GAMMA: Leveraging Gustavson’s Algorithm to Accelerate Sparse Matrix Multiplication

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ABSTRACT
Sparse matrix-sparse matrix multiplication (spMspM) is at the heart of a wide range of scientific and machine learning applications. spMspM is inefficient on general-purpose architectures, making accelerators attractive. However, prior spMspM accelerators use inner- or outer-product dataflows that suffer poor input or output reuse, leading to high traffic and poor performance. These prior accelerators have not explored Gustavson’s algorithm, an alternative spMspM dataflow that does not suffer from these problems but features irregular memory access patterns that prior accelerators do not support.

We present GAMMA, an spMspM accelerator that uses Gustavson’s algorithm to address the challenges of prior work. GAMMA performs spMspM’s computation using specialized processing elements with simple high-radix mergers, and performs many merges in parallel to achieve high throughput. GAMMA uses a novel on-chip storage structure that combines features of both caches and explicitly managed buffers. This structure captures Gustavson’s irregular reuse patterns and streams thousands of concurrent sparse fibers (i.e., lists of coordinates and values for rows or columns) with explicitly decoupled data movement. GAMMA features a new dynamic scheduling algorithm to achieve high utilization despite irregularity. We also present new preprocessing algorithms that boost GAMMA’s efficiency and versatility. As a result, GAMMA outperforms prior accelerators by gmean 2.1x, and reduces memory traffic by gmean 2.2x and by up to 13x.

CCS CONCEPTS
• Computer systems organization → Architectures.

KEYWORDS
sparse matrix multiplication, sparse linear algebra, accelerator, Gustavson’s algorithm, high-radix merge, explicit data orchestration, data movement reduction

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1 INTRODUCTION
Scientific and machine learning applications are increasingly computing on sparse data, i.e., data where a large fraction of values are zeros. In this work, we focus on accelerating sparse matrix-sparse matrix multiplication (spMspM), a key kernel that lies at the heart of many sparse algorithms, like sparse deep neural networks [18, 39], sparse linear and tensor algebra [29, 57], graph analytics [16, 27], and simulation [6].

spMspM has two key characteristics that make it challenging to accelerate. First, spMspM is bottlenecked by memory traffic and data movement: it requires far fewer arithmetic operations per input element than dense matrix multiplication, and its inputs and outputs typically use a compressed representation that omits zeros but is more complicated to traverse, requiring irregular and indirect accesses. Thus, to be effective, accelerators must minimize data movement, rather than compute operations. Second, spMspM has a rich algorithmic diversity: it admits a wide range of dataflows (i.e., computation schedules) with different tradeoffs, and some dataflows have asymptotically worse performance on particular inputs. Thus, accelerators must achieve efficiency though specialization while avoiding the inefficiencies of using an inadequate spMspM dataflow.

Prior work has proposed spMspM accelerators that greatly improve performance over CPUs and GPUs. And yet, these accelerators have focused on one of two spMspM dataflows, inner-product [20, 43] or outer-product [37, 59], which have significant drawbacks (Sec. 2). Inner-product maximizes output reuse but sacrifices reuse of input matrices, and is inefficient with highly sparse matrices, as it is dominated by the cost of intersections that do not produce output values. By contrast, outer-product maximizes input reuse, but sacrifices output reuse, as it suffers from the cost and memory traffic of merging large partial output matrices. Prior accelerators have missed a third spMspM dataflow, Gustavson’s algorithm [17], which is often the most efficient dataflow and is widely used in CPUs and GPUs [15, 29, 52]. Gustavson’s algorithm often achieves the least amount of memory traffic and requires simpler operations because it avoids the extremes of inner- and outer-product. However, Gustavson’s algorithm has more irregular reuse across data structures, demanding a storage organization that can exploit that reuse to reduce memory traffic.

MatRaptor [48], which was published after the submission of this work, is an accelerator that exploits Gustavson’s algorithm. We discuss it briefly in Sec. 7.
To unlock the potential of \texttt{spMspM} acceleration, we propose \textsc{Gamma}, the Gustavson-Algorithm Matrix-Multiplication Accelerator (Sec. 3). \textsc{Gamma} combines three key features:

(1) \textsc{Gamma} uses simple processing elements (PEs) that linearly combine sparse input rows to produce each output row. PEs implement high-radix mergers that combine many input rows (e.g., 64 in our design) in a single pass, reducing work and memory accesses. Instead of expensive high-throughput mergers as in prior work [59], \textsc{Gamma} uses simple scalar mergers, and relies on Gustavson\textquotesingle s row-level parallelism to achieve high throughput efficiently, using tens of PEs to perform many combinations in parallel. Thus, \textsc{Gamma} concurrently processes \textit{thousands of compressed sparse fibers}, variable-sized rows from inputs or partial outputs.

(2) \textsc{Gamma} uses a novel storage structure, \textsc{FiberCache}, to efficiently buffer the thousands of fibers required by PEs. \textsc{FiberCache} is organized as a cache to capture Gustavson\textquotesingle s irregular reuse patterns. However, \textsc{FiberCache} is managed explicitly, like a large collection of buffers, to fetch missing fibers ahead of time and avoid PE stalls. This saves megabytes of dedicated on-chip buffers.

(3) \textsc{Gamma} dynamically schedules work across PEs to ensure high utilization and minimize memory traffic despite the irregular nature of Gustavson\textquotesingle s algorithm.

While Gustavson\textquotesingle s algorithm is an improvement over other dataflows, it still incurs excessive traffic on some inputs. To address this issue, we propose a preprocessing technique (Sec. 4), that combines row reordering and selective tiling of one matrix input. Preprocessing improves \textsc{Gamma}’s performance and avoids pathologies across the full range of inputs.

We synthesize \textsc{Gamma} and evaluate its performance on a wide range of sparse matrices (Sec. 6). Compared to state-of-the-art accelerators, with a similar hardware budget, \textsc{Gamma} reduces total DRAM traffic by 2.2× on average, non-compulsory DRAM traffic by 12× on average, and achieves significantly higher DRAM bandwidth utilization. Moreover, \textsc{Gamma} is effective on a much broader range of sparse matrices.

In summary, we make the following contributions:

- We show that prior \texttt{spMspM} accelerators have missed a key dataflow, Gustavson\textquotesingle s, which is often more efficient but has less regular access patterns than previously used dataflows.
- We build \textsc{Gamma}, a novel \texttt{spMspM} accelerator that combines specialized PEs, a novel cache-based structure to capture Gustavson\textquotesingle s irregular reuse, and dynamic scheduling to achieve high utilization despite irregularity.
- We propose preprocessing techniques that boost \textsc{Gamma}’s effectiveness and avoid Gustavson\textquotesingle s pathologies.
- We evaluate \textsc{Gamma} under a broad range of matrices, showing large performance gains and memory traffic reductions over prior systems, as well as higher versatility.

2 BACKGROUND AND MOTIVATION

Sparse matrix-sparse matrix multiplication (\texttt{spMspM}) is widely used in deep learning inference [18, 39, 54], linear algebra [5, 29, 57], and graph analytics [16, 27] (including breadth-first search [16], maximum matching [44], cycle detection [58], triangle counting [2], clustering [50], and all-pair shortest paths [7]). It is also a key building block for many other workloads, such as parsing [41], searching [25], and optimization [26].

We first describe the data structures used by \texttt{spMspM} and the basic \texttt{spMspM} dataflows; then, we review prior accelerators, the optimizations they introduce, and their limitations, motivating the need for a Gustavson-based accelerator.

2.1 Compressed Sparse Data Structures

\texttt{spMspM} operates on compressed sparse data structures, i.e., structures where only nonzeros are represented. Fig. 1 shows a sparse matrix encoded in two commonly used formats, compressed sparse row (CSR) and compressed sparse column (CSC). In CSR, rows are stored in a compressed format: each row is an ordered list of coordinates (in this case, column indexes) and nonzero values, stored contiguously. Indexing into a particular row is achieved through the offsets array, which stores the starting position of each row. CSC is analogous to CSR, but stores the matrix by compressed columns. In general, we call each compressed row or column a fiber, represented by a list of coordinates and values, sorted by coordinate.

Compressed sparse data structures introduce two challenges. First, certain kinds of traversals, called concordant traversals [49], are more efficient than others. For example, a CSR matrix can be traversed row by row, but traversing it by columns or accessing elements at random coordinates is inefficient. Thus, to be efficient, different \texttt{spMspM} dataflows impose different constraints on the preferred representation of input and output matrices. Second, \texttt{spMspM} relies on indirect accesses (through the offsets array) to variable-sized fibers, and requires combining or intersecting those fibers. These operations are inefficient on CPUs and GPUs.

2.2 \texttt{spMspM} Dataflows

Fig. 2 shows the three basic dataflows for \texttt{spMspM}: \textit{inner-product}, \textit{outer-product}, and Gustavson. Fig. 2 also shows the abstract loop nest corresponding to each dataflow (for simplicity, these loop nests assume dense matrices; with compressed sparse matrices, operations are more complex). \texttt{spMspM} computes $C_{M \times N} = A_{M \times K} \times B_{K \times N}$ using a triply-nested loop that iterates over A\textquotesingle s and B\textquotesingle s independent dimensions, M and N, and co-iterates over their shared dimension, K. The dataflow is determined by the level of this co-iteration: in \textit{inner-product}, co-iteration happens at the innermost loop; in \textit{outer-product}, at the outermost loop; and in Gustavson\textquotesingle s, at the middle loop.\footnote{While Fig. 2 shows three loop nest orders, there are six possible orders. The remaining three stem from swapping the M and N loops; this merely switches the dimensions in which inputs are traversed, but results in an otherwise identical dataflow. For example, Fig. 2 shows an inner-product dataflow where A is traversed by rows and B by columns; swapping the outer two loops results in an inner-product dataflow where A is traversed by columns and B by rows.}
for $k$ in $[0, K)$
for $m$ in $[0, M)$
for $n$ in $[0, N)$
$C[m, n] += A[m, k] \times B[k, n]$

Finally, Gustavson has an additional advantage over the other dataflows: its inputs and outputs are all in a consistent format, CSR. By contrast, inner- or outer-product require one input to be in CSR and the other in CSC, to support efficient concordant traversals. We do not evaluate this issue further, but for compound operations (e.g., matrix exponentiation), having different formats requires expensive operand transformations, e.g., converting CSC to CSR, that rival the cost of accelerated \texttt{spMspM} [11].

\subsection{\texttt{spMspM} Accelerators}

Despite the advantages of Gustavson’s algorithm, prior \texttt{spMspM} accelerators have focused on inner- and outer-product dataflows, seeking to maximize reuse of one operand. These designs incorporate different optimizations over the basic dataflow they adopt to mitigate its inefficiencies.

Accelerators like UCNN [20] and SIGMA [43] implement inner-product \texttt{spMspM}. These designs are built around hardware support to accelerate intersections: UCNN traverses compressed sparse data structures, while SIGMA uses a hardware-friendly bitmap-based fiber representation to further accelerate intersections. To counter poor input reuse, some designs also tile input matrices [19] to fit on-chip. While these designs achieve much higher throughput than CPUs and GPUs when matrices are relatively dense (as is typical in e.g., deep learning inference), they suffer from the algorithmic inefficiencies of ineffectual intersections on sparse matrices.

By contrast, accelerators including OuterSPACE [37], SpArch [59], and SCNN [39] implement an outer-product \texttt{spMspM}, and take different approaches to mitigate its inefficiencies. To reduce merge complexity, OuterSPACE divides partial output matrices in rows, then merges rows individually. However, OuterSPACE produces a large amount of off-chip traffic due to partial outputs, which do not fit on-chip. SpArch, by contrast, is built around a very complex high-throughput, high-radix merger that can merge up to 64 partial matrices per pass, and two main techniques to use this merger: pipelining the production of the partial output matrices and their merging to avoid spilling them off-chip, and using a matrix condensing technique that reduces the number and size of partial output matrices. Scaling up SpArch is inefficient because its throughput is bottlenecked by the merger, and scaling up the merger’s throughput incurs \textit{quadratic} area and energy costs. Instead, GAMMA achieves high throughput with linear cost by performing many independent merges in parallel. On highly sparse matrices, SpArch often achieves nearly perfect off-chip traffic because it can produce fewer than 64 partial output matrices; however, on large or less-sparse matrices, SpArch incurs high traffic as it needs to spill many partial outputs off-chip. SpArch’s matrix condensing technique also sacrifices reuse of the $B$ matrix, which can add significant traffic.

Finally, some prior work adopts a hybrid of inner- and outer-product: ExTensor [19] is a flexible accelerator for tensor algebra that combines outer-product at the chip level, and inner-product

\footnote{We use the “stationary terminology from Chen et al. [9].}

\footnote{Or CSC in the alternative Gustavson dataflow; see footnote 2.}
within individual PEs. This approach requires tiling to be used well, and though this hierarchical design eliminates more ineffectual work than a pure inner-product design (by skipping entire ineffective tiles when possible), it still suffers from the drawbacks of the dataflows it adopts.

Despite these optimizations, prior spMxspM accelerators are saddled by the fundamental inefficiencies of the dataflows they adopt. Fig. 3 shows this by comparing the memory traffic of different accelerators when squaring (multiplying by itself) two representative sparse matrices: gupta2 (49 MB, density $1 \times 10^{-3}$), which is relatively dense, and web-Google (58 MB, density $6 \times 10^{-6}$), which is highly sparse. We compare five accelerators with similar hardware budgets (see Sec. 5 for methodology details): (1) IP uses an inner-product dataflow with optimally tiled input matrices; (2) OS is OuterSPACE; (3) S is SpArch; (4) G is Gamma without preprocessing; and (5) GP is Gamma with preprocessing. Each bar shows traffic normalized to compulsory traffic (i.e., the traffic all designs would incur with unbounded on-chip memory, equivalent to reading the inputs and writing the output matrix). Traffic is broken down by data structure: reads of A and B, writes of the final output C, and reads of partial outputs.

Fig. 3 shows that, despite their optimizations, prior accelerators have significant drawbacks: IP works reasonably well on the denser matrix, but is inefficient on the sparser one because of many sparse tiles resulting from the hard-to-predict distribution of nonzeros. OuterSPACE suffers from partial outputs, while SpArch incurs less traffic on partial outputs, but more on matrix B. They both perform well on the sparser matrix, but not on the denser one. Even without preprocessing, Gamma outperforms them all solely by virtue of using Gustavson’s dataflow. But Gamma supports matrix tiling and reordering techniques like prior work, as we will see in Sec. 4. With these preprocessing techniques, Gamma achieves even larger traffic reductions. Finally, since spMsPM is memory-bound, this lower bandwidth translates to higher performance (Sec. 6).

3 Gamma

Fig. 4 shows an overview of Gamma. Gamma consists of multiple processing elements (PEs) that linearly combine sparse fibers; a scheduler that adaptively distributes work across PEs; and a FiberCache that captures irregular reuse of fibers.

Fig. 5 illustrates Gamma’s operation through a simple example that shows how the first few elements of an output row are produced. Gamma always operates on fibers, i.e., streams of nonzero values and their coordinates sorted by coordinate. First, the scheduler fetches row fibers from matrix A and dispatches them to PEs. Each PE then computes a linear combination of row fibers of B to produce a row fiber of output C. For example, in Fig. 5, the scheduler dispatches row A1 to PE 0. Row A1 has only two nonzeros, at coordinates 3 and 5. Therefore, PE 0 linearly combines rows B3 and B5. Fig. 5 shows how the first few elements of each row are combined. First, the B3 and B5 fibers are streamed from the FiberCache. (The FiberCache retains these fibers, so subsequent uses do not incur off-chip traffic.) Then, these fibers are merged into a single fiber, with elements ordered by their shared (column, i.e., N-dimensional) coordinate. Each element in the merged fiber is then scaled by the coefficient of A’s row corresponding to the fiber element’s row (K) coordinate. Finally, consecutive values with the same column (N) coordinate are summed up, producing the output fiber. Fig. 5 shows the values of these intermediate fibers needed to produce the first three elements of output row C1.

Gamma PEs have a bounded radix, R: PEs can linearly combine up to R input fibers in a single pass (though Fig. 5 illustrates the combination of only two fibers, Gamma PEs have a higher radix, 64 in our implementation). When a row of A has more than R nonzeros, the scheduler breaks the linear combination into multiple rounds. For example, with $R = 64$, processing a row of A with 256 nonzeros would be done using four 64-way linear combinations followed by a 4-way linear combination. Each of the initial linear combinations produces a partial output fiber, which is then consumed by the final linear combination. The FiberCache buffers these partial output fibers, avoiding off-chip traffic when possible.

Gamma PEs use high-radix, modest-throughput mergers: PEs have two key design parameters: radix, i.e., how many input fibers they can take; and throughput, i.e., how many input and output elements they can consume and produce per cycle. These parameters are given by the radix and throughput of the PE’s hardware merger, which takes R input fibers and produces a sequence sorted by coordinate (with repeats) as a step in creating a single output fiber from all the elements of all the input fibers. Radix and throughput choices have a substantial impact on PE and system efficiency, and on memory system design, so we discuss them first.

Implementing high-radix merges is cheap, since merger area grows linearly with radix. A high radix in turn makes computation more efficient: it allows many linear combinations to be done...
in a single pass, and increasing the radix reduces the number of merge rounds and partial output fibers needed. For example, linearly combining 4096 fibers with radix-64 PEs would require 65 PE invocations in a depth-2 tree; using radix-2 PEs would require 4095 PE invocations in a depth-12 tree. The radix-64 PEs would produce one set of partial output fibers, whereas the radix-2 PEs would produce 11, increasing FiberCache traffic by about an order of magnitude.

Since higher-radix mergers are larger, there is a tradeoff between the size and power cost of the merger and both PE performance (measured in number of passes required) and FiberCache traffic (due to partial output fibers). With current technology, the sweet spot balancing overall PE cost and performance occurs around $R = 64$.

Another consideration is the throughput of the merger. Implementing high-throughput mergers is costly, since merger area and energy grow quadratically with throughput. Producing $N$ output elements per cycle requires the merger to consume up to $N$ elements from a single input, and to perform up to $N^2$ comparisons. Thus, Gamma uses simple pipelined merge units that produce one output and consume one input per cycle, and achieves high throughput by doing many independent linear combinations in parallel, e.g., by using multiple PEs to process distinct rows of $A$.

This design tradeoff stands in contrast to SpArch [59], the spM-SPM accelerator that comes closest to Gamma’s efficiency. Because SpArch merges partial output matrices rather than fibers, it cannot exploit row-level parallelism, and implements a single high-throughput merger that dominates area and limits throughput. Gamma and SpArch both implement radix-64 mergers. However, while in Gamma each PE’s merger is about the same area as its floating-point multiplier, SpArch spends 38× more area on the merger than on multipliers.

**Gamma’s on-chip storage captures irregular reuse across many fibers:** Although Gamma’s PEs are efficient, the combination of high-radix and many PEs to achieve high throughput means that Gamma’s memory system must support efficient access to a large number of concurrent fibers. For example, a system using 32 radix-64 PEs can fetch 2048 input fibers concurrently. Gamma relies on a novel on-chip storage idiom, FiberCache, to support the irregular reuse patterns of Gustavson’s algorithm efficiently. FiberCache takes two key design decisions: sharing a single structure for all fibers that may have reuse, and combining caching and explicit decoupled data orchestration [40] to avoid large fetch buffers.

Gamma processes four types of fibers: rows of $A$ and $B$, and partial and final output rows of $C$. Rows of $A$ and final output rows of $C$ have no reuse, so they are streamed from/to main memory. Rows of $B$ and partial output rows of $C$ have reuse, but different access patterns: rows of $B$ are read-only and are accessed potentially multiple times (depending on $A$’s nonzeros), whereas partial output fibers, which need to be further merged to produce a final output row, are produced and consumed by PEs, typically within a short period of time. The FiberCache buffers both types of fibers within a single structure, instead of having separate buffers for inputs and outputs. Sharing capacity across fiber types helps because different matrices demand a widely varying share of footprint for partial outputs, but requires careful management to maximize reuse.

FiberCache is organized as a highly banked cache, which allows it to flexibly share its capacity among many fibers or fiber fragments. However, FiberCache is managed using the explicit data orchestration idioms common in accelerators [40]: the fibers needed by each PE are fetched ahead of time, so that when the PE reads each input fiber element, the data is served from the FiberCache. This avoids PE stalls and lets the FiberCache pull double duty as a latency-decoupling buffer. This feature is important because, due to the large number of concurrent fibers processed, implementing such buffering separately would be inefficient: with 32 radix-64 PEs and an 80 ns main memory, implementing these buffers would require about 2 MB of storage, a large fraction of the 3 MB FiberCache we implement (Sec. 5).

### 3.1 Processing Element

Fig. 6 details the design of Gamma’s PE. The PE linearly combines up to $R$ fibers incrementally. Operation begins with a request from the scheduler, which streams up to $R$ input fiber descriptors: for each input, the scheduler specifies its starting location, size, and a scaling factor. If the input fiber is a row of $B$, $B_k$, the scaling factor is value $a_{mk}$; otherwise, the input fiber is a previously generated partial output, and its scaling factor is 1.0. The PE stores scaling factors in a register file, and input fiber locations in the fiber fetcher.

The fiber fetcher then begins streaming input fibers from the FiberCache. The read elements are streamed into two sets of circular buffers: coordinates ($N$) are staged as inputs to the high-radix merger, while values are buffered separately. Each set has $R$ buffers, one for each way of the merger. Since the FiberCache ensures low access latency, these buffers are small and incur low overheads.

![Gamma’s PE architecture.](image)

The merger consumes the minimum coordinate ($N$) among the heads of its $R$ input buffers, and outputs the coordinate together with its way index, i.e., a value between 0 and $R - 1$ that identifies which input fiber this coordinate came from.

The way index is used to read both the corresponding value from the value buffer and the scaling factor. The PE then multiplies these values. Finally, the coordinate and value are processed by an accumulator that buffers and sums up the values of same-coordinate inputs. If the accumulator receives an input with a different coordinate, it emits the currently buffered element, which is part of the output fiber.
Fig. 7 shows the implementation of the merger. The merger is organized as a balanced binary tree of simple compute units. Each unit has an integer comparator for coordinates, and merges coordinate streams incrementally. This design achieves a small area cost, e.g., 55% of a 64-bit floating point multiplier for a radix of 64, and achieves an adequately high frequency.

Unlike prior mergers [45, 59] with throughputs that are high on average but are very sensitive to coordinate distribution, GAMMA’s merger maintains a constant 1-element-per-cycle throughput. Thus, in steady state, the PE consumes one input fiber element per cycle and performs one scaling operation. This achieves high utilization of its most expensive components, the multiplier and the merger.

### 3.2 FiberCache

Fig. 8 shows the FiberCache design and interface. FiberCache builds upon a cache: it has data and tag arrays, organizes data in lines, and uses a replacement policy tailored to fiber access patterns. But FiberCache has two key distinct features. First, FiberCache extends the usual read-write access interface with primitives that manage data movement more explicitly: fetch and consume. Fetch enables decoupled data orchestration by fetching data from memory ahead of execution. Second, to ensure that read’s hit in most cases, FiberCache ensures that fetched data is unlikely to be evicted. This is achieved through the replacement policy. This effectively turns a dynamic portion of FiberCache into buffer-like storage, but without incurring the high overheads of separate, statically sized buffers.

**Reading rows of B** that are not cached incurs a long latency, stalling the PE and hurting performance. FiberCache addresses this issue by decoupling PE data accesses into two steps: fetch and read. A fetch request is sent ahead of execution and places the data into the FiberCache, accessing main memory if needed, and a read request directs the actual data movement from FiberCache to the PE. This decouples the accesses to memory and the computation on PEs.

Unlike speculative prefetching, a fetch is non-speculative: the data accessed by a fetch is guaranteed to have a short reuse distance. FiberCache exploits this property through the replacement policy. FiberCache assigns each line a priority in replacement. The priority is managed as a counter: e.g., a 5-bit counter for 32 PEs. A fetch request increments the priority, while a read request decrements it. Lower-priority lines are selected for eviction. This guarantees that most read’s hit in the cache; effectively, the priority is a soft lock on lines that are about to be used. FiberCache uses simple 2-bit SRRIP [22] to break ties among same-priority lines.

### 3.3 Scheduler

The scheduler assigns compute tasks to PEs to ensure high utilization and minimize memory traffic.

**From A to tasks**: The scheduler assigns work by traversing the rows of A. Each row of A with fewer nonzeros than the PE radix results in a single task that produces the corresponding output row and writes it directly to main memory.

When a row of A has more nonzeros N than the PE radix R, the scheduler produces a task tree that performs an radix-N linear combination in multiple radix-R steps. Fig. 9 shows an example of a task tree that combines 18 fibers using radix-3 mergers. Each node represents a fiber: the root is the output; leaves are rows of B; and intermediate nodes are the partial output fibers. Edges denote which input fibers (children) contribute to a partial or final output fiber (parent).

The scheduler produces a balanced, top-full tree. Balance improves merge efficiency: in the common case, the rows of B have similar nonzeros, so a balanced tree results in similarly sized input fibers at each tree level. This is more efficient than a linear tree, which would build an overlong fiber. Moreover, a balanced tree enables more PEs to work on the same row in parallel. (SpArch [59] uses more sophisticated dynamic selection of merge inputs based on their lengths; this is helpful in SpArch because it purposefully constructs uneven partial output matrices, but does not help in GAMMA.) Top-fullness keeps footprints of partial output fibers low: by keeping the radix of the top levels full, and allowing only the lowest level to have empty input fibers, partial fibers are kept small, reducing the pressure on FiberCache storage.

Figure 9: Example schedule tree (balanced and top-full) to combine 18 input fibers on PEs with radix 3.
Mapping tasks to PEs: The scheduler dynamically maps tasks to PEs: when a PE becomes ready to receive a new task, the scheduler assigns is the next available one. Tasks are prioritized for execution in row order, to produce the output in an ordered fashion. For multi-task rows, the scheduler follows a dataflow (i.e., data-driven) schedule: it schedules as many leaf tasks from a single row as needed to fill PEs, and schedules each higher-level task as soon as its input fibers become available. The scheduler prioritizes higher-level tasks over lower-level ones to reduce the footprint of partial outputs.

Staging tasks and data: To avoid stalls when starting up a linear combination, PEs can accept a new task while processing the existing one. When a PE receives a new task, it starts staging its data into its merge buffers, so that it can switch from processing the old task to the new task in a single cycle.

The main data structure in a scheduler implementation is a scoreboard that buffers tasks not ready to dispatch and monitors partial fibers that have not been produced. Additional logic and buffers are required to fill tasks in the scoreboard by running the outermost loop of Gustavson’s algorithm. The scheduler is 0.4% of total chip area.

3.4 Memory Management

Prior to the execution, matrices $A$ and $B$ are loaded into memory, and a sufficiently wide range of address space is allocated for $C$ and partial output fibers.

Since the lengths of partial output fibers are unknown ahead of time, GAMMA allocates them dynamically. Upon scheduling a merge that produces a partial output fiber, the scheduler estimates the number of nonzeros of the fiber conservatively, by using the sum of the numbers of nonzeros in all its input fibers. The scheduler then assigns and records the address range of the partial output fiber. This space is only used if the FiberCache needs to evict a partial output, a rare occurrence. The scheduler deallocates the memory when the partial output fiber is consumed. The number of partial outputs is limited to twice the number of PEs, so this dynamic memory management requires negligible on-chip memory.

4 PREPROCESSING FOR GAMMA

Though Gustavson is a more efficient dataflow than inner- and outer-product, it can incur high traffic. Consider Gustavson on dense operands: processing each row of $A$ requires a complete traversal of every row of $B$, and results in high memory traffic. This phenomenon is mitigated for sparse operands, because processing a sparse row of $A$ only touches a subset of rows of $B$, and reuse across those subsets makes the FiberCache effective. Specifically, rows of $B$ enjoy reuse in the FiberCache when multiple nonzeros in $A$ with the same column coordinate appear in nearby rows of $A$. However, there are two reasons this may not happen: either nearby rows of $A$ contain largely disjoint sets of column coordinates (the matrix lacks structure), so there is minimal reuse of rows of $B$, or a single row of $A$ has many nonzeros, which requires many rows of $B$, thrashing the FiberCache.

Prior work has addressed improving such problematic memory access patterns in sparse matrices and graphs using preprocessing techniques like tiling and reordering [21, 23, 42]. Similarly, GAMMA, like prior accelerators, can exploit preprocessing tailored to its memory system and dataflow to further reduce data movement.

To improve data reference behavior, we design two preprocessing techniques for rows of $A$. Affinity-based row-reordering targets disparate adjacent rows of $A$ by reordering rows so that similar rows are processed consecutively. Selective coordinate-space tiling breaks (only) dense rows of $A$ into subrows to avoid thrashing, and is applied before row-reordering to extract affinity among the subrows. Both techniques can be implemented by either relying on auxiliary data for indirections or by modifying the memory layout of $A$. These techniques improve the reuse of sets of rows of $B$, achieving better versatility and efficiency.

4.1 Affinity-Based Row Reordering

Problem definition: We use a score function $S(i, j)$ to represent the affinity of two rows $A_i$ and $A_j$. $S(i, j)$ is the number of coordinates for which both $A_i$ and $A_j$ have a nonzero value.

Because on-chip storage can hold rows of $B$ corresponding to several rows of $A$, we are interested in maximizing the affinity of a row with the previous $W$ adjacent rows:

$$\alpha(i) = \sum_{j=max(0, i-W)}^{i-1} S(i, j)$$

We set the window size $W$ to capture the number of rows of $B$ that fit in the FiberCache on average:

$$W = \frac{\text{max nnz in FiberCache}}{\text{nnz per row}_A \cdot \text{nnz per row}_B}$$

The goal of the algorithm is to find a proper permutation of rows to maximize the affinity of the whole matrix, which we call $\alpha$:

$$\alpha = \sum_{i=1}^{M-1} \alpha(i) = \sum_{i=1}^{M-1} \sum_{j=max(0, i-W)}^{i-1} S(i, j)$$

Algorithm: Algorithm 1 shows the pseudocode for the affinity-based row-reordering algorithm. This algorithm is greedy and uses a priority queue ($Q$) to efficiently find the row with highest affinity. The algorithm produces a permutation $P$ of $A$’s rows. This algorithm has complexity $O(RlogR \cdot N^2)$, where $R$ is the number of rows and $N$ is the average number of nonzeros per row, so it scales well to large matrices as long as they are sparse.

Algorithm 1: Affinity-based row reordering.

Result: Permutation $P$ of row indices
for $r \in$ rows do $Q.insert(r, 0)$;
select some $r$ to start, $P[0] \leftarrow r, Q.remove(r)$;
for $i \in [1, M]$ do
  for $u \in$ column coords of row $P[i - 1]$ do
    if $r \in$ row coords of column $u$ do
      if $r \in Q$ then $Q.incKey(r)$;
    if $i > W$ then
      for $u \in$ column coords of row $P[i - W - 1]$ do
        if $r \in$ row coords of column $u$ do
          if $r \in Q$ then $Q.decKey(r)$;
  $P[i] \leftarrow Q.pop()$;
4.2 Selective Coordinate-Space Tiling

Tiling improves input reuse (as each input tile is sized to fit on-chip) at the expense of additional intermediate outputs that must be merged. Tiling dense matrices is nearly always a good tradeoff [8, 38] because each input contributes to many outputs, and tiling introduces a large gain in input locality for a few extra fetches of intermediate outputs. However, this no longer holds with sparse matrices, because output traffic often dominates. In other words, tiling sparse rows may reduce traffic to B but produce many partial output fibers that must be spilled off-chip and then brought back to be merged.

Therefore, we apply tiling selectively, only to extremely dense rows of A. Specifically, we split rows of A whose footprint to hold rows of B is estimated to be above 25% of the FiberCache capacity (the estimated footprint is the length of A’s row times the average number of nonzeros per row of B). Each subrow resulting from this split contributes to a partial output fiber that must be combined eventually. Because these partial output fibers are not accessed close in time, they are likely to be spilled. To ensure that the partial output fibers generated by subrows can be combined in just one round, we use the merger’s radix R as the tiling factor, i.e., the number of subrows. Rather than splitting rows into evenly-sized subrows, we perform coordinate-space tiling [49]: we split evenly in coordinate space, so if column coordinates are in the range [0, K), we create up to R subrows with the ith subrow having the nonzeros within an even subrange [ik/R, (i + 1)K/R). Experimentally, we find this creates subrows with higher affinity, improving performance. In large matrices, the resulting subrows may still be large, so this process is repeated recursively.

5 METHODOLOGY

System: We evaluate a GAMMA system sized to make good use of high-bandwidth memory and consume similar levels of resources compared to prior accelerators [37, 59], in order to make fair comparisons. Our system has 32 radix-64 PEs, a 3 MB FiberCache, and a 128 GB/s High-Bandwidth Memory (HBM) interface. The system runs at 1 GHz. Table 1 details the system’s parameters. We built a cycle-accurate simulator to evaluate GAMMA’s performance and resource utilization.

Table 1: Configuration of the evaluated GAMMA system.

<table>
<thead>
<tr>
<th>Component</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>PEs</td>
<td>32 radix-64 PEs, 1 GHz</td>
</tr>
<tr>
<td>FiberCache</td>
<td>3 MB, 48 banks, 16-way set-associative</td>
</tr>
<tr>
<td>Crossbars</td>
<td>48x48 and 48x16, swizzle-switch based</td>
</tr>
<tr>
<td>Main memory</td>
<td>128 GB/s, 16 64-bit HBM channels, 8 GB/c/channel</td>
</tr>
</tbody>
</table>

We measure GAMMA’s area by writing RTL for the PEs and scheduler. We then synthesize this logic using Synopsys Design Compiler and yosys [55] on the 45 nm FreePDK45 standard cell library [35], with a target frequency of 1GHz at 1.25V. We use CACTI 7.0 [3] to model the FiberCache at 45 nm. We model the same swizzle-switch networks [46] as in prior work [37]. Table 2 shows GAMMA’s area breakdown, which we contrast with prior work in Sec. 6.

Baselines: We compare GAMMA with two state-of-the-art accelerators, OuterSPACE and SpArch. We built detailed memory traffic models for OuterSPACE and SpArch to understand their key operational differences. We use the same approach as prior work [59] to compare end-to-end performance, by using the same set of matrices used in their evaluations. We use the original designs proposed in OuterSPACE and SpArch papers, rather than scaling them to conduct iso-area or iso-power comparisons. This is because the correct scaling strategy for each baseline is unclear. For instance, scaling SpArch requires carefully tuning various buffer and comparator array sizes. As a result, both baselines used in the comparisons have a larger area than GAMMA at the same technology.

Each accelerator uses inputs in the right format for its dataflow (e.g., CSC and CSR inputs for outer-product), and SpArch uses preprocessed inputs as described by Zhang et al. [59]. We use 32-bit integer coordinates and 64-bit, double-precision floating-point

Table 2: Area breakdown of GAMMA (left) and one PE (right).

<table>
<thead>
<tr>
<th>Component</th>
<th>Area (mm²)</th>
<th>PE component</th>
<th>Area (mm²)</th>
<th>% PE</th>
</tr>
</thead>
<tbody>
<tr>
<td>32 PEs</td>
<td>4.8</td>
<td>Merger</td>
<td>0.045</td>
<td>30%</td>
</tr>
<tr>
<td>Scheduler</td>
<td>0.11</td>
<td>FP Mul</td>
<td>0.082</td>
<td>55%</td>
</tr>
<tr>
<td>FiberCache</td>
<td>22.6</td>
<td>FP Add</td>
<td>0.015</td>
<td>10%</td>
</tr>
<tr>
<td>Crossbars</td>
<td>3.1</td>
<td>Others</td>
<td>0.008</td>
<td>5%</td>
</tr>
<tr>
<td>Total</td>
<td>30.6</td>
<td>PE total</td>
<td>0.15</td>
<td>100%</td>
</tr>
</tbody>
</table>

We use 32-bit integer coordinates and 64-bit, double-precision floating-point

Table 3: Characteristics of the common set of matrices (all square).

<table>
<thead>
<tr>
<th>Matrix</th>
<th>Nnz/row</th>
<th>Rows</th>
<th>Matrix</th>
<th>Nnz/row</th>
<th>Rows</th>
<th>Matrix</th>
<th>Nnz/row</th>
<th>Rows</th>
</tr>
</thead>
<tbody>
<tr>
<td>patents_main</td>
<td>2.33</td>
<td>240,547</td>
<td>web-Google</td>
<td>5.57</td>
<td>916,428</td>
<td>2cubes_sphere</td>
<td>16.23</td>
<td>101,492</td>
</tr>
<tr>
<td>p2p-Gnutella31</td>
<td>2.36</td>
<td>62,586</td>
<td>scircuit</td>
<td>5.61</td>
<td>170,998</td>
<td>offshore</td>
<td>16.33</td>
<td>259,789</td>
</tr>
<tr>
<td>roadNet-CA</td>
<td>2.81</td>
<td>1,971,281</td>
<td>amazon0512</td>
<td>7.99</td>
<td>400,727</td>
<td>cop20k_A</td>
<td>21.65</td>
<td>121,192</td>
</tr>
<tr>
<td>webbase-1M</td>
<td>3.11</td>
<td>1,000,005</td>
<td>ca-CondMat</td>
<td>8.08</td>
<td>23,133</td>
<td>filter3D</td>
<td>25.63</td>
<td>106,437</td>
</tr>
<tr>
<td>m133-b3</td>
<td>4.00</td>
<td>200,200</td>
<td>email-Enron</td>
<td>10.02</td>
<td>36,692</td>
<td>poison3Da</td>
<td>26.10</td>
<td>13,514</td>
</tr>
<tr>
<td>cit-Patents</td>
<td>4.38</td>
<td>3,774,768</td>
<td>wiki-Vote</td>
<td>12.50</td>
<td>8,297</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>mario002</td>
<td>5.38</td>
<td>389,874</td>
<td>eage12</td>
<td>15.61</td>
<td>130,228</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 4: Characteristics of the extended set of matrices.

<table>
<thead>
<tr>
<th>Matrix</th>
<th>Nnz/row</th>
<th>Rows</th>
<th>Matrix</th>
<th>Nnz/row</th>
<th>Rows</th>
<th>Matrix</th>
<th>Nnz/row</th>
<th>Rows</th>
<th>Matrix</th>
<th>Nnz/row</th>
<th>Rows</th>
</tr>
</thead>
<tbody>
<tr>
<td>NotreDame_actors</td>
<td>3.75</td>
<td>392,400</td>
<td>gupta2</td>
<td>68.45</td>
<td>62,064</td>
<td>x104</td>
<td>80.4</td>
<td>108,384</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>relat8</td>
<td>3.86</td>
<td>345,668</td>
<td>vsp_bestk30</td>
<td>50.0</td>
<td>56,12</td>
<td>m_t1</td>
<td>99.96</td>
<td>97,578</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Maragal_7</td>
<td>25.63</td>
<td>46,845</td>
<td>Ge87H76</td>
<td>69.85</td>
<td>112,985</td>
<td>ship_001</td>
<td>11.58</td>
<td>34,920</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>degen3</td>
<td>43.81</td>
<td>185,501</td>
<td>raefsky3</td>
<td>70.22</td>
<td>21,200</td>
<td>msc10848</td>
<td>113.36</td>
<td>10,848</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>EternityII_Etilde</td>
<td>116.42</td>
<td>10,054</td>
<td>sme3D8</td>
<td>71.6</td>
<td>29,067</td>
<td>opt1</td>
<td>124.97</td>
<td>15,449</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>nemsemm1</td>
<td>267.17</td>
<td>3,945</td>
<td>Ge99H100</td>
<td>74.8</td>
<td>112,985</td>
<td>ramege02</td>
<td>170.31</td>
<td>16,830</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
values. Because the outer-product baselines and Gamma always consume coordinates and values, we store them together, as shown in Fig. 1. For the memory traffic comparison in Fig. 3, the inner-product accelerator (IP) uses separate coordinate and value arrays, and values are only fetched on a matching intersection, since this reduces traffic.

We also compare Gamma against the spMGeneral implementation from Intel MKL [52] (mkl_sparse_spmm function), running on a 4-core, 8-thread Skylake Xeon E3-1240 v5, with two DDR4-2400 channels. We do not include GPU results because existing GPU spMGeneral implementations perform similarly to MKL on CPUs [59].

Inputs: We use two sets of matrices. First, the Common set of matrices is the set used in the evaluations of OuterSPACE and SpArch, as shown in Table 3. We use the Common set for direct performance comparisons with these accelerators. However, the Common set covers only a fraction of the space of possible inputs: these matrices are square, and most are very sparse, with a maximum mean of 26 nonzeros per row. This is not representative of other commonly used matrices, and masks the inefficiencies of outer-product designs. To evaluate the designs with a broader range of inputs, we construct the Extended set of matrices, which includes 18 matrices from the SuiteSparse Matrix Collection [30]. Table 4 lists these matrices, which include non-square and square matrices with a wider range of sparsities and sizes. We evaluate $A \times A$ for square matrices (like prior work), and $A \times A^T$ for non-square matrices.

6 EVALUATION

6.1 Performance on Common-Set Matrices

Fig. 10 reports the performance of all accelerators on common-set matrices. Each bar shows the gmean speedup over our software baseline, MKL. Note that common-set matrices are highly sparse and thus well suited for OuterSPACE and SpArch. On these matrices, Gamma (with preprocessing) is gmean 2.1× faster than SpArch, 7.7× faster than OuterSPACE, and 38× faster than MKL. Even without preprocessing, which makes Gamma gmean 16% faster, Gamma outperforms SpArch by 1.84×, OuterSPACE by 6.6×, and MKL by 33×.

Fig. 11 further shows the per-matrix speedups of Gamma (with preprocessing) over MKL. Gamma outperforms MKL by up to 184×.

Fig. 12 and Fig. 13 explain how Gamma outperforms SpArch and OuterSPACE: through a combination of reducing memory traffic and improving memory bandwidth utilization.

Fig. 12 reports the memory traffic of OuterSPACE, SpArch, and Gamma without and with preprocessing. Each group of bars shows results for one matrix. Traffic is normalized to the compulsory traffic, which would be incurred with unbounded on-chip storage:

fetching $A$, the needed rows of $B$, and writing $C$. Each bar is broken down into four categories: reads of $A$ or $B$, writes of $C$, and reads and writes of partial outputs.

Fig. 12 shows that Gamma incurs close-to-optimal traffic: across all inputs, it is only 7% higher than the compulsory (i.e., minimum) traffic with preprocessing, and 26% higher without preprocessing. By contrast, SpArch is 59% higher, and OuterSPACE is 4× higher. OUTERSPACE suffers reads and writes to partial matrices. SpArch reduces partial output traffic over OuterSPACE, but incurs high traffic on $B$ for two reasons. First, to reduce partial output traffic, SpArch preprocesses $A$ to produce a schedule that worsens the access pattern to $B$. Second, SpArch splits its storage resources across data types (e.g., merge and prefetch buffers), leaving only part of its on-chip storage (around half a megabyte) to exploit reuse of $B$. By contrast, Gamma’s shared FiberCache allows $B$’s rows to use more on-chip storage when beneficial. Because Gamma’s partial outputs are rows, it has negligible partial output traffic, and its main overhead comes from imperfect reuse of $B$.

Fig. 13 further illustrates how memory bandwidth translates to performance. Because Gamma’s PEs achieve very high throughput (processing inputs and outputs at a peak rate of 768 GB/s) and Gustavson’s algorithm does not have compute-bound execution phases, Gamma almost always saturates the available 128 GB/s memory bandwidth. By contrast, OuterSPACE and SpArch suffer from the compute bottleneck of merging all the partial matrices, and hence achieve lower bandwidth utilizations of 48.3% and 68.6%, respectively, on the same matrices. Gamma’s higher performance stems from its lower memory traffic and higher bandwidth utilization.

To further illustrate how FiberCache is utilized, for each application, we sample the utilization of FiberCache every 10,000 cycles. Fig. 14 shows the average utilization of FiberCache. On these matrices, $B$ fibers are dominant in FiberCache, while partial result fibers consume non-negligible capacity on some inputs, including wiki-Vote, email-Enron, and webbase-1M.

6.2 Performance on Extended-Set Matrices

To further evaluate the versatility of Gamma, we use the extended set of matrices, which includes non-square matrices and square matrices more diverse than the common set (Sec. 5). Fig. 15 shows the speedups of Gamma (with preprocessing) over MKL. By exploiting hardware specialization, Gamma outperforms MKL by gmean 17× and by up to 50×.
explained, A B C Partial outputs

Fig. 12: Off-chip traffic on common-set matrices of OuterSPACE (O), SpArch(S), and GAMMA without and with preprocessing (G/GP) (lower is better).

Fig. 13: Memory bandwidth utilization on common-set matrices of GAMMA without and with preprocessing (G/GP).

Fig. 14: Cache utilization on common-set matrices of GAMMA without and with preprocessing (G/GP).

6.3 Effectiveness of GAMMA Preprocessing

Preprocessing improves the performance of GAMMA by 18% on average. Fig. 19 further illustrates the effects of affinity-based row reordering and selective coordinate-space tiling in two cases. Affinity-based row reordering improves the reuse of B. For instance, it contributes to a 6x reduction of traffic on sme3db. As Sec. 4.2 explained, tiling all rows of A (T in Fig. 19) may hurt: it does little harm to Maragal_7 but causes 13x extra traffic on sme3db due to excessive partial outputs. This is why GAMMA selectively tiles long rows only. Selective coordinate-space tiling reduces traffic of B drastically by

Fig. 15: Speedups of GAMMA with preprocessing over MKL on extended-set matrices.
tiling dense rows (e.g., on Maragal_7), and also avoids performance pathologies by not tiling sparse rows (e.g., on sme3Db).

Preprocessing takes an average time of 44 seconds and 208 seconds on the common-set matrices and the extended-set matrices, respectively. On average, the preprocessing time for a matrix is 4600X longer than using Gamma to execute spMpsM on the same matrix. Thus, preprocessing is beneficial only when the A matrix will be reused frequently.

6.4 Gamma Scheduling
Gamma’s scheduling algorithm (Sec. 3.3) uses multiple PEs to process the tasks produced by the same row of A (or, if preprocessing tiles the row, the same subrow of A). To demonstrate its effectiveness, we compare it against a less dynamic algorithm that always uses a single PE to process all the tasks for each row of A. Fig. 20 shows the off-chip memory traffic on input matrix email-Enron. With the single-PE approach, all the tasks from the same row are serialized, so partial result fibers stay resident in FiberCache for a longer time. By contrast, Gamma’s multi-PE algorithm allows partial result fibers to be consumed as early as possible. On email-Enron, this multi-PE scheduling algorithm reduces memory traffic by 18%, and hence improves performance by 17%.

6.5 Gamma Roofline Analysis
To show that Gamma uses resources well, Fig. 21 presents its roofline analysis plot. The plot presents arithmetic intensity (x-axis) versus Memory traffic (y-axis). The lines represent different scheduling algorithms: using Multiple PEs (the default) or a Single PE for each row.
Figure 21: Performance of GAMMA without and with preprocessing (G/GP) in a roofline model.

in FLOPs per byte of off-chip memory traffic, and performance (y-axis) in GFLOPs (as is usual, one multiply-accumulate is counted as a single FLOP, despite being performed by a separate multiplier and adder in GAMMA PEs). Note that the plot uses a logarithmic scale for both axes. Each dot represents a single matrix; results without preprocessing are shown in blue, while results with preprocessing are shown in yellow. The plot also shows the design’s roofline at 32 GFLOPs, which caps the maximum achievable performance: the sloped (left) part of the roofline is set by memory bandwidth, while the flat (right) part is set by compute (PE) throughput.

Fig. 21 shows that most matrices have low arithmetic intensity and are memory bandwidth-bound, while some have higher arithmetic intensity and are compute-bound. More importantly, this shows that GAMMA uses its resources well: almost all matrices are right at or very close to the roofline, showing that the system is driven to saturation all the time. Only three matrices are noticeably below the roofline (gupta2, Ge87H76, and Ge99H100). By inspection, we have found that these matrices have memory-bound and compute-bound phases, so while their average compute intensity falls past the sloped part of the roofline, they do not saturate PEs all the time due to memory-bound phases. Nonetheless, compared to prior accelerators, which have memory-bound and compute-bound phases (e.g., partial output matrix generation vs. merging in OuterSPACE and SpArch), this result shows that Gustavson’s algorithm yields a more consistent behavior that uses resources better.

6.6 GAMMA Area Analysis

As shown in Table 2, the total area of GAMMA is 30.6 mm², synthesized with a 45 nm standard cell library. Scaled down to 40 nm, GAMMA’s area is 24.2 mm², smaller than the 28.5 mm² of SpArch at 40 nm and the 87 mm² of OuterSPACE at 32 nm. The vast majority of area is used by the FiberCache. This is a good tradeoff for SpMSPM, since the key bottleneck is memory traffic and data movement. The PEs are simple, taking 16% of chip area, and the merger and multiplier are its main components. By contrast, SpArch and OuterSPACE spend far more area on compute resources, e.g., 60% on SpArch’s merger.

6.7 Scalability Studies

Fig. 22 and 23 show GAMMA’s performance and traffic on common-set and extended-set matrices, respectively, when the number of PEs is swept from 8 to 128 (the default is 32 PEs). For common-set matrices, 32 PEs are the right tradeoff, as all are memory-bound at 32 PEs. Since some extended-set matrices have higher reuse
and thus arithmetic intensity, GAMMA continues to improve performance past 32 PEs: at 128 PEs (which would increase accelerator area by about 50%), GAMMA is gmean 65% faster than at 32 PEs.

Fig. 24 and Fig. 25 show GAMMA’s performance and traffic on common-set and extended-set matrices, respectively, when FIBERCACHE size is swept from 0.75 MB to 12 MB (the default is 3 MB). At and after 1.5 MB, performance improves smoothly with FIBERCACHE size, showing that GAMMA can leverage additional storage to gracefully improve performance on inputs where non-compulsory traffic is high. However, performance is significantly degraded at 0.75 MB. This performance cliff occurs because FIBERCACHE is used as decoupling buffers, and at this size, there is little capacity left to capture irregular reuse. These results show that FIBERCACHE does indeed save significant storage on dedicated buffers.

7 ADDITIONAL RELATED WORK

Much prior work has proposed optimized CPU and GPU implementations for spMspM, e.g., using autotuning [51], input characteristics [56], or code generation [29] to pick a well-performing spMspM implementation. Intel’s MKL [52], which we use in our evaluation, is generally the fastest, or close to the fastest, across input matrices [56]. Although GPUs have higher compute and memory bandwidth than CPUs, spMspM is a poor match to the regular data parallelism supported in current GPUs, so GPU frameworks [13, 32, 36] achieve similar spMspM performance to CPUs [56, 59].

Most CPU and GPU implementations follow Gustavson’s dataflow; variants differ in how they merge rows of B, e.g., using sparse accumulators [15, 28], bitmaps [24], unordered associative containers [33, 34, 36], trees [47], or heaps [1] to hold outputs. This algorithmic diversity arises because merging fibers is an expensive operation in general-purpose architectures. At a high level, heaps are space-efficient but slow, and the other data structures trade lower compute for higher space costs. GAMMA’s high-radix merges are both space-efficient and make merges very cheap, avoiding this dichotomy.

As explained in Sec. 2.3, to the best of our knowledge, accelerators earlier than GAMMA did not exploit Gustavson’s dataflow. However, MatRaptor [48], which is concurrent with GAMMA, does exploit Gustavson’s dataflow. Nonetheless, MatRaptor and GAMMA are very different. MatRaptor does not exploit the reuse of B fibers: it streams such fibers from DRAM and uses them once. By contrast, GAMMA exploits the reuse of B fibers with FIBERCACHE. This adds area costs, but since reusing B fibers is the key way by which Gustavson’s dataflow minimizes traffic, GAMMA improves performance significantly. Consequently, on the common-set matrices, MatRaptor outperforms OUTERSPACE by only 1.8× [48], worse than SpArch’s improvement over OUTERSPACE (3.6×), while GAMMA outperforms OUTERSPACE by 6.6× even without preprocessing.

Preprocessing of sparse matrices [10, 12, 14, 53] has been studied extensively on CPUs and GPUs. Matrix preprocessing on CPUs and GPUs typically targets creating dense tiles [42] to reduce irregularity of partial outputs, disjoint tiles [4] to minimize communication, or balanced tiles [21, 23] to ease load balancing. These techniques differ from GAMMA’s: our goal is to improve the locality of B, whereas CPUs and GPUs lack high-radix mergers and have more on-chip storage, making B’s locality a less pressing concern.

To classify on-chip storage structures, we can use the two-dimensional taxonomy from Pellauer et al. [40]. Specifically, the content of an on-chip storage structure can be managed in two styles: explicit or implicit. Explicitly orchestrated structures allow applications to directly control what to retain or remove, while implicitly orchestrated structures infer such decisions implicitly based on read/write accesses. A storage structure can be used in either coupled or decoupled manner depending on whether the data needed is pre-staged ahead of processing to hide the memory access latency. Caches are implicit and coupled. GAMMA’s FIBERCACHE combines features of caches and explicitly managed buffers to both exploit irregular reuse and hide memory latency through explicit decoupled data orchestration. Stash [31] is also a hybrid of caches and scratchpads, but with different goals: Stash maps data regions and accesses them explicitly, with a scratchpad interface, to reduce addressing power. Stash fetches accessed data lazily, which saves traffic when not all mapped data is accessed, but leaves accesses coupled to users. By contrast, GAMMA knows precisely which data will be accessed so its decoupled design hides long access latency. Following the taxonomy above, Stash is explicit and coupled, whereas FIBERCACHE is implicit and decoupled.

Finally, while we focus on spMspM, many applications use high-dimensional tensors. For instance, TACO [28, 29] introduces workspace and proposes compiler machinery to handle complex tensor operations. GAMMA can be combined with such techniques to support a broader range of applications.

8 CONCLUSION

spMspM is the basic building block of many emerging sparse applications, so it is crucial to accelerate it. However, prior spMspM accelerators use inefficient inner- and outer-product dataflows, and miss Gustavson’s more efficient dataflow. We have presented GAMMA, an spMspM accelerator that leverages Gustavson’s algorithm. GAMMA uses dynamically scheduled PEs with efficient high-radix mergers and performs many merges in parallel to achieve high throughput, reducing merger area by about 15× over prior work [59]. GAMMA uses a novel on-chip storage structure, FIBERCACHE, which supports Gustavson’s irregular reuse patterns and streams thousands of concurrent sparse fibers with explicitly decoupled data movement. We also devise new preprocessing algorithms that boost GAMMA’s efficiency and versatility. As a result, GAMMA outperforms prior accelerators by gmean 2.1×, and reduces memory traffic by 2.2× on average and by up to 13×.

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