Low-Latency Graph Streaming using Compressed Purely-Functional Trees*

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Abstract

There has been a growing interest in the graph-streaming setting where a continuous stream of graph updates is mixed with graph queries. In principle, purely-functional trees are an ideal fit for this setting as they enable safe parallelism, lightweight snapshots, and strict serializability for queries. However, directly using them for graph processing leads to significant space overhead and poor cache locality.

This paper presents C-trees, a compressed purely-functional search tree data structure that significantly improves on the space usage and locality of purely-functional trees. We design theoretically-efficient and practical algorithms for performing batch updates to C-trees, and also show that we can store massive dynamic real-world graphs using only a few bytes per edge, thereby achieving space usage close to that of the best static graph processing frameworks.

To study the applicability of our data structure, we designed Aspen, a graph-streaming framework that extends the interface of Ligra with operations for updating graphs. We show that Aspen is faster than two state-of-the-art graph-streaming systems, Stinger and LLAMA, while requiring less memory, and is competitive in performance with the state-of-the-art static graph frameworks, Galois, GAP, and Ligra+. With Aspen, we are able to efficiently process the largest publicly-available graph with over two hundred billion edges in the graph-streaming setting using a single commodity multicore server with 1TB of memory.

CCS Concepts • Computing methodologies → Vector / streaming algorithms.

Keywords • streaming graph processing, purely-functional data structures, parallel graph algorithms

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1 Introduction

In recent years, there has been growing interest in programming frameworks for processing streaming graphs due to the fact that many real-world graphs change in real-time (e.g., [21, 29, 30, 34, 47, 83]). These graph-streaming systems receive a stream of queries and a stream of updates and must process both updates and queries with low latency, both in terms of query processing time and the time it takes for updates to be reflected in new queries. There are several existing graph-streaming frameworks, such as STINGER, based on maintaining a single mutable copy of the graph in memory [29, 30, 34]. Unfortunately, these frameworks require either blocking queries or updates so that they are not concurrent, or giving up serializability [83]. Another approach is to use snapshots [21, 47]. Existing snapshot-based systems, however, are either very space-inefficient, or suffer from high latency on updates. Therefore, an important question is whether we can design a data structure that supports lightweight snapshots which can be used to concurrently process queries and updates, while ensuring that the data structure is safe for parallelism and achieves good asymptotic and empirical performance.

In principle, representing graphs using purely-functional balanced search trees [1, 57] can satisfy both criteria. Such a representation can use a search tree over the vertices (the vertex-tree), and for each vertex store a search tree of its incident edges (an edge-tree). Because the trees are purely-functional, acquiring an immutable snapshot is as simple as acquiring a pointer to the root of the vertex-tree. Updates can then happen concurrently without affecting the snapshot. In fact, any number of readers (queries) can concurrently acquire independent snapshots without being affected by a writer. A writer can make an individual or bulk update and then set the root to make the changes immediately and atomically visible to the next reader without affecting current active readers. A single update costs $O(\log n)$ work, and because the trees are purely-functional it is relatively easy and safe to parallelize a bulk update.
However, there are several challenges that arise when comparing purely-functional trees to compressed sparse row (CSR), the standard data structure for representing static graphs in shared-memory graph processing [64]. In CSR, the graph is stored as an array of vertices and an array of edges, where each vertex points to the start of its edges in the edge-array. Therefore, in the CSR format, accessing all edges incident to a vertex \( v \) takes \( O(\text{deg}(v)) \) work, instead of \( O(\log n + \text{deg}(v)) \) work for a graph represented using trees. Furthermore, the format requires only one pointer (or index) per vertex and edge, instead of a whole tree node. Additionally, as edges are stored contiguously, CSR has good cache locality when accessing the edges incident to a vertex, while tree nodes could be spread across memory. Finally, each set of edges can be compressed internally using graph compression techniques [71], allowing massive graphs to be stored using just a few bytes per edge [25]. This approach cannot be used directly on trees. This would all seem to put a search tree representation at a severe disadvantage.

In this paper, we describe a compressed purely-functional tree data structure that we call a C-tree, which addresses the poor space usage and locality of purely-functional trees. The C-tree data structure allows us to take advantage of graph compression techniques, and thereby store very large graphs on a single machine. The key idea of a C-tree is to chunk the elements represented by the tree and store each chunk contiguously in an array. Because elements in a chunk are stored contiguously, the structure achieves good locality. By ensuring that each chunk is large enough, we significantly reduce the space used for tree nodes. Although the idea of chunking is intuitive, designing a chunking scheme which admits asymptotically-efficient algorithms for batch-updates and also performs well in practice is challenging. We note that our chunking scheme is independent of the underlying balancing scheme used, and works for any type of element.

In the context of graphs, because each chunk in a C-tree stores a sorted set of integers, we can compress by applying difference coding within each block and integer code the differences. We compare to some other chunking schemes, including B-trees [4] and ropes [2, 9, 15, 31] in Section 3.3.

To address the asymptotic complexity issue, we observe that for many graph algorithms the \( O(\log n) \) work overhead to access vertices can be handled in one of two ways. The first is for global graph algorithms, which process all vertices and edges. In this case, we can afford to compute a flat snapshot, which is an array of pointers to the edge-tree for each vertex. We show how to create a flat snapshot using \( O(n) \) work, \( O(\log n) \) depth, and \( O(n) \) space. A flat snapshot can be created concurrently with updates and other reads since it copies from the persistent functional representation. Once a flat snapshot is created, the work for accessing the edges for a vertex \( v \) is only \( O(\text{deg}(v)) \), as with CSR. The second case is for local graph algorithms, where we cannot afford to create a flat snapshot. In this setting, we note that many local algorithms examine all edges incident to a vertex after retrieving it. Furthermore, although real-world graphs are sparse, their average degree is often in the same range or larger than \( \log n \). Therefore, the cost of accessing a vertex in the vertex-tree can be amortized against the cost of processing its edges.

To evaluate our ideas, we describe a new graph-streaming framework called Aspen that enables concurrent, low-latency processing of queries and updates on graphs with billions of vertices and hundreds of billions of edges, all on a relatively modest shared-memory machine equipped with 1TB of RAM. Our system is fully serializable and achieves high throughput and performance comparable to state-of-the-art static graph processing systems. Aspen extends the interface proposed by Ligra [70] with operations for updating the graph. As a result, all of the algorithms implemented using Ligra, including graph traversal algorithms, local graph algorithms [72], algorithms using bucketing [24], and others [25], can be run using Aspen with minor modifications. We have made Aspen publicly-available at https://github.com/ldhulipala/aspen.

Compared to state-of-the-art graph-streaming frameworks, Aspen provides significant improvements both in memory usage (8.5–11.4x more memory-efficient than Stinger [29] and 1.9–3.3x more memory-efficient than LLAMA [47]), and algorithm performance (1.8–10.2x faster than Stinger and 2.8–15.1x faster than LLAMA). Aspen is also comparable to the fastest static graph processing frameworks, including GAP [6] (Aspen is 1.4x faster on average), Ligra+ [71] (Aspen is 1.4x slower on average), and Galois [56] (Aspen is 12x faster on average). Compared to Ligra+, which is one of the fastest static compressed graph representations, Aspen only requires between 1.8–2.3x more space.

Our experiments show that adding a continuous stream of edges while running queries does not affect query performance by more than 3%. Furthermore, the latency is well under a millisecond, and the update throughput ranges from 11K–78K updates per second when performing one update at a time to 105M–442M updates per second when performing batches of updates. We show that our update rates are an order of magnitude faster than the update rates achievable by Stinger, even when using very small batches.

The contributions of this paper are as follows:

1. A practical compressed purely-functional data structure for search trees, called the C-tree, with operations that have strong theoretical bounds on work and depth.
2. The approach of flat-snapshotting for C-trees to reduce the cost of random access to the vertices of a graph.
3. Aspen, a multicore graph-streaming framework built using C-trees that enables concurrent, low-latency processing of queries and updates, along with several algorithms using the framework.
4. An experimental evaluation of Aspen in a variety of regimes over graph datasets at different scales, including the largest publicly-available graphs (graphs with billions
of vertices and hundreds of billions of edges), showing significant improvements over state-of-the-art graph-streaming frameworks, and modest overhead over static graph processing frameworks.

2 Preliminaries

Notation and Primitives. We denote a graph by $G(V, E)$, where $V$ is the set of vertices and $E$ is the set of edges in the graph. For weighted graphs, the edges store real-valued weights. The number of vertices in a graph is $n = |V|$, and the number of edges is $m = |E|$. Vertices are assumed to be indexed from 0 to $n − 1$. For undirected graphs, we use $N(v)$ to denote the neighbors of vertex $v$ and $\text{deg}(v)$ to denote its degree. We assume that we have access to a family of uniformly (purely) random hash functions which we can draw from in $O(1)$ work [22, 58]. In functions from such a family, keys are mapped to elements in the range with equal probability, independent of the values that other keys hash to, and the function can be evaluated for a key in $O(1)$ work.

Work-Depth Model. We analyze algorithms in the work-depth model, where the work is the number of operations used by the algorithm and the depth is the length of the longest sequential dependence in the computation [14, 39].

Purely-Functional Trees. Purely-functional (mutation-free) data structures preserve previous versions of themselves when modified and yield a new structure reflecting the update [57]. The trees studied in this paper are binary search trees, which represent a set of ordered elements. In a purely-functional tree, each element is used as a key, and is stored in a separate tree node. The elements can be optionally associated with a value, which is stored in the node along with the key. Trees can also be augmented with an associative function $f$ (e.g., $+$), allowing the sum with respect to $f$ in a range of the tree to be queried in $O(\log n)$ work and depth, where $n$ is the number of elements in the tree.

Interfaces for Graphs. We will extend the interface defined by Ligra [70] and so we review its interface here. We use the vertexSubset data structure which represents subsets of vertices, and the edgemap primitive which is used for mapping over edges incident to sets of vertices. edgemap takes as input a subset of vertices and applies a function over the edges incident to the subset that satisfy a condition (e.g., edges to vertices that have not yet been visited by a breadth-first search). Specifically, edgemap takes as input a graph $G(V, E)$, a vertexSubset $U$, and two boolean functions $F$ and $C$; it applies $F$ to $(u, v) \in E$ such that $u \in U$ and $C(v) = \text{true}$ (call this subset of edges $E_a$), and returns a vertexSubset $U'$ where $u \in U'$ if and only if $(u, v) \in E_a$ and $F(u, v) = \text{true}$.

3 Compressed Purely-Functional Trees

In this section, we describe a compressed purely-functional search tree data structure called a C-tree. After describing the data structure (Section 3.1), we argue that C-trees improve locality and reduce space-usage relative to ordinary purely-functional trees (Section 3.2). Finally, we compare C-trees to other possible design choices, such as B-trees (Section 3.3).

3.1 C-tree Definition

The main idea of C-trees is to apply a chunking scheme over the tree to store multiple elements per tree-node. The chunking scheme takes the ordered set of elements to be represented and "promotes" certain elements to be heads, which are stored in a tree. The remaining elements are stored in tails associated with each tree node. To ensure that the same keys are promoted in different trees, a hash function is used to choose which elements are promoted. An important goal for C-trees is to maintain similar asymptotic cost bounds as for the uncompressed trees while improving space and cache performance, and to this end we describe theoretically efficient implementations of tree primitives in Section 4.

More formally. For an element type $K$, fix a hash function, $h : K \rightarrow \{1, \ldots, N\}$, drawn from a uniformly random family of hash functions ($N$ is some sufficiently large range). Let $b$ be a chunking parameter, a constant which controls the granularity of the chunks. Given a set $E$ of $n$ elements, we first compute the set of heads $H(E) = \{e \in E \mid h(e) \mod b = 0\}$. For each $e \in H(E)$ let its tail be $t(e) = \{x \in E \mid e < x < \text{next}(H(E), e)\}$, where $\text{next}(H(E), e)$ returns the next element in $H(E)$ greater than $e$. We then construct a purely-functional tree with keys $e \in H(E)$ and associated values $t(e)$.

Thus far, we have described the construction of a tree over the head elements, and their tails. However, there may be a "tail" at the beginning of $E$ that has no associated head, and is therefore not part of the tree. We refer to this chunk of elements as the prefix. We refer to either a tail or prefix as a chunk. We represent each chunk as a (variable-length) array of elements. As described later, when the elements are integers we can use difference encoding to compress each of the chunks. The overall C-tree data structure consists of the tree over head keys and tail values, and a single (possibly empty) prefix. Figure 1 illustrates the C-tree data structure over a set of integer elements.

Properties of C-trees. The expected size of chunks in a C-tree is $b$ as each element is independently selected as a head under $h$ with probability $1/b$. Furthermore, the chunks are unlikely to be much larger than $b$—in particular, a simple calculation shows that the chunks have size at most $O(b \log n)$ with high probability (w.h.p.),\footnote{We use \emph{with high probability (w.h.p.)} to mean with probability $1 − 1/n^c$ for some constant $c > 0$.} where $n$ is the number of elements in the tree. Notice that an element chosen to be a head will be a head in any C-trees containing it, a property that simplifies the implementation of primitives on C-trees.

Our chunking scheme has the following bounds, which we prove in the full version of the paper [26].
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When using a balanced binary tree for the
corollary 3.2.

In this section, we first discuss the improved space usage
by a factor of b can significantly reduce the size of the tree.

**Lemma 3.1.** The number of heads (keys) in a C-tree over a set E of n elements is O(n/b) w.h.p. Furthermore, the maximum size of a tail (the non-head elements associated with a head) or prefix is O(b log n) w.h.p.

**Corollary 3.2.** When using a balanced binary tree for the heads (one with O(log n) height for n keys), the height of a C-tree over a sequence E of n elements is O(log(n/b)) w.h.p.

**3.2 C-tree Compression**

In this section, we first discuss the improved space usage of C-trees relative to purely-functional trees without any assumption on the underlying type of elements. We then discuss how we can further reduce the space usage of the data structure in the case where the elements are integers.

**Space Usage and Locality.** Consider the layout of a C-tree compared to a purely-functional tree. By Lemma 3.1, the expected number of heads is O(n/b). Therefore, compared to a purely-functional tree, which allocates n tree nodes, we reduce the number of tree nodes allocated by a factor of b. As each tree node is quite large (in our implementation, each tree node is at least 32 bytes), reducing the number of nodes by a factor of b can significantly reduce the size of the tree. Experimental results are given in Section 7.1.

In a purely-functional tree, in the worst case, accessing each element will incur a cache miss, even in the case where elements are smaller than the size of a cache line. In a C-tree, however, by choosing b, the chunking parameter, to be slightly larger than the cache line size (≈ 128), we can store multiple elements contiguously within a single chunk and amortize the cost of a cache miss across all elements read from the chunk. Furthermore, note that the data structure can provide locality benefits even in the case when the size of an element is larger than the cache line size, as a modest value of b will ensure that reading all but the heads, which constitute an O(1/b) fraction of the elements, will be contiguous loads from the chunks.

**Integer C-trees.** In the case where the elements are integers, the C-tree data structure can exploit the fact that elements are stored in sorted order in the chunks to further compress the data structure. We apply a difference encoding scheme to each chunk. Given a chunk containing d integers, \{i_1, ..., i_d\}, we compute the differences \{i_1, i_2 - i_1, ..., i_d - i_{d-1}\}. The differences are then encoded using a byte-code [71, 81]. We applied byte-codes due to the fact that they are fast to decode while achieving most of the memory savings that are possible using a shorter code [12, 81].

Note that in the common case when b is a constant, the size of each chunk is small (O(log n) w.h.p.). Therefore, despite the fact that each chunk must be processed sequentially, the cost of the sequential decoding does not affect the overall work or depth of parallel tree methods. For example, mapping over all elements in the C-tree, or finding a particular element have the same asymptotic work as purely-functional trees and optimal (O(log n)) depth. To make the data structure dynamic, chunks must also be recompressed when updating a C-tree, which has a similar cost to decompressing the chunks. In the context of graph processing, the fact that methods over a C-tree are easily parallelizable and have low depth lets us avoid designing and implementing a more complex parallel decoding scheme, like the parallel byte-code in Ligra+ [71].

**3.3 Other Approaches**

Our data structure is loosely based on a previous sequential approach to chunking [11]. That approach was designed to be a generic addition to any existing balanced tree scheme for a dictionary and has overheads due to this goal.

Another option is to use B-trees [4]. However, the objective of a B-tree is to reduce the height of a search tree to accelerate searches in external memory, whereas our goal is to build a data structure that stores contiguous elements in a single node to make compression possible. The problem
with B-trees in our purely-functional setting is that we require path copying during functional updates, as illustrated in Figure 2. In our trees, this only requires copying a single binary node (32 or 40 bytes in our implementation) per level of the tree. For a B-tree, it would require copying B pointers (potentially thousands of bytes) per level of the tree, adding significant overhead in terms of memory and running time.

There is also work on chunking of functional trees for representing strings or (unordered) sequences [2, 9, 15, 31]. The motivation is similar (decrease space and increase locality), but the fact they are sequences rather than search trees makes the tradeoffs different. None of this work uses the idea of hashing or efficiently searching the trees. Using hashing to select the heads has an important advantage in simplifying the code, and proving asymptotic bounds. Keeping the elements with internal nodes and using a prefix allows us to access the first b elements (or so) in constant work.

4 Operations on C-trees

In this section, we show how to support various tree operations over C-trees, such as building, searching and performing batch-updates to the data structure. These are operations that we will need for efficiently processing and updating graphs. We argue that the primitives are theoretically efficient by showing bounds on the work and depth of each operation. We also describe how to support augmentation in the data structure using an underlying augmented purely-functional tree. We note that the C-tree interfaces defined in this section operate over element-value pairs, whereas the C-trees defined in Section 3.1 only stored a set of elements for the sake of illustration. The algorithm descriptions elide the code, and proving asymptotic bounds. Keeping the elements with internal nodes and using a prefix allows us to access the first b elements (or so) in constant work.

4.1 Algorithms for Batch Insertions and Deletions

Our MultiInsert and MultiDelete algorithms are based on more fundamental algorithms for Union, Intersection, and Difference on C-trees. Since we can simply build a tree over the input sequence to MultiInsert and call Union (or Difference for MultiDelete), we focus only on the set operations. Furthermore, because the algorithms for Intersection and Difference are conceptually very similar to the algorithm for Union, we only describe in detail the Union algorithm, and Split, an important primitive used to implement Union.

Union. Our Union algorithm (Algorithm 1) is based on the recursive algorithm for Union given by Blelloch et al. [13]. The main differences between the implementations are how to split a C-tree by a given element, and how to handle elements in the tails and prefixes. The algorithm takes as input two C-trees, C1 and C2, and returns a C-tree C containing the elements in the union of C1 and C2. Figure 3 provides an illustration of how our Union algorithm computes the union of two C-trees. The algorithms use the following operations defined on C-trees and chunks. The EXPOSE operation takes as input a tree and returns the left subtree, the element and prefix at the root of the tree, and the right subtree. The SPLIT operation takes as input a C-tree B and an element k, and returns two C-trees B1 and B2, where B1 (resp. B2) is a C-tree containing all elements less than (resp. greater than) k. It can also optionally return a boolean indicating whether k was found in B, which is used when implementing DIFFERENCE and INTERSECTION. SMALLEST operation returns the smallest head in a tree. UNIONBC merges a C-tree consisting of a prefix and empty tree, and another C-tree. We also use the SPLITCHUNK and UNIONCHUNK operations, which are defined similarly to SPLIT and UNION for chunks.

The idea of the algorithm is to call EXPOSE on the tree of one of the two C-trees (C2), and split the other C-tree (C1) based on the element exposed at the root of C2’s tree (Line 7). The split on C1 returns the trees B1 and B2 (Line 8). The

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### Algorithm 1 Union

1. function Union(C1, C2)
2. case (C1, C2) of
3. (null, _) → UNIONBC(C1, C2)
4. (_, null) → UNIONBC(C2, C1)
5. (T1, P1), (T2, P2) →
6. let
7. val (L2, k2, v2, R2) = EXPOSE(T2)
8. val (B1, (BT2, BP2)) = SPLIT(C1, k2)
9. val (vP2', vR') = SPLITCHUNK(v2, SMALLEST(BT2))
10. val (PL', PR) = SPLIT(BP2, vR')
11. val vP2' = UNIONCHUNK(vP2', PL')
12. val (C1, CR) = UNION(B1, L2, P2, ||
13. UNION(BT2, PR), (R2, vR'))
14. ctreeJoin(C2.Tree, CR.Tree, k2, vP2', C1.Prefix)
15. end

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algorithm then recursively calls \texttt{Union} on the $C$-trees constructed from $L_2$ and $R_2$, the left and right subtrees exposed in $C_2$’s tree with the $C$-trees returned by \texttt{Split}, $B_1$, and $B_2$.

However, some care must be taken, since elements in $k_2$’s tail, $v_2$, may come after some heads in $B_2$. Similarly, elements in $B_2$’s prefix may come after some heads of $R_2$. In both cases, we should merge these elements with their corresponding heads’ tails. We handle these cases by splitting $v_2$ by the leftmost element of $B_2$ (producing $v_{2l}$ and $v_{2g}$), and splitting $B_2$’s prefix by the leftmost element of $R_2$ (producing $P_L$ and $P_R$). The left recursive call to \texttt{Union} just takes the $C$-trees $B_1$ and $(L_2, P_2)$. The right recursive call takes the $C$-trees $(B_2, Tree, P_R)$, and $(R_2, v_{2g})$. Note that all elements in the prefixes $P_R$ and $v_{2g}$ are larger than the smallest head in $B_2$ and $R_2$. Therefore, the $C$-tree returned from the right recursive call has an empty prefix. The output of \texttt{Union} is the $C$-tree formed by joining the left and right trees from the recursive calls, $k_2$, and the tail $v_{2t}$ formed by unisoning $v_{2l}$ and $P_L$, with the prefix from $C_L$.

\textbf{Union-BC}. Recall that the \texttt{Union-BC} algorithm merges a $C$-tree consisting of a prefix and empty tree, and another $C$-tree. We give a detailed description and pseudocode of the algorithm in the full version of the paper due to space constraints [26]. The idea of \texttt{Union-BC} is to split the prefix based on the leftmost element of $P$’s tree into two pieces, $P_L$ and $P_R$ containing elements less than and greater than the leftmost element respectively. $P_L$ is merged with $P$’s prefix to generate $P'$. The elements in $P_R$ find the heads they correspond to by searching the tree for the largest head that is smaller than them. We then construct a sequence of head-tail pairs by inserting each element in $P_R$ into its corresponding elements. Finally, we generate a new tree, $T'$, by performing a \texttt{MultiInsert} into $C$’s tree with the updated head-tail pairs. The return value is the $C$-tree $(T', P')$.

\textbf{Split}. \texttt{Split} takes a $C$-tree, $C = (T, P)$, and an element $k$ and returns a triple consisting of a $C$-tree of all elements less than $k$, whether the element was found, and a $C$-tree of all elements greater than $k$. We provide a high-level description of the algorithm here and defer the pseudocode and details to the full version of the paper [26].

The algorithm works by enumerating cases for how the split key can split $C$. If $k$ is less than the first element in $P$, then we return an empty $C$-tree, false, indicating that $k$ was not found, and $C$ as the right $C$-tree. Similarly, if $k$ splits $P$ (it lies between the first and last elements of $P$) then we split $P$, and return the list of elements less than the split key as the left $C$-tree, with the boolean and right tree handled similarly. Otherwise, if the above cases did not match, and the tree is null, we return $C$ as the left $C$-tree. The recursive cases are similar to how \texttt{Split} is implemented in Blelloch et al. [13], except for the case where $k$ splits the tail at the root of the tree. Another important detail is how we compute the first and last elements of a chunk. Instead of scanning the chunk, which will cause us to do work proportional to the sum of chunks on a root-to-leaf path in the tree, we store the first and last elements at the head of each chunk to perform this operation in $O(1)$ work and depth. This modification is important to show that \texttt{Split} can be done in $O(b \log n)$ work and depth w.h.p. on a $C$-tree.

### 4.2 Work and Depth Bounds

We provide details, correctness proofs, and analysis for our $C$-tree primitives in the full version of the paper [26], and state the work and depth bounds below.

\textbf{Building}. Building $(\texttt{Build}(S, f))$ a $C$-tree can be done in $O(n \log n)$ work and $O(b \log n)$ depth w.h.p. where $|S| = n$.

\textbf{Searching}. Searching $(\texttt{Find}(T, e))$ for an element $e$ in a $C$-tree can be implemented in $O(b \log n)$ work and depth w.h.p.

\textbf{Mapping}. Mapping $(\texttt{Map}(T, f))$ over a $C$-tree containing $n$ elements with a constant-work function $f$ can be done in $O(n)$ work and $O(b \log n)$ depth w.h.p.

\textbf{Batch Updates}. Batch updates $(\texttt{MultiInsert}(T, f, S)$ and $\texttt{MultiDelete}(T, S))$ can be performed in $O(b^2(k \log(n/k) + 1))$ expected work and $O(b \log k \log n)$ depth w.h.p. where $k = \min(|T|, |S|)$ and $n = \max(|T|, |S|)$.

### 5 Representing Graphs as Trees

\textbf{Representation}. An undirected graph can be implemented using purely functional tree-based data structures by representing the set of vertices as a tree, which we call the \textbf{vertex-tree}. Each vertex in the vertex-tree represents its adjacency information by storing a tree of identifiers of its adjacent neighbors, which we call the \textbf{edge-tree}. Directed graphs can be represented in the same way by simply storing two edge-trees per vertex, one for the out-neighbors, and one for the in-neighbors. The resulting graph data structure is a tree-of-trees that has $O(\log n)$ overall depth using any balanced tree implementation (w.h.p. using a treap). Figure 4
The direction optimization \( [\text{vertex-tree}, \text{which takes value of the vertex-tree respectively, which can both be done vertices and number of edges in the graph by querying the } G \text{ in true} \) can also be compressed using a \( C \text{-tree but defer evaluating this idea for future work.} \)

**Basic Graph Operations.** We can compute the number of vertices and number of edges in the graph by querying the size (number of keys) in the vertex-tree and the augmented value of the vertex-tree respectively, which can both be done in \( O(1) \) work. Weighted graphs can be represented using the same structure, with the only difference being that the elements in the edge-trees are modified to store an associated edge weight. Note that computing associative functions over the weights (e.g., aggregating the sum of all edge-weights) could be easily done by augmenting the edge and vertex-trees. We also note that the vertex-tree could also be compressed using a \( C \)-tree but defer evaluating this idea for future work.

**EDGEMAP.** We implement EDGEMAP (defined in Section 2) by mapping over the vertices in the input vertexSubset \( U \) in parallel and for each vertex \( u \in U \) searching the vertex-tree for its edge-tree, and then mapping over \( u \)'s incident neighbors, again in parallel. For each of \( u \)'s neighbors \( v \), we apply the map function \( F(u, v) \) if the filter function \( C(v) \) returns true. Otherwise than finding vertices in the input vertexSubset in \( G \) and traversing edges via the tree instead of an array, the implementation is effectively the same as in Ligra [70].

The direction optimization [5, 70] can also be implemented, and we describe more details later in this section. Assuming the functions \( F \) and \( C \) take constant work, EDGEMAP takes \( O(\sum_{u \in U} \deg(u) + |U| \log n) \) work and \( O(\log n) \) depth.

**Batch Updates.** Inserting and deleting edges are defined similarly, and so we only provide details for INSERTEDGES. Note that updates (e.g., to the weight) of existing edges can be done within this interface. Let \( A \) be the sequence (batch) of edge updates and let \( k = |A| \).

We first sort the batch of edge pairs using a comparison sort. Next, we compute an array of source vertex IDs that are being updated and for each ID, in parallel, build a tree over its updated edges. We can combine duplicate updates in the batch by using the duplicate-combining function provided by the \( C \)-tree constructor. As the sequence is sorted, the build costs \( O(k \log k) \) work and \( O(3 n \log n) \) depth. Next, in the update step, we call \text{ MultiInsert } over the vertex-tree with each \( (source, tree) \) pair in the previous sequence. The combine function for \text{ MultiInsert } combines existing values (edge-trees) with the new edge-trees by calling \text{ Union } on the old edge-tree and new edge-tree.

We give a simple worst-case analysis of the algorithm and show that the algorithm performs \( O(k \log n) \) work overall, and has \( O(3 n \log n) \) depth. All steps before the \text{ MultiInsert } cost \( O(k \log k) \) work, and \( O(3 n \log n) \) depth in total, as they sort and apply parallel sequence operations to sequences of length \( k \) [39]. As the depth of both the vertex-tree and edge-tree is \( O(\log n) \), the overall work of updating both the vertex-tree and each affected edge-tree can be upper bounded by \( O(k \log n) \). The depth of \text{ MultiInsert } is \( O(3 n \log n + D_{\text{Union}}) \), where \( D_{\text{Union}} \) is the depth of union. This simplifies to \( O(3 n \log n) \) by upper-bounding \( D_{\text{Union}} \) as \( O(\log n) \), as shown in the full version of the paper [26].

### 5.1 Efficiently Implementing Graph Algorithms

We now address how to efficiently implement graph algorithms using a tree of \( C \)-trees, mitigating the increase in access times due to using trees. We first describe a technique for handling the asymptotic increase in work for global graph algorithms due to using trees. We then consider local algorithms, and argue that for many local algorithms, the extra cost of searching the vertex-tree can be amortized. Finally, we describe how direction optimization [5] can be easily implemented over the \( C \)-tree data structure.

**Flat Snapshots.** Notice that algorithms in our framework that use EDGEMAP incur an extra \( O(K \log n) \) factor in their work, where \( K \) is the total number of vertices accessed by EDGEMAP over the course of the algorithm. For an algorithm like breadth-first search, which runs in \( O(m + n) \) work and
$O(D \log n)$ depth for a graph with diameter $D$ using a static-graph processing framework [25], a naive implementation using our framework will require performing $O(m + n \log n)$ work (the depth is the same, assuming that $b$ is a constant).

Instead, for global graph algorithms, which we loosely define as performing $\Omega(n)$ work, we can afford to take a flat snapshot of the graph, which reduces the $O(K \log n)$ term to $O(K)$. The idea of a flat snapshot is very simple—instead of accessing vertices through the vertex-tree, and calling $\text{Find}$ for each $v$ supplied to $\text{edgeMap}$, we just precompute the pointers to the edge-trees for all $v \in V$ and store them in an array of size $n$. This can be done in linear work and $O(\log n)$ depth by traversing the vertex-tree once to fetch the pointers. By providing this array, which we call a flat snapshot to each call to $\text{edgeMap}$, we can directly access the edges tree in $O(1)$ work and reduce the work of $\text{edgeMap}$ on a vertexSubset, $U$, to $O(\sum_{u \in U} \text{deg}(u)) + |U|$. In practice, using a flat snapshot speeds up BFS queries on our input graphs by an average of 1.26x (see Table 4).

**Local Algorithms.** For local graph algorithms we often cannot afford to create a flat snapshot without a significant increase in the work. We observe, however, that after retrieving a vertex many local algorithms will process all edges incident to it. Because the average degree in real-world graphs is often in the same range or larger than $\log n$ (see Table 1), the logarithmic overhead of accessing a vertex in the vertex-tree in these graphs can be amortized against the cost of processing the edges incident to the vertex, on average.

**Direction Optimization.** Direction optimization is a technique first described for breadth-first search in Beamer et al. [5], and later generalized as part of Ligra in its $\text{edgeMap}$ implementation [70]. It combines a sparse traversal, which applies the $F$ function in $\text{edgeMap}$ to the outgoing neighbors of the input vertexSubset $U$, with a dense traversal, which applies $F$ to the incoming neighbors $u$ of all vertices $v$ in the graph where $C(v) = \text{true}$ and $u \in U$. The dense traversal improves locality for large input vertexSubsets, and reduces edge traversals in some algorithms, such as breadth-first search. The traversal mode on each iteration is selected based on the size of $U$ and its out-degrees. We implemented the optimization by implementing a sparse traversal and a dense traversal that traverses the underlying C-trees.

6 Aspen Graph-Streaming Framework

In this section, we outline the Aspen interface and implementation for processing streaming graphs, and provide the full interface in the full version of the paper [26]. The Aspen interface is an extension of Ligra’s interface. It includes the full Ligra interface—vertexSubsets, $\text{edgeMap}$, and various other functionality on a fixed graph. On top of Ligra, we add a set of functions for updating the graph—in particular, for inserting or deleting sets of edges or sets of vertices. We also add a flat-snapshot function. Aspen currently does not support weighted edges, but we plan to add this functionality using a similar compression scheme for weights as used in Ligra+ in the future. All of the functions for processing and updating the graph work on a fixed and immutable version (snapshot) of the graph. The updates are functional, and therefore instead of mutating the version, return a handle to a new graph. The implementation of these operations follow the description given in the previous sections.

The Aspen interface supports three functions, $\text{acquire}$, $\text{set}$, and $\text{release}$, for acquiring the current version of a graph, setting a new version, and releasing a the version. The interface is based on the recently defined version maintenance problem and implemented with the corresponding lock-free algorithm to solve it [8]. $\text{release}$ returns whether it is the last copy on that version, and if so we garbage collect it. The three functions each act atomically. The framework allows any number of concurrent readers (i.e., transactions that $\text{acquire}$ and $\text{release}$ but do not set) and a single writer ($\text{acquires}$, $\text{sets}$, and then $\text{releases}$). Multiple concurrent readers can acquire the same version, or different versions depending on how the writer is interleaved with them. The implementation of this interface is non-trivial due to race conditions between the three operations. Importantly, however, no reader or writer is ever blocked or delayed by other readers or writers. Aspen guarantees strict serializability, which means that the state of the graph and outputs of queries are consistent with some serial execution of the updates and queries corresponding to real time.

Aspen is implemented in C++ and uses PAM [74] as the underlying purely-functional tree data structure for storing the heads. Our C-tree implementation requires about 1400 lines of C++, most of which are for implementing $\text{Union}$, $\text{Difference}$, and $\text{Intersect}$. Our graph data structure uses an augmented purely-functional tree from PAM to store the vertex-tree. Each node in the vertex tree stores an integer C-tree storing the edges incident to each vertex as its value. We note that the vertex-tree could also be compressed using a C-tree, but we did not explore this direction in the present work. To handle memory management, our implementations use a parallel reference counting garbage collector along with a custom pool-based memory allocator. The pool-allocator is critical for achieving good performance due to the large number of small memory allocations in the the functional setting. Although C++ might seem like an odd choice for implementing a functional interface, it allows us to easily integrate with PAM and Ligra. We also note that although our graph interface is purely-functional (immutable), our global and local graph algorithms are not. They can mutate local state within their transaction, but can only access the shared graph through an immutable interface.

7 Experiments

**Algorithms.** We implemented five algorithms in Aspen, consisting of three global algorithms and two local algorithms.
Our global algorithms are breadth-first search (BFS), single-source betweenness centrality (BC), and maximal independent set (MIS). Our BC implementation computes the contributions to betweenness scores for shortest paths emanating from a single vertex. The algorithms are similar to the algorithms in [25] and required only minor changes to acquire a flat snapshot and include it as an argument to edgeMap. As argued in Section 5.1, the cost of creating the snapshot does not asymptotically affect the work or depth of our implementations. The work and depth of our implementations of BFS, BC, and MIS are identical to the implementations in [25]. Our local algorithms are 2-hop and Local-Cluster. 2-hop computes the set of vertices that are at most 2 hops away from the vertex using edgeMap. The worst-case work is $O(m + n \log n)$ and the depth is $O(\log n)$. Local-Cluster is a sequential implementation of the Nibble-Serial clustering algorithm [72, 73], run using $\epsilon = 10^{-6}$ and $\tau = 10$.

In our experiments, we run the global queries one at a time due to their large memory usage and significant internal parallelism, and run the local queries concurrently (many at the same time).

**Experimental Setup.** Our experiments are performed on a 72-core Dell PowerEdge R930 (with two-way hyper-threading) with $4 \times 2.4$GHz Intel 18-core E7-8867 v4 Xeon processors (with a 4800MHz bus and 45MB L3 cache) and 1TB of main memory. Our programs use a work-stealing scheduler that we implemented. The scheduler is implemented similarly to Cilk for parallelism. Our programs are compiled with the g++ compiler (version 7.3.0) with the -O3 flag. All experiments involving balanced-binary trees use weight-balanced trees as the underlying balanced tree implementation [13, 74]. We use Aspen to refer to the system using C-trees and difference encoding within each chunk and explicitly specify other configurations of the system if necessary.

**Graph Data.** Table 1 lists the graphs we use. LiveJournal is a directed graph of the LiveJournal social network [16]. com-Orkut is an undirected graph of the Orkut social network. Twitter is a directed graph of the Twitter network, where edges represent the follower relationship [45]. ClueWeb is a Web graph from the Lemur project at CMU [16]. Hyperlink2012 and Hyperlink2014 are directed hyperlink graphs obtained from the WebDataCommons dataset where nodes represent web pages [51]. Hyperlink2012 is the largest publicly-available graph, and we show that Aspen is able to process it on a single multicore machine. We symmetrized the graphs in our experiments, as the running times for queries like BFS and BC are more consistent on undirected graphs due to the majority of vertices being in a single large component.

**Overview of Results.** We show the following experimental results in this section.

- The most memory-efficient representation of C-trees saves between 4–11x memory over using uncompressed trees, and improves performance by 2.5–2.8x compared to using uncompressed trees (Section 7.1).
- Algorithms implemented using Aspen are scalable, achieving between 32–78x speedup across inputs (Section 7.2).
- Updates and queries can be run concurrently in Aspen with only a slight increase in latency (Section 7.3).
- Parallel batch updates in Aspen are efficient, achieving between 105–442M updates/sec for large batches (Section 7.4).
- Aspen outperforms Stinger by 1.8–10.2x while using 8.5–11.4x less memory (Section 7.5).
- Aspen outperforms LLAMA by 2.8–7.8x while using 1.9–3.5x less memory (Section 7.6).
- Aspen is competitive with state-of-the-art static graph processing systems, ranging from being 1.4x slower to 30x faster (Section 7.7).
to the uncompressed format. Using difference encoding provides between 1.2–2.3x reduction in memory usage compared to storing the chunks in an uncompressed format. We observe that both using C-trees and compressing within the chunks is crucial for storing and processing our largest graphs in a reasonable amount of memory.

**Comparison with Uncompressed Trees.** Next, we study the performance improvement gained by the improved locality of the C-tree data structure. Due to the memory overheads of representing large graphs using the uncompressed format (see Table 1), we are only able to report results for our three smallest graphs, LiveJournal, com-Orkut, and Twitter, as we cannot store the larger graphs even with 1TB of RAM in the uncompressed format. We ran BFS on both the uncompressed and C-tree formats (using difference encoding) and show the results in the full version of our paper due to space constraints [26]. The results show that using the compressed representation improves the running times of these applications from between 2.5–2.8x across these graphs.

**Choice of Chunk Size.** Next, we consider how Aspen performs as a function of the expected chunk size, $b$. Table 3 reports the amount of memory used, and the BFS, BC, and MIS running times as a function of $b$. In the rest of the paper, we fixed $b = 2^8$, which we found gave the best tradeoff between the amount of memory consumed (it requires 5% more memory than the most memory-efficient configuration) while enabling good parallelism across different applications.

<table>
<thead>
<tr>
<th>Application</th>
<th>LiveJournal</th>
<th>com-Orkut</th>
<th>Twitter</th>
<th>ClueWeb</th>
<th>Hyperlink2014</th>
<th>Hyperlink2012</th>
</tr>
</thead>
<tbody>
<tr>
<td>BFS</td>
<td>0.981</td>
<td>0.021</td>
<td>0.690</td>
<td>0.015</td>
<td>0.138</td>
<td>52.6</td>
</tr>
<tr>
<td>BC</td>
<td>4.66</td>
<td>0.075</td>
<td>62.1</td>
<td>4.58</td>
<td>0.078</td>
<td>81.2</td>
</tr>
<tr>
<td>MIS</td>
<td>3.38</td>
<td>0.054</td>
<td>62.0</td>
<td>4.19</td>
<td>0.069</td>
<td>71.5</td>
</tr>
</tbody>
</table>

Table 2. Running times (in seconds) of our algorithms over symmetric graph inputs where (1) is the single threaded time (72h) is the 72-core time (with hyper-threading, i.e., 144 threads), and (SU) is the self-relative speedup.

<table>
<thead>
<tr>
<th>$b$ (Exp. Chunk Size)</th>
<th>Memory USE (SU)</th>
<th>BFS (72h)</th>
<th>BC (72h)</th>
<th>MIS (72h)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2**1</td>
<td>68.83</td>
<td>0.309</td>
<td>2.72</td>
<td>2.17</td>
</tr>
<tr>
<td>2**2</td>
<td>41.72</td>
<td>0.245</td>
<td>2.09</td>
<td>1.71</td>
</tr>
<tr>
<td>2**3</td>
<td>26.0</td>
<td>0.217</td>
<td>1.68</td>
<td>1.41</td>
</tr>
<tr>
<td>2**4</td>
<td>17.7</td>
<td>0.172</td>
<td>1.45</td>
<td>1.24</td>
</tr>
<tr>
<td>2**5</td>
<td>13.3</td>
<td>0.162</td>
<td>1.32</td>
<td>1.14</td>
</tr>
<tr>
<td>2**6</td>
<td>11.1</td>
<td>0.152</td>
<td>1.25</td>
<td>1.07</td>
</tr>
<tr>
<td>2**7</td>
<td>9.97</td>
<td>0.142</td>
<td>1.22</td>
<td>1.01</td>
</tr>
<tr>
<td>2**8</td>
<td>9.42</td>
<td>0.138</td>
<td>1.18</td>
<td>0.99</td>
</tr>
<tr>
<td>2**9</td>
<td>9.17</td>
<td>0.141</td>
<td>1.20</td>
<td>0.99</td>
</tr>
<tr>
<td>2**10</td>
<td>9.03</td>
<td>0.152</td>
<td>1.19</td>
<td>0.98</td>
</tr>
<tr>
<td>2**11</td>
<td>8.96</td>
<td>0.163</td>
<td>1.20</td>
<td>0.98</td>
</tr>
<tr>
<td>2**12</td>
<td>8.89</td>
<td>0.170</td>
<td>1.21</td>
<td>0.98</td>
</tr>
</tbody>
</table>

Table 3. Memory usage (gigabytes) and performance (seconds) for the Twitter graph as a function of the (expected) chunk size. All times are measured on 72 cores using hyper-threading. Bold text marks the best value in each column. We use $2^8$ in the other experiments.

**Algorithm Performance.** Table 2 reports experimental results including the single-threaded time and 72-core time (with hyper-threading) for Aspen using compressed C-trees. For BFS, we achieve between 46–70x speedup across all inputs. For BC, our implementations achieve between 50–78x speedup across all inputs. Finally, for MIS, our implementations achieve between 60x–78x speedup across all inputs. We observe that the experiments in [25] report similar speedups for the same graphs. For local algorithms, we report the average running time for performing 2048 queries sequentially and in parallel. We achieve between 41–43x speedup for 2-hop, and between 35–49x speedup for Local-Cluster.

**Flat Snapshots.** Table 4 shows the running times of BFS with and without the use of a flat snapshot. Our BFS implementation is between 1.12–1.3x faster using a flat snapshot, including the time to compute a flat snapshot. The table also reports the time to acquire a flat snapshot, which is between 15–24% of the overall BFS time across all graphs. We observe that acquiring a flat snapshot is already an improvement for a single run of an algorithm, and quickly becomes more profitable as multiple algorithms are run over a single snapshot of the graph (e.g., multiple BFS’s or betweenness centrality computations).

**7.3 Simultaneous Updates and Queries**

In this sub-section, we experimentally verify that Aspen can support low-latency queries and updates running concurrently. In these experiments, we generate an update stream by randomly sampling 2 million edges from the input graph to use as updates. We sub-sample 90% of the sample to use as edge insertions, and immediately delete them from the input graph. The remaining 10% are kept in the graph, as
we will delete them over the course of the update stream. The update stream is a random permutation of these insertions and deletions. We believe that sampling edges from the input graph better preserves the properties of the graph and ensures that edge deletions perform non-trivial work, compared to using random edge updates.

After constructing the update stream, we spawn two parallel jobs, one which performs the updates sequentially and one which performs global queries. We maintain the undirectedness of the graph by inserting each edge as two directed edge updates, within a single batch. For global queries, we run a stream of BFS’s from random sources one after the other and measure the average latency. We note that for the BFS queries, as our inputs are symmetrized, a random vertex is likely to fall in the giant connected component which exists in all of our input graphs. The global queries therefore process nearly all of the vertices and edges.

Table 5 shows the throughput in terms of directed edge updates per second, the average latency to make an undirected edge visible, and the latency of global queries both when running concurrently with updates and when running in isolation. We note that when running global queries in isolation, we use all of the threads in the system (72-cores with hyper-threading). We observe that our data structure achieves between 22–157 thousand directed edge updates per second, which is achieved while concurrently running a parallel query on all remaining threads. We obtain higher update rates on smaller graphs, where the small size of the graph enables it to utilize the caches better. In all cases, the average latency for making an edge visible is at most 86 microseconds, and is as low as 12.7 microseconds on the smallest graph.

The last two columns in Table 5 show the average latency of BFS queries from random sources when running queries concurrently with updates, and when running queries in isolation. We see that the performance impact of running updates concurrently with queries is less than 3%, which could be due to having one fewer thread. We ran a similar experiment, where we ran updates on 1 core and ran multiple concurrent local queries (Local-Cluster) on the remaining cores, and found that the difference in average query times is even lower than for BFS.

### 7.4 Performance of Batch Updates

In this sub-section, we show that the batch versions of our primitives achieve high throughput when updating the graph, even on very large graphs and for very large batches. As there are insufficient edges on our smaller graphs for applying the methodology from Section 7.3, we sample directed edges from an rMAT graph generator with $a = 0.5, b = c = 0.1, d = 0.3$ to perform the updates. To evaluate our performance on a batch of size $B$, we generate $B$ directed edge updates from the stream (note that there can be duplicates), repeatedly call `InsertEdges` and `DeleteEdges` on the batch, and report the median of three such trials. The costs that we report include the time to sort the batch and combine duplicates.

Table 6 shows the throughput (the number of edges processed per second) of performing batch edge insertions in parallel on varying batch sizes. The throughput for edge deletions are within 10% of the edge insertion times, and are usually faster (see Figure 5). The running time can be calculated by dividing the batch size by the throughput. We illustrate the throughput obtained for both insertions and deletions in Figure 5 for the largest and smallest graph, and note that the lines for other graphs are sandwiched between these two lines. The only exception of com-Orkut, where batch insertions achieve about 2% higher throughput than soc-LiveJournal at the two largest batch sizes.

We observe that Aspen’s throughput seems to vary depending on the graph size. We achieve a maximum throughput of 442M updates per second on com-Orkut when processing batches of 2B updates. On the other hand, on the Hyperlink2012 graph, the largest graph that we tested on, we achieve 105M updates per second for this batch size. We believe that the primary reason that small graphs achieve much better throughput at the largest batch size is that nearly all of the vertices in the tree are updated for the small graphs.

### Table 5. Throughput and average latency achieved by Aspen when concurrently processing a sequential stream of edge updates along with a sequential stream of breadth-first search queries (each BFS is internally parallel). Latency (C) reports the average latency of the query when running the updates and queries concurrently, while Latency (I) reports the average latency when running queries in isolation on the modified graph.

<table>
<thead>
<tr>
<th>Graph</th>
<th>Update Edges/sec</th>
<th>Latency (BFS)</th>
<th>Latency (C)</th>
<th>Latency (I)</th>
</tr>
</thead>
<tbody>
<tr>
<td>LiveJournal</td>
<td>7.86e4</td>
<td>1.27e-5</td>
<td>0.0190</td>
<td>0.0185</td>
</tr>
<tr>
<td>com-Orkut</td>
<td>6.02e4</td>
<td>1.66e-5</td>
<td>0.0179</td>
<td>0.0176</td>
</tr>
<tr>
<td>Twitter</td>
<td>4.44e4</td>
<td>1.73e-5</td>
<td>0.155</td>
<td>0.155</td>
</tr>
<tr>
<td>ClueWeb</td>
<td>2.06e4</td>
<td>4.83e-5</td>
<td>4.83</td>
<td>4.82</td>
</tr>
<tr>
<td>Hyperlink2014</td>
<td>1.42e4</td>
<td>7.04e-5</td>
<td>6.17</td>
<td>6.15</td>
</tr>
<tr>
<td>Hyperlink2012</td>
<td>1.16e4</td>
<td>8.57e-5</td>
<td>15.8</td>
<td>15.5</td>
</tr>
</tbody>
</table>

### Table 6. Throughput (directed edges/second) obtained when performing parallel batch edge insertions on different graphs with varying batch sizes, where inserted edges are sampled from an rMAT graph generator. The times for batch deletions are similar to the time for insertions. All times are on 72 cores with hyper-threading.

<table>
<thead>
<tr>
<th>Graph</th>
<th>Batch Size</th>
<th>10</th>
<th>10^3</th>
<th>10^5</th>
<th>10^7</th>
<th>2 - 10^9</th>
</tr>
</thead>
<tbody>
<tr>
<td>LiveJournal</td>
<td></td>
<td>8.26e4</td>
<td>2.88e6</td>
<td>2.29e7</td>
<td>1.56e8</td>
<td>4.13e8</td>
</tr>
<tr>
<td>com-Orkut</td>
<td></td>
<td>7.14e4</td>
<td>2.79e6</td>
<td>2.22e7</td>
<td>1.51e8</td>
<td>4.21e8</td>
</tr>
<tr>
<td>Twitter</td>
<td></td>
<td>6.32e4</td>
<td>2.63e6</td>
<td>1.23e7</td>
<td>5.68e7</td>
<td>3.04e8</td>
</tr>
<tr>
<td>ClueWeb</td>
<td></td>
<td>6.57e4</td>
<td>2.38e6</td>
<td>7.19e6</td>
<td>2.64e7</td>
<td>1.33e8</td>
</tr>
<tr>
<td>Hyperlink2014</td>
<td></td>
<td>6.17e4</td>
<td>2.12e6</td>
<td>6.66e6</td>
<td>2.28e7</td>
<td>9.90e7</td>
</tr>
<tr>
<td>Hyperlink2012</td>
<td></td>
<td>6.46e4</td>
<td>2.04e6</td>
<td>4.97e6</td>
<td>1.84e7</td>
<td>8.26e7</td>
</tr>
</tbody>
</table>
In this case, due to the asymptotic work bound for the update algorithm, the work for our updates become essentially linear in the tree size.

7.5 Comparison with Stinger

In this sub-section, we compare Aspen to Stinger [29], a state-of-the-art graph-streaming system.

Stinger Design. Stinger’s data structure for processing streaming graphs is based on adapting the CSR format to support dynamic updates. Instead of storing all edges of a vertex contiguously, it chunks the edges into a number of blocks, which are chained together as a linked list. Updates traverse the list to find an empty slot for a new edge, or to determine whether an edge exists. Therefore, updates take $O(deg(v))$ work and depth for a vertex $v$ that is updated. Furthermore, updates use fine-grained locking to perform edge insertions, which may result in contention when updating very high degree vertices. As Stinger does not support compressed graph inputs, we were unable to run the system on our input graphs that are larger than Twitter.

Memory Usage. We list the sizes of the three graphs that Stinger was able to process in Table 7. The Stinger interface supports a function which returns the size of its in-memory representation in bytes, which is what we use to report the numbers in this paper.

We found that Stinger has a high memory usage, even in the memory-efficient settings used in our experiments. The memory usage we observed appears to be consistent with [29], which reports that the system requires 313GB of memory to store a scale-free (RMAT) graph with 268 million vertices and 2.15 billion edges, making the cost 145 bytes per edge. This number is on the same order of magnitude as the numbers we report in Table 7. We found that Aspen is between 8.5–11.4x more memory efficient than Stinger.

Batch Update Performance. We measure the batch update performance of Stinger by using an rMAT generator provided in Stinger to generate the directed updates. We set $n = 2^{30}$ for updates in the stream. The largest batch size supported by Stinger is 2M directed updates. The update times for Stinger were fastest when inserting into nearly-empty graphs. For each batch size, we insert 10 batches of edges of that size into the graph, and report the median time.

The results in Table 8 show the update rates for inserting directed edge updates in Stinger and Aspen. We observe that the running time for Stinger is reasonably high, even on very small batches, and grows linearly with the size of the batch. The Aspen update times also grow linearly, but are very fast for small batches. Perhaps surprisingly, our update time on a batch of 1M updates is faster than the update time of Stinger on a batch of 10 edges.

Algorithm Performance. Lastly, we show the performance of graph algorithms implemented using the Stinger data structures. We use the BFS implementation for Stinger developed in McColl et al. [49]. We used a BC implementation that is available in the Stinger code base. Unfortunately, this implementation is entirely sequential, and so we compare Stinger’s BC time to our single-threaded time. Neither of the Stinger implementations perform direction-optimization, so to perform a fair comparison, we used an implementation of BFS and BC in Aspen that disables direction-optimization. Table 9 shows the parallel running times of of Stinger and Aspen for these problems. For BFS, which is run in parallel, we achieve between 6.7–10.2x speedup over Stinger.

![Figure 5. Throughput (edges/sec) when performing batches of insertions (I) and deletions (D) with varying batch sizes on Hyperlink2012 and LiveJournal in a log-log scale. All times are on 72 cores with hyper-threading.](image-url)
the parallel running times of our three algorithms implemented in Stinger (ST), LLAMA (LL), and Aspen. A is the parallel time using Aspen without direction-optimization. (A(I)) is the one-thread time of Aspen which is only relevant for comparing with Stinger’s BC implementation. A† is the parallel time using Aspen with direction-optimization. (ST/A) is Aspen’s speedup over Stinger and (LL/A) is Aspen’s speedup over LLAMA.

For BC, which is run sequentially, we achieve between 1.8–4.2x speedup over Stinger. A likely reason that Aspen’s BFS is significantly faster than Stinger’s is that it can process edges incident to high-degree vertices in parallel, whereas traversing a vertex’s neighbors in Stinger requires sequentially traversing a linked list of blocks.

### 7.6 Comparison with LLAMA

In this sub-section, we compare Aspen to LLAMA [47], another state-of-the-art graph-streaming system.

**LLAMA Design.** Like Stinger, LLAMA’s streaming graph data structure is motivated by the CSR format. However, like Aspen, LLAMA is designed for batch-processing in the single-writer multi-reader setting and can provide serializable snapshots. In LLAMA, a batch of size is generates a new snapshot which uses space to store a vertex array, and space to store edge updates in a dynamic CSR structure. The structure creates a linked list over the edges incident to a vertex that is linked over multiple snapshots. This design can cause the depth of iterating over the neighbors of a vertex to be large if the edges are spread over multiple snapshots.

Unfortunately, the publicly-available code for LLAMA does not provide support for evaluating streaming graph algorithms or batch updates. However, we were able to load static graphs and run several implementations of algorithms in LLAMA for which we report times in this section. As LLAMA does not support compressed graphs, we were unable to run the system on the graphs larger than Twitter.

**Memory Usage.** Unfortunately, we were not able to get LLAMA’s internal allocator to report correct memory usage statistics for its internal allocations. Instead, we measured the lifetime memory usage of the process and use this as an estimate for the size of the in-memory data structure built by LLAMA. The memory usage in bytes for the three graphs that LLAMA was able to process is shown in Table 7. The cost in terms of bytes/edge for LLAMA appears to be consistent, which matches the fact that the internal representation is a flat CSR, since there is a single snapshot. Overall, Aspen is between 1.9–3.5x more memory efficient than LLAMA.

### 7.7 Static Graph Processing Systems

We compared Aspen to Ligra+, a state-of-the-art shared-memory graph processing system, GAP, a state-of-the-art graph processing benchmark [6], and Galois, a shared-memory parallel programming library for C++ [56].

**Ligra+.** Table 10 the parallel running times of our three global algorithms expressed using Aspen and Ligra+. The results show that Ligra is 1.43x faster than Aspen for global algorithms on small inputs. We also performed a more extensive experimental comparison between Aspen and Ligra+, comparing the parallel running times of all of our algorithms on all of our inputs, which we include in the full version of the paper [26]. Compared to Ligra+, across all inputs, algorithms in Aspen are 1.51x slower on average (between 1.2x–1.7x) for the global algorithms, and 1.45x slower on average (between 1.0–2.1x) for the local algorithms. We report the local algorithms in the full version of the paper [26].

The local algorithms have a modest slowdown compared to their Ligra+ counterparts, due to logarithmic work vertex accesses being amortized against the relative high average degrees (see Table 1).

**GAP.** Table 10 shows the parallel running times of the BFS and BC implementations from GAP. On average, our implementations in Aspen are 1.4x faster than the implementations from GAP over all problems and graphs. We note that the
code in GAP has been hand-optimized using OpenMP scheduling primitives. As a result, the GAP code is significantly more complex than our code, which only uses the high-level primitives defined by Ligra+.

Galois. Table 10 shows the running times of using Galois, a shared-memory parallel programming library that provides support for graph processing [56]. Galois’ algorithms (e.g., for BFS and MIS) come with several versions. In our experiments, we tried all versions of their algorithms, and report times for the fastest one. On average, our implementations in Aspen are 12x faster than Galois. For BFS, Aspen is between 3.3–4.4x faster than Galois. We note that the Galois BFS implementation is synchronous, and does not appear to use Beamer’s direction-optimization. We omit BC as we were not able to obtain reasonable numbers on our inputs using their publicly-available code (the numbers we obtained were much worse than the ones reported in [56]). For MIS, our implementations are between 9–30x faster than Galois.

8 Related Work

We have mentioned some other schemes for chunking in Section 3.3. Although we use functional trees to support snapshots, many other systems for supporting persistence and snapshots use version lists [7, 27, 62]. The idea is for each mutable value or pointer to keep a timestamped list of versions, and reading a structure to go through the list to find the right one (typically the most current is kept first). LLAMA [47] uses a variation of this idea. However, it seems challenging to achieve the low space that we achieve using such systems since the space for such a list is large.

Graph Processing Frameworks. Many processing frameworks have been designed to process static graphs (e.g., [23, 33, 46, 48, 56, 59, 61, 70, 79], among many others). We refer the reader to [50, 82] for surveys of existing frameworks. Similar to Ligra+ [71], Log(Graph) [10] supports running parallel algorithms on compressed graphs. Their experiments show that they have a moderate performance slowdown on real-world graphs, but sometimes get improved performance on synthetic graphs [10].

Existing dynamic graph streaming frameworks can be divided into two categories based on their approach to ingesting updates. The first category processes updates and queries in phases, i.e., updates wait for queries to finish before updating the graph, and queries wait for updates to finish before viewing the graph. Most existing systems take this approach, as it allows updates to mutate the underlying graph without worrying about the consistency of queries [3, 18, 19, 29, 30, 34, 54, 65–67, 69, 75, 78, 80]. Hornet [18], one of the most recent systems in this category, reports a throughput of up to 800 million edges per second on a GPU with 3,840 cores (about twice our throughput using 72 CPU cores for similarly-sized graphs); however the graphs used in Hornet are much smaller that what Aspen can handle due to memory limitations of GPUs. The second category enables queries and updates to run concurrently by isolating queries to run on snapshots and periodically have updates generate new snapshots [21, 37, 38, 47].

GraphOne [44] is a system developed concurrently with our work that can run queries on the most recent version of the graph while processing updates concurrently by using a combination of an adjacency list and an edge list. They report an update rate of 66.4 million edges per second on a Twitter graph with 2B edges using 28 cores; Aspen is able to ingest 94.5 million edges per second on a larger Twitter graph using 28 cores. However, GraphOne also backs up the update data to disk for durability.

There are also many systems that have been built for analyzing graphs over time [32, 35, 36, 41, 42, 52, 53, 63, 76, 77]. These systems are similar to processing graph streams in that updates to the graph must become visible to new queries, but are different in that queries can performed on the graph as it appeared at any point in time. Although we do not explore historical queries in this paper, functional data structures are particularly well-suited for this task since it is easy to keep any number of persistent versions by keeping their roots.

Graph Databases. There has been significant research on graph databases (e.g., [17, 28, 40, 43, 55, 60, 68]). The main difference between graph-streaming and graph databases is that graph databases support transactions, i.e., multi-writer concurrency. A graph database running with snapshot isolation could be used to solve the same problem we solve. However, due to the need to support transactions, graph databases have significant overhead even for graph analytic queries such as PageRank and shortest paths. McColl et al. [49] show that Stinger is orders of magnitude faster than state-of-the-art graph databases.

9 Conclusion

We have presented a compressed fully-functional tree data structure called the C-tree that has theoretically-efficient operations, low space usage, and good cache locality. We use C-trees to represent graphs, and design a graph-streaming framework called Aspen that is able to support concurrent queries and updates to the graph with low latency. Experiments show that Aspen outperforms state-of-the-art graph-streaming frameworks, Stinger and LLAMA, and only incurs a modest overhead over state-of-the-art static graph processing frameworks. Future work includes designing incremental graph algorithms and historical queries using Aspen, and using C-trees in other applications. Although our original motivation for designing C-trees was for representing compressed graphs, we believe that they are of independent interest and can be used in applications where ordered sets of integers are dynamically maintained, such as compressed inverted indices in search engines.


