination vertex IDs. Then all adjacency lists (arrays of size equal to the out-degree of each vertex  $d_u$ , where  $\sum_{u \in \mathcal{V}} d_u = |\mathcal{E}|$ ) are concatenated to form two large arrays of size  $|\mathcal{E}|$ . A ternary array (i.e., row pointers) marks the start and end index of each adjacency list (i.e., each vertex's neighbors). CSR's representation requires  $O(|\mathcal{E}| + |\mathcal{V}|)$  elements, which is considerably more memory efficient than the  $O(\mathcal{V}^2)$  required for adjacency matrix. However, since it is a packed data structure, it is not possible to update it (i.e., delete or insert a new edge or vertex) without rebuilding the entire data structure.

In this work, we design a graph data structure that is both memory-efficient (like CSR) and also update-friendly (like the adjacency matrix). Suppose  $\mathcal{A}_u$  represents the adjacency list of a particular vertex  $u \in \mathcal{V}$ , where  $\mathcal{A}_u$  contains destinations of all outbound edges connected to  $u$ . Then a dynamic graph data structure should support the following operations:

- 1) Retrieving the adjacency list of vertex  $u$ : returns  $\mathcal{A}_u$  if it exists,  $\perp$  otherwise.
- 2) Inserting a new vertex  $u$ :  $\text{Insert}(u)$ , where  $\mathcal{A}_u$  is initialized with all connected vertices to  $u$ .
- 3) Deleting a vertex  $u$ :  $\text{Delete}(u)$ , where  $\mathcal{A}_u$  can no longer be located.  $\mathcal{A}_u$  is deleted. All edges that have  $u$  as their destination should be removed either immediately or in a lazy fashion. After a deletion, no edge query involving  $u$  may have a false positive result. Querying  $\mathcal{A}_u$  returns no edges.
- 4) Inserting a new edge:  $\text{Insert}(\langle u, v, w \rangle)$ , where  $\mathcal{A}_u$  is first located, then a new pair  $\langle v, w \rangle$  is inserted into  $\mathcal{A}_u$ .

<sup>1</sup>In general, we would like to support arbitrary meta-data assigned to each vertex or edge ( $\mathcal{D}_\mathcal{V}$  and  $\mathcal{D}_\mathcal{E}$ ). Here, for the sake of clarity, we assume  $\mathcal{W}$  represents any sort of meta-data associated with vertices or edges.

- 5) Deleting an edge:  $\text{Delete}(\langle u, v, w \rangle)$ , where  $\mathcal{A}_u$  is first located and then the pair  $\langle v, w \rangle$  is deleted.<sup>2</sup>

We assume that for each vertex  $u \in \mathcal{V}$ , the adjacency list  $\mathcal{A}_u$  is stored in a data structure that supports the following three operations:

- 1)  $\text{Search}_{\mathcal{A}_u}(v)$ : search through adjacency list  $\mathcal{A}_u$  and returns  $\langle v, w \rangle$  if present,  $\perp$  otherwise.
- 2)  $\text{Insert}_{\mathcal{A}_u}\langle v, w \rangle$ : inserts a new entry  $\langle v, w \rangle$  into the adjacency list  $\mathcal{A}_u$ .<sup>3</sup>
- 3)  $\text{Delete}_{\mathcal{A}_u}(v)$ : delete any entries  $\langle v, w \rangle$  from the adjacency list  $\mathcal{A}_u$ .

In general we assume that all of the operations are batched and performed in a *phase-concurrent* fashion (i.e., updating the data structure does not happen concurrently with any kind of read-only search query from the data structure).

### B. Previous Work

The major challenge for an efficient dynamic graph data structure on the GPU is the design of the adjacency list data structure to best accommodate potential updates. Memory management of this data structure is inherent in this challenge. Sha et al. proposed GPMA as a GPU-friendly data structure for dynamic graphs [3] based on the Packed Memory Array data structure (PMA) [5]. PMA is a kind of balanced binary search tree, where nodes are sorted arrays with some anticipated empty gaps to support potential updates. PMA uses lower and upper bound density thresholds for each node, and these thresholds are used to make decisions to either copy a node into a larger newly allocated node (high density), or properly merge multiple nodes into a single node (low density). In GPMA, a batch of updates is first sorted. The sorted batch is further partitioned into several continuous parts, where each part will only belong to a single node in the tree. Then each node is properly updated based on its partition's size in three granularities of warp/block/device. Sha et al. proposed a method to store a CSR format for a graph in a GPMA data structure. There is little discussion of memory management. Most of the experiments on updating the data structure are around edge insertions, but lazy edge deletions are also briefly discussed.

Hornet [1] divides the allocated available memory into blocks that can store a number of edges up to a specific power of two. Initially an adjacency list is stored inside the smallest power-of-two memory block that can contain it. During edge insertion, if the newly inserted edges exceed the capacity of a memory block, the vertex adjacency list is copied to the next smallest power-of-two memory block. For each array of blocks, a B-Tree tracks the free and used ones. Memory management is done on the CPU. Hornet achieves a compact representation for an adjacency list at the expense of

<sup>2</sup>One can define more general edge deletion operations such that all instances of  $\langle u, v, \cdot \rangle$  are deleted regardless of their weights. This version might be useful if we allow multiple edges from a source to a destination, each with a different weight or meta-data.

<sup>3</sup>Duplicates are not allowed: first search for  $v$  (i.e.,  $\text{search}_{\mathcal{A}_u}(v)$ ), and replace a previously inserted element if it exists. Otherwise, insert a new pair.

memory fragmentation. Moreover, it supports vertex insertion (or deletion) through a series of corresponding edge insertions (or deletions).

faimGraph [2] uses a single memory pool on the GPU for both the data structure and the algorithm that solves a graph problem. In contrast to Hornet, faimGraph's memory management is entirely on the GPU. Queues are used for memory reclamations of pages and deleted vertex IDs. faimGraph maintains a mapping between vertex IDs on the GPU and CPU. It also offers both structure-of-arrays (SoA) and array-of-structures (AoS) representations to store edge data, where the former is used for edges with a single property and the latter is used for edges with multiple properties. Memory pages configurable in size contain pointers to next pages when the adjacency list size exceeds a single page size. Using different GPU and CPU vertex IDs allows for flexible memory reclamation.

Edge duplication is not allowed in either Hornet or faimGraph. Both take preventive measures during updates to ensure edge uniqueness in the data structure.

We discuss and compare our results to faimGraph and Hornet. faimGraph is the state-of-the-art dynamic graph data structure and Hornet is a maintained graph processing library. In terms of similarities, our work is similar to faimGraph as our hash table is represented using fixed-size memory pages. In other words, if the hash table consists of a single bucket, which is true in road-network-like graphs (but not scale-free graphs), our work and faimGraph are similar. Similar to Hornet, addition of new vertices in our system requires overallocation of the graph data structure capacity to avoid reallocation, but we keep in mind that the cost of reallocation only requires copying of adjacency-list pointers and not the entire data structure.

### III. OUR GPU DYNAMIC GRAPH

The key challenge in designing a dynamic graph data structure is storing per-vertex adjacency lists. Our graph representation uses a separate data structure for each vertex adjacency list together with associated handles to reach those adjacency lists as necessary to perform various operations. The choice of the data structure used to store adjacency lists must be based on tradeoffs between what operations the data structure supports, the performance of individual operations, and the requirements of the graph library for solving specific problems. For instance, the simplest data structure choice is to use a variable-sized list data structure per adjacency list, with the designer either choosing to keep that list unsorted or maintain it as a sorted list. Hornet [1] embodies this approach. This representation is the most compact, but incurs a large maintenance overhead when compared to other options. Another possibility is the approach taken by faimGraph [2], which relaxes the variable-sized list constraint and instead breaks lists into fixed-size pages to simplify the maintenance of the data structure. Our position is that these primitive GPU data structures can be replaced by more sophisticated ones such as hash tables [6] or B-Trees [7], depending on the

requirements of the problem in terms of the performance and availability of data structure operations. In this work our goal is to provide a high throughput of both updates and lookups, thus we pick hash tables.

*Advantages of a hash table representation:* The primary advantage of hash tables is their efficient operations (both mutations and queries). Supporting efficient queries is an essential requirement in a graph data structure. Not only do graph applications perform read-only queries into graphs, but even mutation operations typically incorporate queries. For instance, an insertion while maintaining unique edges first requires a query determining whether the edge exists or not, followed by the insertion process itself (this is simply writing the new pair into an empty location). For a list-based data structure, performing a query operation is either  $O(n)$  (for an unsorted list) or  $O(\log n)$  (for a sorted list) in the size of the list. For a hash table with a suitable load factor, queries are instead  $O(1)$ . In a dynamic setting, hash table performance can decrease as the chain-length increases (i.e., load factor increases). In practice we can maintain low-cost metrics per vertex to determine the chain-length and periodically perform rehashing if it exceeds a given threshold.

Fig. 1 shows a high-level representation of our graph data structure, which is divided into two parts:

a) *Vertex dictionary:* We store vertices,  $\mathcal{V}$ , in a simple fixed-size array, indexed by vertex ID. The array size can be increased if needed, but frequent reallocation should be avoided to minimize the performance costs commonly associated with memory allocation. Selecting a large-enough initial capacity based on graph problem requirements ensures good performance during vertices insertion.

b) *Adjacency lists:* We use one hash table per vertex to store its associated adjacency list  $\mathcal{A}_u$ . Given a load factor and number of edges in an adjacency list, we calculate the number of buckets in a hash table. Note that the load factor, directly related to the number of buckets, provides a tradeoff between two main operations: 1) reading a complete adjacency list associated with a vertex and 2) performing an edge-exists query in a vertex's adjacency list. In practice we select a single load factor for all hash tables (in this work, we use a load factor of 0.7). This is not strictly necessary, but determining an ideal load factor per-vertex (per-hash-table) a priori is difficult. Our dynamic graph data structure can make use of given connectivity information along with the choice of the load factor to determine the number of necessary buckets to allocate. This decision results in significant performance gains by reducing memory allocation overhead. Using a dynamic memory allocator, any hash table can dynamically allocate additional slabs as needed (Fig. 1). If the connectivity information for a vertex is not available, we construct a hash table with a single bucket for this vertex.

### IV. IMPLEMENTATION

Our dynamic graph data structure's adjacency lists are stored as hash tables. In this work, we use *Slab Hash*, a dynamic hash table data structure for the GPU [6], as the basis

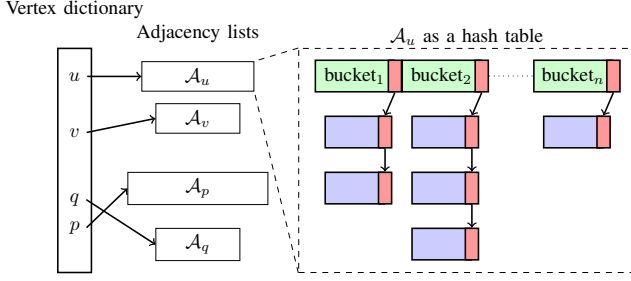


Fig. 1: High level schematic of our graph data structure. Each adjacency list is represented using a slab hash. The number of base slabs per adjacency list depends on the load factor used per adjacency list. Base slabs are statically allocated in consecutive memory locations, while the slabs used to resolve collisions are allocated dynamically and reached through pointers.

of our underlying hash tables.<sup>4</sup> We have significantly improved the functionality of the original slab hash in order to meet our requirements.<sup>5</sup> We offer two variants of our dynamic graph data structure. One uses Slab Hash’s *concurrent map*, which should be used if storing a value per edge is required. The second variant uses Slab Hash’s new *concurrent set*, which should be used if edge values are not required. Any hash table design can be used for this underlying data structure, as long as it is efficient in both searching (for queries) and updating the data structure itself (for insertions and deletions). In the end, the performance of our graph data structure directly depends on the performance of its underlying hash tables.

We integrate our dynamic graph data structure into the Gunrock GPU graph analytics framework [4]. In order to take advantage of the high-performance operations Slab Hash offers, all our operations are implemented based on the Warp Cooperative Work Sharing (WCWS) strategy [6]. In WCWS, each thread has an independent task assigned to it, but all threads within a warp cooperate with each other to collectively perform one independent task at a time. This is the right design decision because it better matches the memory access pattern desired by the GPU hardware (coalesced memory accesses), and hence it provides better performance for updates. On the downside, it requires all threads within a warp to be active. In other words, an operation on the data structure can not be performed within a branch where threads (in a warp) diverge when executing it.

#### A. Memory Management

Our memory management is divided into two parts: 1) vertex dictionary memory, and 2) hash table memory.

<sup>4</sup><https://github.com/owensgroup/SlabHash>

<sup>5</sup>The original slab hash only provided a *concurrent map* data structure, without any restrictions on duplicate keys. To name a few of our recent additions: maintaining key-uniqueness, proper iterator access, and design and implementation of a new concurrent set (keys only, and no values).

1) *Vertex dictionary memory*: Defining a graph requires defining the graph’s vertex capacity. The vertex dictionary stores pointers to the hash table associated with each vertex’s adjacency list. When inserting more vertices than the vertex dictionary’s capacity, we copy the vertex dictionary to a new memory location after increasing its capacity. This process only requires shallow copying of the pointers to each of the hash tables (including pointers to the hash tables associated with the new vertices).

2) *Adjacency list hash table memory management*: Constructing a hash table requires choosing and allocating a number of buckets (base slabs) that are required for insertion processes. The initial number of buckets for a vertex  $u$  is  $\lceil |\mathcal{A}_u| / (lf \times B_c) \rceil$ , where  $lf$  is the load factor and  $B_c$ , the bucket capacity per slab, is either 15 or 30 for Slab Hash map or set, respectively. During insertion, if a bucket’s slab becomes full (capacity achieved), Slab Hash dynamically allocates a new slab for that bucket that is singly linked to the tail of the list; a dynamic memory allocator handles these dynamic allocations [6]. Only when we perform vertex deletion (essentially this deletes an entire hash table) do we free this dynamically allocated memory (Section IV-D2).

Our graph data structure handles the memory allocation required for the initial buckets by statically allocating all the memory required for the initial buckets in bulk. This is more desirable than requiring each hash table to independently allocate a small number of buckets with different `cudaMalloc` calls. We initialize a vertex’s hash table with its initial number of buckets, the memory address for its first bucket, and the number of neighbors (to zero). In cases where the number of neighbors is not defined, we allocate a single bucket.

#### B. Query operations

To iterate over a vertex’s adjacency list, we provide a vertex adjacency list iterator. For a given vertex, the iterator loops over all of the hash table buckets associated with the vertex as well as additional slabs used to resolve hash collisions. The iterator loads one slab at a time and moves from one slab to the next using a `next` operator.

We also provide an `edgeExist` query that checks if the destination  $v$  of a given pair  $\langle u, v \rangle$  exists in the hash table associated with  $u$ . `edgeExist` simply performs a search query [6] in  $u$ ’s hash table.

#### C. Edge operations

We interpret edge operations (insertion and deletion) as modifications to the source vertex’s adjacency list. As discussed in Sec. III, we use a hash table to represent each vertex’s adjacency list. We discuss this in more depth below in the context of a directed graph. In an undirected graph, inserting (or deleting) an edge between a source and destination is similar but also requires an operation on the edge in the other direction. Our semantics for edge operations follows the semantics for hash table operations, which we discuss below.

1) *Edge insertion*: Algorithm 1 shows high-level pseudocode for an edge insertion. We assume that each thread has a single edge to insert. Initially, in Line 3, we ensure no self-edges are allowed. A work queue is constructed within a warp (Line 4) through a ballot instruction on all threads' remaining tasks. All threads locate the next task to perform (through finding the first set bit in the work queue as in Line 5) and then get the corresponding source vertex of the chosen task (through a shuffle instruction as in Line 6). Since multiple edges within a warp might share the same source vertex (Lines 7 and 8), all these insertions are grouped together to be performed in one single coalesced call to the hash table associated with the source vertex. We implemented and used a new slab-hash *replace* operation to ensure key-uniqueness in the hash table (i.e., unique destination vertices). In this operation, if a key (a destination vertex) already exists in the hash table, it will be replaced with the most recent value. Otherwise, a new key-value pair is added to the hash table. If the batch of edges contains the same unique edge but with different weights, only the most recent edge and its weight will be stored in the graph. The replace operation returns a boolean value indicating whether a new key (i.e., an edge) was added to the hash table or the key previously existed and was hence just replaced. We use this returned boolean variable to maintain an exact number of edges per vertex (population count on all successful additions within a warp in Line 10). The thread whose task was just completed, as well as all threads that shared the same source vertex (i.e., the coalesced insertion group), mark themselves as completed (Line 11). We repeat this procedure until the work queue is completely empty, i.e., no more edges remain to be inserted.

2) *Edge deletion*: Edge deletion is similar to edge insertion with two major differences: 1) instead of using the replace operation in Algorithm 1, we use the *delete* operation; 2) the delete operation also returns a boolean variable as to whether the key already existed. This boolean variable is used to decrement the number of edges that belongs to the adjacency list of the vertex. Note that in order to maintain uniqueness within the slab hash (due to how insertion/replace operations are designed and implemented), deleted edges (i.e., keys) are only marked as deleted (i.e., by replacing a key with a tombstone) and not explicitly removed. Tombstones are disregarded in edge insertion (as if that particular location is not empty). Tombstones can later be completely flushed out of the data structure, if required. Together, these design decisions ensure that empty locations can only exist at the end of each bucket's list in the slab hash. Moreover, not overwriting tombstones results in faster insertion rates (since new edges are only added to the end of the bucket's linked list). This comes at the expense of having unused memory locations. A different approach would be to break down the insertion process into two stages: 1) traversing the bucket's linked list to ensure uniqueness, then 2) for unique keys, a follow-up insertion that overwrites tombstones. We use the former approach during insertions, but the latter could be used to optimize for memory usage on the expense of decreased

insertion throughput.

#### D. Vertex Operations

1) *Vertex insertion*: We define a vertex insertion operation as the operation of inserting edges connected to a vertex that has an empty adjacency list. As discussed earlier, if the new vertex count exceeds the capacity of the vertex dictionary, we first extend the vertex dictionary. Once the vertex is entered into the dictionary, we then insert all attached edges using Algorithm 1.

2) *Vertex deletion*: Algorithm 2 summarizes this process for an undirected graph. Each warp deletes one vertex at a time. Because each vertex in a multi-vertex deletion operation may have a different number of edges, a straightforward implementation would suffer from load imbalance. We address this imbalance with a simple technique. We maintain a queue of deleted vertices with an atomic counter (Line 4). A single thread inside the warp acquires a new vertex from the queue (Line 3). The new vertex queue location is broadcast for all threads in the warp (Line 6). The vertex-deletion kernel only exits after deleting all the required vertices (Line 8). A warp reads the vertex index (Line 10) and requests an edge iterator over all the slabs associated with the vertex (Line 11). Using the iterator, we loop over all of the vertex destinations and delete the vertex from their adjacency lists (Line 16). Additionally, all dynamically allocated memory (i.e., memory used to resolve collisions) is freed and reclaimed by the memory allocator (Line 19). Finally, the count of edges connected to the vertex is set to zero (Line 22). Statically allocated memory is not reclaimed. For a directed graph, the only requirement is to free the memory. To clean up, we end with a follow-up lookup and delete all of the deleted vertices in all of the hash tables.

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#### Algorithm 1 Graph edge insertion algorithm.

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```

1: procedure INSERTEDGES(GpuGraph graph, Edges edges)
2:   thread_edge  $\leftarrow$  edges[threadIdx]
3:   to_insert  $\leftarrow$  thread_edge.src  $\neq$  thread_edge.dst
4:   while work_queue  $\leftarrow$  ballot(to_insert) do
5:     current_lane  $\leftarrow$  find_first_set_bit(work_queue)
6:     current_src  $\leftarrow$  shuffle(thread_edge.src, current_lane)
7:     same_src  $\leftarrow$  thread_edge.src == current_src
8:     success  $\leftarrow$  graph[current_src].replace(thread_edge, same_src & to_insert)
9:     added_count  $\leftarrow$  popc(ballot(success))
10:    graph[current_src].incrementEdgesCount(added_count)
11:    if same_src & to_insert then
12:      to_insert  $\leftarrow$  false
13:    end if
14:  end while
15: end procedure

```

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#### V. EVALUATION STRATEGY

Our community has not yet defined consistent standards for evaluating a dynamic graph data structure. In part this is because of the very recent development of these data structures as a topic for study. As a result, we lack a broad set of applications or workloads that require dynamic data structures. We believe the evaluation we present below improves on previous work by identifying and characterizing a set of operations, workloads, and applications that together encompass the wide

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**Algorithm 2** Graph vertex deletion algorithm.

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```
1: procedure DELETEVERTICES(GpuGraph graph, Vertices vertices, Count count,
   Queue queue)
2:   while true do
3:     if laneId == 0 then
4:       queueId ← atomicAdd(queue, 1)
5:     end if
6:     queueId ← shuffle(queueId, 0)
7:     if queueId ≥ count then
8:       return
9:     end if
10:    warp_vertex ← vertices[queueId]
11:    vertex_edges_it ← GpuGraph::EdgeIterator(warp_vertex)
12:    while vertex_edges_it.next() do
13:      lane_dst ← vertex_edges_it.getDst(laneId)
14:      for lane in lanes do
15:        current_dst ← shuffle(lane_dst, lane)
16:        graph[current_dst].delete(warp_vertex)
17:      end for
18:      if vertex_edges_it.current() is not base slab then
19:        free(vertex_edges_it.getAddress())
20:      end if
21:    end while
22:    graph[warp_vertex].setEdgesCount(0)
23:  end while
24: end procedure
```

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range of use cases that will be addressed with dynamic graph data structures.

We believe a comprehensive evaluation requires three components:

**Operations** Dynamic data structures support particular operations, e.g., edge and vertex deletion and insertion. We enumerate these operations and measure their throughput.

**Workloads** Because dynamic graph data structures on the GPU are not yet in significant use, applications that use them are few. However, we present a set of workloads—common patterns of how we will use the data structure—that we believe will underlie future applications in this area.

**Applications** Finally, prior work has identified particular applications on which they evaluate their data structure. We evaluate our work on these specific applications as well.

#### A. Low-Level Operations on a Dynamic Graph Data Structure

We begin with measuring throughput for the important low-level operations on our data structure:

1) *Edge Insertion and Deletion*: Starting from a static graph stored in a dynamic data structure, we measure the throughput of edge insertion and deletion operations for different batch sizes. Edges are inserted or deleted between existing vertices in the graph. Duplicate edges are allowed within a batch and across the batch and the graph. The graph data structure only maintains unique edges.

2) *Vertex Insertion and Deletion*: Similar to edge insertion and deletion, we start from a static graph and measure the throughput of inserting and deleting vertices in different batch sizes.

#### B. Workloads on a Dynamic Graph Data Structure

To evaluate a dynamic graph data structure we propose the following different set of workloads. Each one of these workloads targets a different scenario, not specific to any

particular application, that we expect to be a pattern that can be used in real-world applications.

1) *Static Workloads / Bulk-build*: We start by comparing our dynamic graph data structure to other alternatives in a static setting. Specifically, we evaluate the performance of building a static graph. We assume that the number of edges per vertex and the number of vertices is known a priori.

Given a static graph, we measure the time required for building the graph in bulk and compare this with previous work. We assume that the input is given in a COO format (i.e., a list of edges each defined by source vertex, destination vertex, and edge value).

2) *Dynamic Workloads / Incremental Build*: Starting with an empty graph, we incrementally build a graph using different batch sizes. In general, to avoid memory reallocation we assume that a suitable vertex capacity (i.e., maximum number of vertices) is known.

#### C. Applications with a Dynamic Graph Data Structure

We again emphasize that the existing set of graph applications that use dynamic data structures is small. We thus primarily evaluate work on the applications developed in and described by previous work where possible. For the *static application* case, we use static triangle counting to evaluate and compare our dynamic graph data structure performance to static CSR [4] and the two dynamic graph representations [1, 2]. In the *dynamic application* case, we evaluate based on a dynamic triangle counting application that performs triangle counting after each batch insertion. Note that in this work we only focus on the performance of the dynamic graph data structure. Optimizing a static or dynamic graph algorithm (e.g., triangle counting) is beyond the scope of this paper.

## VI. RESULTS

We evaluate and compare our dynamic graph data structure to Hornet<sup>6</sup> and faimGraph<sup>7</sup>. faimGraph is the state of the art in dynamic GPU graph data structures. Hornet is an actively maintained GPU data structure for sparse graphs and matrices. In our tests, faimGraph’s page size is configured to be 128 bytes to match our slab page size. Our tests do not require that either faimGraph or Hornet maintain a sorted adjacency list. All of our measured performance timings for all libraries only include the time to perform the operation and do not include the time required to transfer memory between CPU and GPU. We perform our benchmarking using the datasets shown in Table I on an NVIDIA TITAN V (Volta) GPU with 12 GB DRAM and an Intel Xeon CPU E5-2637.

#### A. Operations

1) *Batched Edge Insertion*: We perform a batched edge insertion (Sec. V-A1) for different batch sizes and measure the average throughput for all the given datasets. faimGraph only supports batch updates of sizes less than 1M. Table II

<sup>6</sup><https://github.com/hornet-gt/hornet/tree/a5c754d9616f54404a7b2a15c11143d52a346ab9>

<sup>7</sup><https://bitbucket.org/mwinter92/faimgaph>

TABLE I: Datasets

Dataset	Vertices	Edges	Degree			
			Min.	Max.	Avg.	$\sigma$
luxembourg_osm	114K	239K	1	6	2.1	0.41
germany_osm	11.5M	24.7M	1	13	2.1	0.51
road_usa	23.9M	57.71M	1	9	2.4	0.85
delaunay_n23	8.4M	50.3M	3	28	6.0	1.33
delaunay_n20	1M	6.3M	3	23	6.0	1.33
rgg_n_2_20_s0	1M	13.8M	0	36	13.1	3.62
rgg_n_2_24_s0	16.8M	265.1M	0	40	16.0	3.99
coAuthorsDBLP	299K	1.9M	1	336	6.4	9.80
ldoor	952K	45.5M	27	76	47.7	11.97
soc-LiveJournal1	4.8M	85.7M	0	20K	17.2	50.65
soc-orkut	3M	212.7M	1	27K	70.9	139.72
hollywood-2009	1.1M	112.8M	0	11K	98.9	271.70

TABLE II: Mean edge insertion rates (in MEdge/s) for different batch sizes.

Batch size	Hornet	faimGraph	Ours
$2^{16}$	33.67	92.47	501.33
$2^{17}$	44.71	133.97	513.56
$2^{18}$	51.43	157.15	591.06
$2^{19}$	70.81	188.98	641.25
$2^{20}$	83.54	—	664.52
$2^{21}$	97.41	—	658.09
$2^{22}$	110.89	—	646.01

compares our edge insertion throughput to Hornet and faimGraph. Our speedup ranges between 5.8–14.8x compared to Hornet and 3.4–5.4x compared to faimGraph.

2) *Batched Edge Deletion*: Similar to batched edge insertion, we run a batched edge deletion. Table III shows the result of this benchmark. This is where Hornet performance becomes competitive with ours. Deletion is a simple process and does not require cross-duplicate checking between the graph and the input batch. Note that for small datasets, the true number of deleted edges (i.e., unique edges within the batch) is much lower than the number of randomly generated edges, hence resulting in less work in general. Our performance is as fast as Hornet for a large batch size and almost 7x faster for a smaller batch size of  $2^{16}$ . Our deletion rates are between 3.6–7.8x faster than faimGraph’s.

TABLE III: Mean edge deletion rates (in MEdge/s) for different batch sizes.

Batch size	Hornet	faimGraph	Ours
$2^{16}$	91.73	111.71	640.63
$2^{17}$	159.69	112.96	886.92
$2^{18}$	259.31	171.75	947.60
$2^{19}$	377.79	257.66	939.98
$2^{20}$	537.73	—	988.85
$2^{21}$	739.82	—	1,007.16
$2^{22}$	1,024.87	—	1,015.47

TABLE IV: Mean vertex deletion throughput (in MVertex/s) for different batch sizes.

Batch size	faimGraph	Ours
$2^{16}$	0.44	5.35
$2^{17}$	0.71	9.23
$2^{18}$	1.12	12.66
$2^{19}$	1.62	19.21
$2^{20}$	2.96	26.49

3) *Vertex Deletion*: Beginning with an undirected graph, we delete a batch of vertices and measure the throughput of vertex deletion. Table IV shows the throughput for different batch sizes averaged over four datasets: soc-orkut, soc-LiveJournal1, delaunay\_n23, and germany\_osm. Our vertex deletion throughput is between 8.9–12.2x faster than faimGraph (Hornet does not implement vertex deletion). Both we and faimGraph delete vertices from neighbor adjacency lists and free the memory used to store the vertex adjacency list, but faimGraph implements one operation that we do not: it places the deleted vertex into a vertex queue and can thus reuse identifiers of deleted vertices during subsequent vertex insertions. This allows faimGraph to be more memory efficient compared to our approach. It would be straightforward to implement the same strategy with our data structure but we have not yet done so. If we compare only delete and free operations common to both data structures, our speedup is 8.5–11.56x over faimGraph. As with query operations, the dominant factor in vertex-deletion performance is looking up a deleted vertex in its neighbors’ adjacency lists; this is faster in a hash table than in a list.

## B. Workloads

1) *Bulk Build*: We perform the bulk build benchmark from Sec. V-B1. Bulk build is simply inserting all edges from a graph into the graph data structure in one single batch. We implemented the bulk build functionality in Hornet only. Table V shows the time required to bulk build the datasets. For two datasets—rgg\_n\_2\_24\_s0 and soc-orkut—Hornet runs out of memory. We believe that this is due to the memory overhead of sorting and duplicate checking. Our dynamic graph data structure is 2–30x faster. Note that for a large dataset, hollywood-2009, 45% of Hornet’s insertion time is spent in duplication checking alone, which is the same time as our entire build.

2) *Incremental Build*: In this benchmark we begin with an empty graph and incrementally insert edges (Sec. V-B2). The goal is to test and measure the edge throughput when building a graph data structure given a known bound on the number of vertices a priori, but an unknown number of edges. For our dynamic graph data structure, this means that each hash table is given only one bucket (i.e., a single linked list). Note that in this experiment our data structure is similar to faimGraph, but differs from Hornet in our use of a linked list of pages versus a single block containing the adjacency list. For our

TABLE V: Bulk build elapsed time (ms).

Dataset	Hornet	Ours
luxembourg_osm	5.562	0.184
germany_osm	330.311	12.407
road_usa	644.308	27.910
delaunay_n23	273.532	19.590
delaunay_n20	37.68	2.494
rgg_n_2_20_s0	37.084	5.053
rgg_n_2_24_s0	—	0.697
coAuthorsDBLP	11.672	0.835
ldoor	46.486	15.936
soc-LiveJournal1	179.879	26.176
soc-orkut	—	39.907
hollywood-2009	90.705	42.387

TABLE VI: Incremental build mean edge insertion rates (in MEdge/s) for different batch sizes.

Batch size	Hornet	Ours
2 <sup>20</sup>	164.44	841.31
2 <sup>21</sup>	176.96	945.64
2 <sup>22</sup>	184.75	993.82

hash table based graph data structure, this represents the worst-case scenario.

Table VI shows the average throughput for building graphs with a similar number of edges (ldoor, delaunay\_n23, road\_usa, soc-LiveJournal1) using different batch sizes. We implemented incremental build in Hornet only. On average our data structure is 5x faster than Hornet. For the two low-variance graph datasets (delaunay\_n23 and road\_usa), our speedups are between 15–25x. We believe that the main reason for our performance advantage is that Hornet maintains its adjacency list in a single fixed-size block. When an added edge exceeds the size of the block, the entire adjacency list must be copied to an existing or newly allocated empty block of the appropriate size. In contrast, our linked lists avoid copying and simply allocate new pages to accommodate new edges as needed. With Hornet, copying into larger-sized blocks is expected to happen more often for low-variance datasets. For high-variance datasets, Hornet’s doubling adjacency list strategy becomes more efficient because the need for copying to new blocks decreases. We see speedups of 1.6–2.5x for the ldoor dataset, but for the soc-LiveJournal1 dataset our throughput is 0.92x slower.

### C. Applications

We pick triangle counting as a simple application to explore the interaction between a dynamic data structure and solving a graph problem. Our goal here is not to provide an optimal solution to the dynamic triangle counting problem; rather, we would like to explore the performance of triangle counting’s main query operation, intersect. The intersect operation inputs two adjacency lists and counts the number of edges in common. If the adjacency list is stored as a list, to perform an intersect operation efficiently, the list must be sorted. The

TABLE VII: Static triangle counting time in ms.

Dataset	Hornet	faimGraph	Ours
luxembourg_osm	0.57	1.01	0.31
germany_osm	26.31	16.61	29.69
road_usa	51.57	39.19	66.20
delaunay_n23	25.74	21.14	56.38
delaunay_n20	3.35	2.92	6.43
rgg_n_2_20_s0	6.42	7.35	23.24
rgg_n_2_24_s0	154.75	165.86	493.6
coAuthorsDBLP	1.20	4.75	6.98
ldoor	22.09	48.04	222.07
soc-LiveJournal1	482.98	705.71	1526
soc-orkut	3832	8986	6758
hollywood-2009	9784	57311	11060

hash-based data structure we present here does not have this constraint. We thus compare the cost of maintaining the sorted list-based data structures used by Hornet and faimGraph with our approach.

1) *Static*: In this experiment, we compare dynamic graph data structures to solve the static graph problem of triangle counting. Since triangle counting only requires maintaining the destinations of edges and not their values, we use the set variant of the dynamic graph data structure. The Hornet and faimGraph data structures require sorted adjacency lists to efficiently compute set intersections. Table VII shows the time required to perform triangle counting on different datasets. On most datasets, our dynamic data structure performs worse than either Hornet or faimGraph, because their intersection operation between two sorted lists is efficient. They find the starting location of one list in the other and then (serially) walk to the end of the lists, accumulating the number of matches. While this exhibits little parallelism, it is cheaper and faster than a hash-table-based solution. In our hash table representation, we perform an `edgeExist` query for all edges.

Note, the sort in the list-based data structures is not free, and is not counted in the results above. Table VIII summarizes sort cost on these datasets. Hornet does not provide a GPU sort for their data structure, so we substitute CUB’s segmented sort by key [8]. Interestingly, faimGraph’s sorting is faster than CUB’s when the maximum vertex degree of the graph is small, but for a large maximum vertex degree, faimGraph’s sort is much slower than CUB’s. These results raise the question of the overhead of maintaining a sorted Hornet or faimGraph data structure in order to perform a dynamic application that requires a sorted list, such as triangle counting. We further investigate this in the following section.

2) *Dynamic*: For this experiment we pick two datasets, one with a small largest-vertex degree (road\_usa) and one with a large largest-vertex degree (hollywood-2009). We perform triangle counting after incrementally inserting edges five times. This scenario was not previously implemented in either Hornet or faimGraph; we implemented it for Hornet only. Table IX shows the result of this experiment. For road\_usa, our implementation offers a 1.8x speedup over Hornet’s, largely



TABLE VIII: CSR-sort (with CUB) and faimGraph-sort time in ms. Sort time is in general comparable to (and often considerably larger than) the time for triangle counting (Table VII).

Dataset	Sort CSR	Sort faimGraph
luxembourg_osm	58.13	0.07
germany_osm	5260	4.84
road_usa	10875	12.65
delaunay_n23	3854	18.98
delaunay_n20	503.29	2.48
rgg_n_2_20_s0	496.85	8.62
rgg_n_2_24_s0	7753	178.37
coAuthorsDBLP	136.89	7.36
ldoor	442.15	175.12
soc-LiveJournal1	2226	20428
soc-orkut	1404	41833
hollywood-2009	540.30	8504

TABLE IX: Cumulative time required to perform triangle counting and inserting a batch of size  $2^{22}$  into the graph.

Iter.	Insert	Ours TC	Total	Insert	Hornet TC	Total	Speedup
road_usa							
1		65.5	64.1		51.6	116.0	1.81
2	14.8	135.8	129.5	214.1	110.2	235.1	1.82
3	29.7	201.7	195.0	438.4	174.6	356.4	1.83
4	44.5	267.5	260.5	652.6	243.8	476.2	1.83
5	59.4	333.2	325.8	855.7	319.7	597.8	1.83
hollywood-2009							
1		11151	11151		9893	9893	0.89
2	12.4	22539	22551	73.4	19982	20056	0.89
3	24.9	33921	33946	149.0	30174	30323	0.89
4	37.3	45297	45335	229.3	40552	40781	0.90
5	49.8	56724	56774	313.0	51090	51403	0.91

due to our faster insertion. For hollywood-2009, although our insertion performance is around 6x faster than Hornet, Hornet's faster triangle counting is still fast enough to cover the cost of maintaining sorted adjacency lists. We are 0.9x slower than Hornet on this dataset.

#### D. Effect of the load factor on our graph data structure

To measure the effect of load factor on our hash table, we perform two experiments. Fig. 2 shows our first experiment where we manipulate the chain length of our hash tables by building a graph with the suitable average degree and load factor. As we expect, the insertion throughput of our data structure drops as the chain length increases. On the other hand, the memory utilization increases. This is due to the fact that buckets are now more full. Moreover, the amount of memory used decreases as the average chain length increases. This is due to the fact that fewer buckets are now needed. Fig. 3 shows our second experiment. Similar to the first one, we explore the query performance as the average chain length increases (we use static triangle counting to provide the query workload). The results show the optimal average chain length for our hash table, which is around 0.7.

## VII. CONCLUSION AND FUTURE WORK

Our dynamic GPU graph data structure uses hash tables to represent per-vertex adjacency lists. This representation suits operations that require fast insertion, deletion, and edge lookups, and is superior in performance to previous work that focuses on list data structures for adjacency lists.

In regards to future work, we note that other data structures can be used to represent adjacency lists. For instance, a B-Tree [7] provides a different set of operations as well as maintaining a sorted adjacency list, an optimization that is useful in certain graph algorithms. We also note that supporting hash tables with varying slab sizes may better suit load balancing in scale-free graphs where vertex degrees vary over several orders of magnitude. This would complicate the update procedure at the expense of providing better scalability in graph processing. Moreover, it would require a more complicated dynamic memory allocator design (compared to SlabAlloc used in slab hash [6]) that can support variable-sized memory allocations efficiently.

Although we only discussed phase-concurrent updates and queries in this work, both the slab hash and the B-Tree provide concurrent queries and updates. These concurrent operations can be exposed to a dynamic graph algorithm. The key challenge here is to carefully consider the semantics of these operations. For instance, a graph problem might need to lock a graph node to ensure a consistent view of the adjacency list for the different query and update operations.

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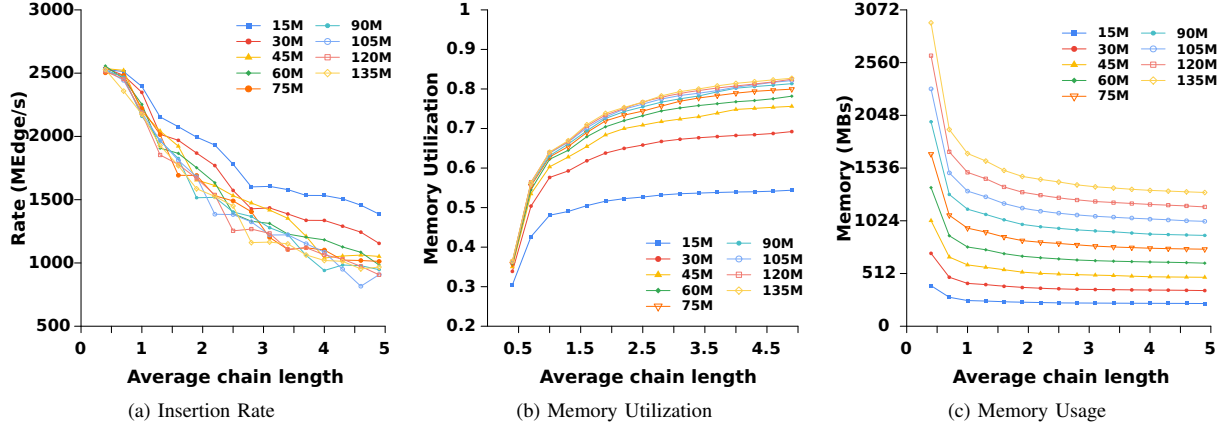


Fig. 2: For different directed RMAT graphs with  $2^{20}$  vertices but different average degree (different number of edges), we build the graph using different load factors. (a) The insertion throughput drops by a factor of 2.5 if the hash tables have, on average, chains of length 5. On the other hand, the memory utilization increases (b) and the memory usage decreases (c) as the average chain-length increases.

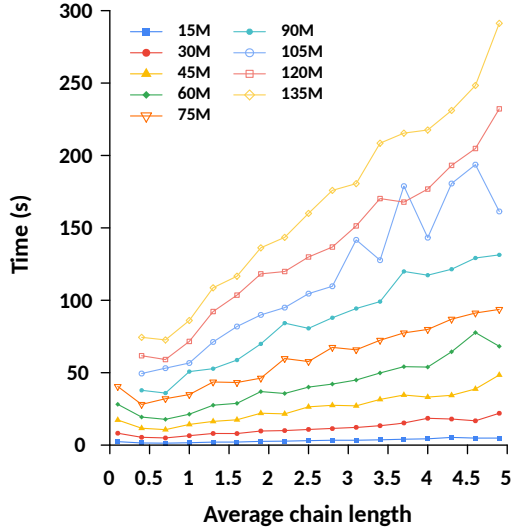


Fig. 3: Static triangle counting performance for different undirected RMAT graphs with  $2^{20}$  vertices, but different average degree using hash tables with different load factors. Our data structure achieves its optimal performance when the load factor is around 0.7.

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