Composing Loop-carried Dependence with Other Loops

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Abstract
Sparse fusion is a compile-time loop transformation and runtime scheduling implemented as a domain-specific code generator. Sparse fusion generates efficient parallel code for the combination of two sparse matrix kernels where at least one of the kernels has loop-carried dependencies. Available implementations optimize individual sparse kernels. When optimized separately, the irregular dependence patterns of sparse kernels create synchronization overheads and load imbalance, and their irregular memory access patterns result in inefficient cache usage, which reduces parallel efficiency. Sparse fusion uses a novel inspection strategy with code transformations to generate parallel fused code for sparse kernel combinations that is optimized for data locality and load balance. Code generated by Sparse fusion outperforms the existing implementations ParSy and MKL on average 1.6× and 5.1× respectively and outperforms the LBC and DAGP coarsening strategies applied to a fused data dependence graph on average 5.1× and 7.2× respectively for various kernel combinations.

1 Introduction
Numerical algorithms [41] and optimization methods [5, 9, 46] are typically composed of numerous consecutive sparse matrix computations. For example, in iterative solvers [41] such as Krylov methods [10, 42], sparse kernels that apply a preconditioner or update the residual are repeatedly executed inside and between iterations of the solver. Sparse kernels with loop-carried dependencies, i.e. kernels with partial parallelism, are frequently used in numerical algorithms, and the performance of scientific simulations relies heavily on efficient parallel implementations of these computations. Sparse kernels that exhibit partial parallelism often have multiple wavefronts of parallel computation where a synchronization is required for each wavefront, i.e. wavefront parallelism [16, 54]. The amount of parallelism varies per wavefront and often tapers off towards the end of the computation, which results in load imbalance. Figure 1 shows with dark lines the nonuniform parallelism for the sparse incomplete Cholesky (SpIC0) and the sparse triangular solve (SpTRSV) kernels when SpTRSV executes after SpIC0 completes. Separately optimizing such kernels exacerbates this problem by adding even more synchronization. Also, opportunities for data reuse between two sparse computations might not be realized when sparse kernels are optimized separately.

Instead of scheduling iterations of sparse kernels separately, they can be scheduled jointly. Wavefront parallelism can be applied to the joint DAG of two sparse computations. A data flow directed acyclic graph (DAG) describes dependencies between iterations of a kernel [7, 19, 50]. A joint DAG includes all of the dependencies between iterations within and across kernels. The joint DAG of sparse kernels with partial parallelism with the DAG of another sparse kernel provides slightly more parallelism per wavefront without increasing the number of wavefronts. The yellow line in Figure 1 shows how scheduling the joint DAG of SpIC0 and SpTRSV provides more parallelism per wavefront and significantly reduces the number of wavefronts (synchronizations). However, the load balance issues remain, and there are still several synchronizations.

Wavefronts of the joint DAG can be aggregated to reduce the number of synchronizations. DAG partitioners such as Load-Balanced Level Coarsening (LBC) [8] and DAGP [20] apply aggregation, however, when applied to the joint DAG because they aggregate iterations from consecutive wavefronts, load imbalance might still occur. Also, by aggregating iterations from wavefronts in the joint DAG, DAG partitioning methods potentially improve the temporal locality between the two kernels but, this can disturb spatial locality within each kernel. For example, for two sparse kernels that only share a small array and operate on different sparse matrices, optimizing temporal locality between kernels will not
be profitable. Finally, even when applied to the DAG of an individual kernel, DAGP and LBC are slow for large DAGs because of the overheads of coarsening [20]. This problem exacerbates when applied to the joint DAG because the joint DAG is typically 2-4x larger than an individual kernel’s DAG.

We present sparse fusion that creates an efficient schedule and fused code for when a sparse kernel with loop-carried dependencies is combined with another sparse kernel. Sparse fusion uses an inspector to apply a novel Multi-Sparse DAG Partitioning (MSP) runtime scheduling algorithm on the DAGs of the two input sparse kernels. MSP uses a vertex dispersion strategy to balance workloads in the fused schedule, uses two novel iteration packing heuristics to improve the data locality due to spatial and temporal locality of the merged computations, and uses vertex pairing strategies to aggregate iterations without joining the DAGs.

Figure 2 compares the schedule created by sparse fusion (sparse fusion schedule) with the schedules created by applying LBC to the individual DAGs of each sparse kernels (LBC unfused schedule) and LBC applied to the joint DAG (LBC joint DAG schedule). All approaches take the input DAGs in Figure 2b. Solid purple vertices are the DAG of sparse triangular solve (SpTRSV) and the dash-dotted yellow correspond to Sparse Matrix-Vector multiplication (SpMV). LBC is a DAG partitioner that partitions a DAG into a set of aggregated wavefronts called s-partitions that run sequentially, each s-partition is composed of some independent w-partitions. In the LBC unfused schedule in Figure 2c, LBC is used to partition the SpTRSV DAG and will create two s-partitions, i.e. s1 and s2. The vertices of SpMV are scheduled to run in parallel in a separate wavefront s3. This implementation is not load balanced because the number of partitions that can run in parallel differs for each s-partition. In the LBC joint DAG, the DAGs are first joint using the dependency information between the two kernels shown with blue dotted arrows and then LBC is applied to create the two s-partitions in Figure 2d. These s-partitions are also not load balanced, for example s2 only has one partition. Sparse fusion uses MSP to first partition the SpTRSV DAG and then disperses the SpMV iterations to create load-balanced s-partitions, e.g. the two s-partitions in Figure 2e have three closely balanced partitions.

SpTRSV solves \( Lx = b \) to find \( x \) and SpMV performs \( y = Ax \) where \( L \) is a sparse lower triangular matrix, \( A \) is a sparse matrix, and \( x, b, \) and \( y \) are vectors. The LBC joint DAG schedule interleaves iterations of two kernels to reuse \( x \). However, this can disturb spatial locality within each kernel because the shared data between the two kernels, \( x \), is smaller than the amount of data used within each kernel, \( A \) and \( L \). With the help of a reuse metric, Sparse fusion realizes the larger data accesses inside each kernel and hence packs iterations to improve spatial locality within each kernel.

We implement sparse fusion as an embedded domain-specific language in C++ that takes the specifications of the sparse kernels as input, inspects the code of the two kernels, and transforms code to generate an efficient and correct parallel fused code. The primary focus of sparse fusion is to fuse two sparse kernels where at least one of the kernels has loop-carried dependence. Sparse fusion is tested on seven of the most commonly used sparse kernel combinations in scientific codes which include kernels such as sparse triangular solver, incomplete Cholesky, incomplete LU, diagonal scaling, and matrix-vector multiplication. The generated code is evaluated against MKL and ParSy with average speedups of 5.1x and 1.6x respectively. Sparse fusion compared to fused implementations of LBC, DAGP, and wavefront techniques applied to the joint DAG provides on average 5.1x, 7.2x and 2.5x speedup respectively.
Sparse Fusion

Sparse fusion is implemented as a code generator with an inspector-executor technique that can be used as a library. It takes the input specification shown in Figure 3a and generates the inspector and the executor in Figure 3b. The inspector includes the MSP algorithm and functions that generate its inputs, i.e., dependency DAGs, reuse ratio, and the dependency matrix. The executor is the fused code that is created by the fused transformation.

2.1 Code Generation

Sparse fusion is implemented as an embedded domain-specific language. It takes as input the specification shown in Figure 3a and generates the driver code in Figure 3b. At compile-time, the data types and kernels in Figure 3a are converted to an initial Abstract Syntax Tree (AST) using TM.gen_c() in line 14. Lines 11 and lines 12 in Figure 3a demonstrate how the user specifies the two kernels for the running example in Figure 2 as inputs to Sparse fusion. The corresponding AST for the example is shown in Figure 2a.

At runtime by running the driver code in Figure 3b, the inspector creates a fused schedule, and the executor runs the fused schedule. The inspector first builds inputs to MSP using functions intra_DAG, inter_DAG, and compute_reuse in lines 6–10 in Figure 3b and then calls MSP in line 11 to generate FusedSchedule for r threads. Then the executor code, fused_code in line 14 in Figure 3b, runs in parallel using the fused schedule.

2.2 The Inspector in Sparse Fusion

The MSP algorithm requires kernel-specific inputs. Its inputs are the dependency matrix between kernels, the DAG of each kernel, a reuse ratio. Sparse fusion analyzes the kernel code, available from its AST, to generate inspector components that create these inputs.

Dependency DAGs: Lines 6–7 in Figure 3b use an internal domain-specific library to generate the dependency DAG of each kernel. General approaches such as work by Mohammadi et al. [33] can also be used to generate the DAGs, however, that will lead to higher inspection times compared to a domain-specific approach. For example, with domain knowledge, sparse fusion will use the L matrix as the SpTRSV DAG $G_i$ in Figure 2b. Each nonzero $L_{ij}$ represents a dependency from iteration $i$ to $j$.

Dependency Matrix $F$: MSP uses the dependency information between kernels to create a correct fused schedule. By running the inter_DAG function, sparse fusion creates this information and stores it in matrix $F$. To generate inter_DAG, sparse fusion finds dependencies between statements of the two kernels by analyzing the AST. Each nonzero $F_{ij}$ represents a dependency from iteration $j$ of the first loop, i.e. column $j$ of $F$, to iteration $i$ of the second loop, i.e. row $i$ of $F$. In Figure 2a, there exists a read after write (flow) dependency between statements $x[i1]$ in line 5 and $x[j1]$ in line 13. As a result, sparse fusion generates the function shown in Listing 1. The resulting $F$ matrix, generated at runtime, is shown in Figure 2b.

```c
for(i1=0; i1<n; i1++){
    j1 = i1;
    if(A.p[j1] < A.p[j1+1])
        F[j1].append(i1);
}
```

Listing 1. inter_DAG function for the example in Figure 2a.

Reuse Ratio: MSP uses a reuse ratio based on the memory access patterns of the kernels to decide whether to improve locality within each kernel or between the kernels. The inspector in line 9 in Figure 3b computes the reuse ratio metric. The metric represents the ratio of common to total memory accesses of the two kernels, i.e., $rac{\text{common memory access}}{\max(\text{kernels1 accesses, kernels2 accesses})}$. For a reuse ratio larger than one, the number of common memory accesses between the two kernels is larger than the accesses inside a kernel. Sparse fusion estimates memory accesses using the ratio of the size of common variables over the maximum of the total size of variables amongst the kernels. For the running example, the code generated for compute_reuse is $2*x.n / \max(A.size+x.n+y.n,L.size+x.n+b.n)$. Since $x$ is smaller than $L$ or $A$, the reuse ratio is less than one.

2.3 Fused Code

To generate the fused code, a fused transformation is applied to the initial AST at compile-time and two variants of the fused code are generated, shown in Figure 4. The transformation variants are separated and interleaved. The fused code uses the reuse ratio at runtime to select the correct variant for the specific input. The variable fusion in line 1 of Figure 4b and 4c is set to False if MSP determines fusion is not profitable. Figure 4a shows the sequential loops in the AST, which are annotated with Fuse, and are transformed to the separated and interleaved code variants as shown in order in Figures 4b and 4c. The separated variant is selected when
the reuse ratio is smaller than one. In this variant, iterations of one of the loops run consecutively without checking the loop type. The interleaved variant is chosen when the reuse ratio is larger than one. In this variant, iterations of both loops should run interleaved, and the variant checks the loop type per iteration as shown in lines 6 and 10 in Figure 4c.

### 3 Multi-Sparse DAG Partitioning

Sparse fusion uses the multi-sparse DAG partitioning (MSP) algorithm to create an efficient fused partitioning that will be used to schedule iterations of the fused code. MSP partitions vertices of the DAGs of the two input kernels to create parallel load-balanced workloads for all cores while improving locality within each thread. This section describes the inputs, output, and three steps of the MSP algorithm using the running example in Figures 2 and 5.

#### 3.1 Inputs and Output to MSP

The inputs to MSP (shown in Algorithm 1) are two DAGs \( G_1 \) and \( G_2 \) from in order lexicographically first and second input kernels, and the inter-DAG dependency matrix \( F \) that stores the dependencies between kernels. A DAG shown with \( G_j(V_j, E_j, c) \) has a vertex set \( V_j \) and an edge set \( E_j \) and a non-negative integer weight \( c(v_i) \) for each vertex \( v_i \in V_j \). The vertex \( v_i \) of \( G_j \) represents iteration \( i \) of a kernel and each edge shows a dependency between two iterations of a kernel. \( c(v_i) \) is the computational load of a vertex and is defined as the total number of nonzeros touched to complete its computation. Because sparse matrix computations are generally memory bandwidth-bound, \( c(v_i) \) is a good metric to evaluate load balance in the algorithm [8]. \( F \) is stored in the compressed sparse row (CSR) format and \( F_1 \) is used to extract the set of vertices in \( G_1 \) that \( v_i \in V_1 \) depends on. Other inputs to the algorithm are the number of requested partitions \( r \), which is set to the number of cores, and the reuse ratio discussed in section 2.2.

The output of MSP is a *fused partitioning* \( \mathcal{V} \) that has \( b \geq 1 \) s-partitions, each s-partition contains up to \( k \geq 1 \) w-partitions, where \( k \leq r \). MSP creates \( b \) disjoint s-partitions from vertices of both DAGs, shown with \( \mathcal{V}_s \) where \( \bigcup_{b=1}^{b} \mathcal{V}_s = \mathcal{V}_1 \cup \mathcal{V}_2 \). Each s-partition includes vertices from a lower bound and upper bound of wavefront numbers shown with \( s_i = [l_i, ub_i] \) as well as some slack variables. For each s-partition \( \mathcal{V}_s \), MSP creates \( m_i \leq k \) independent w-partitions \( \mathcal{V}_{s_i w} \) where \( \bigcup_{i=1}^{b} \mathcal{V}_{s_i w} = \mathcal{V}_s \). Since w-partitions are independent, they can run in parallel.

**Example.** In Figure 2b, the SpTRSV DAG \( G_1 \), the SpMV DAG \( G_2 \), the inter-DAG dependency matrix \( F \) are inputs to MSP. Other inputs to MSP are \( r=3 \) and the *reuse ratio*. The fused partitioning shown in Figure 2e has two s-partitions (\( b=2 \)). The first s-partition has three w-partitions (\( m_1=3 \)) shown with \( \mathcal{V}_{s_1} = \{ [1, 2, 3, 4] : [5, 6, 5, 6, 7, 8, 9, 9] \} \), the underscored vertices belong to \( G_1 \).

#### 3.2 The MSP Algorithm

Algorithm 1 shows the MSP algorithm. It takes the inputs and goes through three steps of (1) vertex partitioning and partition pairing with the objective to aggregate iterations without joining the DAGs of the inputs kernels; (2) merging and slack vertex assignment to reduce synchronization and to balance workloads; and (3) packing to improve locality.

**3.2.1 Vertex Partitioning and Partition Pairing.** The first step of MSP partitions one of the input DAGs \( G_1 \) or \( G_2 \), and then uses that partitioning to partition the other DAG. The created partitions are stored in \( \mathcal{V} \). Partitioning the joint DAG is complex and might not be efficient because of the significantly larger number of edges and vertices added compared to the individual DAG of each kernel. Instead, MSP ignores the dependencies across kernels and first creates a partitioning from one of the DAGs with the help of *vertex partitioning*. Then the other DAG is partitioned using a *partition pairing* strategy. The DAG that is partitioned first is the head DAG and the other is the tail DAG. A *head DAG choice strategy* is used to select the head DAG.
Vertex partitioning. MSP uses the LBC DAG partitioner [8] to construct a partitioning of the head DAG in lines 2 and 11 of Algorithm 1 by calling the function LBC. The resulting partitioning has a set of disjoint s-partitions. Each s-partition contains k disjoint w-partitions which are balanced using vertex weights. Disjoint w-partitions ensure all w-partitions within s-partitions are independent. The created partitions are stored in a two-dimensional list H using 1st, e.g., w-partition w_j of s-partition s_i is stored in H_{i,j}.

Partition pairing. The algorithm then partitions the tail DAG with forward pairing, if G_1 is the head DAG, or with backward pairing, if G_2 is the head DAG. With the pairing strategy, some of the partitions of the tail DAG are paired with the head DAG partitions. Pair-partitions are self-contained so that they execute in parallel if assigned to the same s-partition. The created partitions are put in the fused partitioning V to be used in step two. The first algorithm first describes the condition for partitions to be self-contained and then explains the forward and backward pairing strategies. Pair partitions H_{i,j} and T_{i,j} are called self-contained if all reachable vertices from a breadth-first search (BFS) on ∀v ∈ H_{i,j} ∪ T_{i,j} through vertices of G_1 and G_2 are in H_{i,j} ∪ T_{i,j}. Self-contained pair partition (H_{ip}, T_{ip}) and pair partition (H_{iq}, T_{iq}) can execute in parallel without synchronization if in the same wavefront i, i.e., ∀1 ≤ i ≤ b ∧ (1 ≤ p, q ≤ m_i). Partitions that do not satisfy this condition create synchronizations in the final schedule.

The backward pairing strategy visits every partition H_{i,j} and performs a BFS (line 5) from vertex v_i ∈ H_{i,j} to its dependent vertices in G_1 which are reachable through F_1. Reachable vertices are stored in T_{i,j}. The partitions in H and T are assigned a w- and s-partition and are then put into the fused partitioning V (via add in line 6). The assigned s- and w-partitions for H_{i,j} are s_{i+1} and w_j respectively, i.e., V_{s_{i+1},w_j}. T_{i,j} should be executed before H_{i,j} thus is placed in s-partition S or V_{s_{i+1},w_{m+1}}, where m_i is number of w-partitions in V_{s_i} at this point. If a vertex in H_{i,j} depends on more than one vertex in G_1, some vertices are replicated in different T partitions. While replication leads to redundant computation, it ensures that the pair partition (H_{i,j}, T_{i,j}) is self-contained because vertices that depend on the vertices in H_{i,j} will be included in T_{i,j}. MSP performs fusion only if profitable, hence fusion is disabled (by setting fusion to False) if the number of redundant computations go beyond a threshold. This threshold is 2 × (|V_1| + |V_2|) in line 9 and is defined as the sum of vertices of both DAGs.

The forward pairing strategy iterates over every partition H_{i,j} and performs a BFS from vertex v_i ∈ H_{i,j} to its reachable vertices in G_2 through F_2^T, see lines 12–18 in Algorithm 1. The list of reachable vertices are stored in T_{i,j} via BFS in line 14. If a vertex v_m in T_{i,j} depends on vertex v_j in G_1 and v_j does not exist in H_{i,j} then v_m should be removed to ensure (H_{i,j}, T_{i,j}) is self contained. The remove_uncontained function in line 15 removes vertex v_m and puts it in partition U_{i,j}. Finally, the created partitions are assigned to the fused partitioning V via add in line 16 as follows: V_{s_i,w_j} = H_{i,j}, V_{s_{i+1},w_{m+1+1}} = T_{i,j}, V_{s_{i+1},w_{m+1+1}} = U_{i,j}.

The head DAG choice. MSP chooses the DAG with edges as the head DAG to improve locality. Locality is improved because the head DAG is partitioned with LBC. LBC creates well-balanced partitions with good locality when applied to DAGs with edges. Selecting G_2 as the head DAG reduces inspector overhead. If both G_1 and G_2 are DAGs of kernels with dependency, then G_2 is chosen as the head DAG to reduce inspector overhead. When G_2 is partitioned first, MSP chooses backward pairing which is more efficient compared to forward pairing. Forward pairing traverses F and its transpose F^T and thus performs 2 * nz_F + 2 * n operations where nz_F is the number of nonzeros in F. However, backward pairing only traverses F and performs nz_F + n operations.
Algorithm 1: The MSP algorithm.

```plaintext
Input : \( G_1(\{V_1, E_1, c_1\}, G_2(\{V_2, E_2, c_2\}, F, r, \text{reuse\_ratio}) \)
Output: \( V' \)
/* (i) Vertex partitioning and partition pairing */
1. if \( |E_2| > 0 \) then
2. \([H, k]\) = LBC\((G_2, r)\).list(), \( T = 0, V = 0 \)
/* Backward pairing */
3. for \((i = 1 : H.size())\) do
4. for \((j = 1 : H_i.size())\) do
5. \( T_{i,j} = \text{BFS}(H_i, F, G_1) \)
6. \( V' = \text{add}(T_{i,j}, H_i) \)
7. end
8. end
9. if \(|V| > 2 \times (|V_1| + |V_2|)\) then
10. \( V' = \text{fusion} = \text{False, exit()} \)
else
11. \([H, k]\) = LBC\((G_1, r)\).list(), \( T = 0, V = 0 \)
/* Forward pairing */
12. for \((i = 1 : H.size())\) do
13. for \((j = 1 : H_i.size())\) do
14. \( T_{i,j} = \text{BFS}(H_i, F^T, G_2) \)
15. \( U_{i,j} = T_{i,j}.\text{remove}\_\text{uncontained}(F) \)
16. \( V' = \text{add}(H_i, T_{i,j}, U_{i,j}) \)
17. end
18. end
/* (ii) Merging and slacked vertex assignment */
19. \( S = \text{slack\_info}(V) \)
20. for \((\text{every } w\text{-partition pair } \{w, w'\} \in V'.\text{pairs})\) do
21. if \((SN(w) = 0) \land (SN(w') = 0)\) then \( V'.\text{merge}(w, w') \)
22. end
23. \( V' = V - S, \epsilon = |V| \times 0.001 \)
24. for \((i = 1 : V.b)\) do
25. for \((j = 1 : m_i)\) do
26. if \( \text{max\_diff}(V_i, V_{s_i, w}) > \epsilon \land S \neq 0\) then
27. \( S = V_{s_i, w}.\text{balance}\_\text{with\_pair}(S) \)
28. if \( \text{max\_diff}(V_i, V_{s_i, w}) > \epsilon \land S \neq 0\) then
29. \( S = V_{s_i, w}.\text{balance}\_\text{with\_slacks}(S) \)
30. end
31. end
32. if \( S \neq 0\) then \( S = V_{s_i}.\text{assign\_even}(S) \)
33. end
/* (iii) Packing */
34. if \( \text{reuse\_ratio} \geq 1\) then \( V'.\text{interleaved\_pack}(F) \)
35. else \( V'.\text{separated\_pack()} \)
```

Example. Figures 5b shows the output of MSP after the first step for the inputs in Figure 2b. MSP chooses \( G_1 \) as the head DAG because it has edges \((|E_1| > 1)\), \( G_2 \) has no edges. In vertex partitioning, \( G_1 \) is partitioned with LBC to create up to three w-partitions (because \( r = 3 \)) per s-partition. The created partitions are shown in Figure 5a and are stored in \( H \). The first s-partition \( V_{s_1} \) is stored in \( H_1 \) and its three w-partitions are indexed with \( H_{1,1}, H_{1,2}, \) and \( H_{1,3} \). Similarly, \( V_{s_2} \) is stored in \( H_2 \) and its only w-partition is in \( H_{2,1} \). Figure 5b shows the output of partition pairing.

Since \( G_1 \) is the head DAG, MSP uses forward pairing and performs a BFS from each partition in \( H \) to create self-contained pair partitions stored in \( T \). For example, a BFS from \( H_{1,1} = \{1, 2, 3, 4\} \) creates \( T_{1,1} = \{1, 2, 3, 4\} \). Since \( T_{1,1} \) and \( H_{1,1} \) are self-contained, no vertices are removed from \( T_{1,1} \) and thus \( U_{1,1} = \emptyset \). Finally, MSP puts \( H_{1,1} \) and \( T_{1,1} \) in \( V_{s_1, w_1} \) and \( V_{s_2, w_2} \) respectively, and adds \((V_{s_1, w_1}, V_{s_2, w_2})\) to \( V'.\text{pairs} \). The final pair partitions and pairings as shown in Figure 5b are: \( V' = \{\{H_{1,1}, H_{1,2}, H_{1,3}\}, \{H_{2,1}, T_{1,1}, T_{1,2}, T_{1,3}\}, \{T_{2,1}\}\} = \{\{1, 2, 3, 4\}, \{5, 6\}, \{7, 8, 9\}\}, \{\{10, 11\}\} \} \) and the pairing information is: \( V'.\text{pairs} = \{(V_{s_1, w_1}, V_{s_2, w_2}), (V_{s_1, w_2}, V_{s_2, w_3}), (V_{s_1, w_3}, V_{s_2, w_1}), (V_{s_2, w_2}, V_{s_3, w_1})\} \).

3.2.2 Merging and Slack Vertex Assignment. The second step of MSP reduces the number of synchronizations by merging some of the pair partitions in a merging phase. It also improves load balance by dispersing vertices across partitions using slacked vertex assignment.

Slack definitions: A vertex \( v \) can always run in its wavefront number \( l(v) \). However, the execution of vertex \( v \) can sometimes be postponed up to \( SN(v) \) wavefronts without having to move its dependent vertices to later wavefronts. \( SN(v) \) is the slack number of \( v \) and is defined as \( SNP(v) = \text{PC} - l(v) - \text{height}(v) \) where \( \text{height}(v) \) is the maximum path from a vertex \( v \) to a sink vertex (a sink vertex is a vertex without any outgoing edge), \( \text{PC} \) is the critical path of \( G \), and \( l(v) \) is the wavefront number of \( v \). A vertex with a positive slack number is a slack vertex. To compute vertex slack numbers efficiently, instead of visiting all vertices, MSP iterates over partitions and computes the slack number of each partition in the partitioned DAG, i.e. partition slack number. The computed slack number for a partition is assigned to all vertices of the partition. As shown in line 20 of Algorithm 1, all partition slack numbers of \( V' \) are computed via \( \text{slack\_info} \) and are stored in \( S \). For example, because vertices in \( V_{s_2, w_2} \) can be postponed one wavefront, from s-partition 2 to 3, their slack number is 1. Vertices in w-partitions \( V_{s_2, w_1} \) and \( V_{s_3, w_1} \) can not be moved because their slack numbers are zero.

Merging. MSP finds pair partitions with partition slack number of zero and then merges them as shown in lines 21-23. Since pair partitions are self contained, merging them does not affect the correctness of the schedule. Algorithm 1 visits all pair partitions \((w, w')\) in \( V'.\text{pairs} \) and merges them using the merge function in line 22 if their slack numbers are zero, i.e. \( SNP(w) = 0 \) and \( SNP(w') = 0 \). The resulting merged partition is stored in \( V' \) in place of the w-partition with the smaller s-partition number.

Slack vertex assignment. The algorithm then uses slacked vertex assignment to approximately load balance the w-partitions of an s-partition using a cost model. The cost of w-partition \( w \in V_{s_i} \) is defined as \( \text{cost}(w) = \sum_{w \in V_{s_i}} e(v) \). A w-partition is balanced if the maximal difference of its cost and the cost of other w-partitions in its s-partition is smaller than a threshold \( \epsilon \). The maximal difference for a w-partition...
inside a s-partition is computed by subtracting its cost from the cost of the w-partition (from the same s-partition) with the maximum cost.

MSP first removes all slacked vertices $S$ from the fused partitioning $V$ in line 24. It then goes over every s-partition $i$ and w-partition $j$ and balances $V_{s_i, w_j}$ by assigning a slacked vertex to it where possible. W-partition $V_{s_i, w_j}$ becomes balanced with vertices from its pair partition using the function balance_with_pair in line 27. If $V_{s_i, w_j}$ is still imbalanced, balance_with_slacks in line 28 balances the w-partition using the slacked vertices $v_l \in S$ that satisfy the following condition $l(v_l) < i < (l(v_l) + SN(v_l))$. Slack vertices in $S$ that depend on each other are dispersed as a group to the same w-partition for correctness. In line 30, slacked vertices in $S$ that are not postponed to later s-partitions are evenly divided between the w-partitions of the current s-partition ($V_s$) using the assign_even function.

Example. Figure 5d shows the output of the second step of MSP from the partitioning in Figure 5b. First pair partitions ($V_{s_2, w_1}$, $V_{s_3, w_2}$), shown with red dash-dotted circles in Figure 5b, are merged because their slack numbers are zero. The resulting merged partition is placed in $V_{s_1, w_0}$ to reduce synchronization as shown in Figure 5c. Then slacked vertex assignment balances the w-partitions in Figure 5c. The balanced partitions are shown in Figure 5d. The slacked vertices $S$, are shown with dotted blue circles in Figure 5c. The w-partitions in $V_s$ are balanced using vertices of their pair partitions, e.g. the yellow dash-dotted vertices 5 and 6 are moved to $w_2$ in $V_s$ as shown in Figure 5d. balance_with_slacks is used to balance partitions in $V_s$. This is because the vertices in $S$ do not belong to the pair partitions of the w-partitions in $V_s$. However, since the slack vertices in $S$ can execute in either s-partition two or three because they are from s-partition one and have a slack number of one, they are used to balance the w-partitions in $V_{s_2}$.

3.2.3 Packing. The third step of MSP reorders the vertices inside a w-partition to improve data locality for a thread within each kernel or between the two kernels. The previous steps of the algorithm create w-partitions that are composed of vertices of one or both kernels however the order of execution is not defined. Using the reuse ratio, the order at which the nodes in a w-partition should be executed is determined with a packing strategy. MSP has two packing strategies: (i) in interleaved packing, the vertices of the two DAGs in a w-partition are interleaved for execution and (ii) in separated packing the vertices of each kernel are executed separately. Interleaved packing improves temporal locality between kernels while separated packing enhances spatial and temporal locality within kernels. When the reuse ratio is greater than one, in line 32 of Algorithm 1 function interleaved_pack is called to interleave iterations of the two kernels based on F. Otherwise, separated_pack is called (line 33) to pack iterations of each kernel separately.

| ID   | Name          | Nonzeros     |
|-----|---------------|--------------|
| 1   | Flan_1565     | 117.4×10^6   |
| 2   | bone010       | 71.7×10^6    |
| 3   | Hook_1498     | 60.9×10^6    |
| 4   | af_shell10    | 52.3×10^6    |
| 5   | Emilia_923    | 41×10^6      |
| 6   | StocF-1465    | 21×10^6      |
| 7   | af_0_k101     | 17.6×10^5    |
| 8   | ted_B_unscal  | 0.14×10^5    |

Table 1. The list of sparse matrices.

Example. Figure 2e shows the output of MSP’s third step from the partitioning in Figure 5d. Since the reuse ratio is smaller than one separated packing is chosen thus $V_{s_1, w_0}$ is stored as $V_{s_1, w_0} = \{10, 11, 10, 11\}$. Vertices are ordered to keep dependent iterations of SpTRSV and consecutive iterations SpMV next to each other.

4 Experimental Results

We compare the performance of sparse fusion to MKL [56] and ParSy [8], two state-of-the-art tools that accelerate individual sparse kernels, which we call unfused implementations. Sparse fusion is also compared to three fused implementations that we create. To our knowledge, sparse fusion is the first work that provides a fused implementation of sparse kernels where at least one kernel has loop-carried dependencies. For comparison, we also create three fused implementations of sparse kernels by applying LBC, DAGP, and a wavefront technique to the joint DAG of the two input sparse kernels and create a schedule for execution using the created partitioning, the methods will be referred to as fused LBC, fused DAGP, and fused wavefront in order.

Setup. The set of symmetric positive definite matrices listed in Table 1 are used for experimental results. The matrices are from [12] and with real values in double precision. The testbed architecture is a multicore processor with 12 cores of a Xeon E5-2680v3 processor with 30MB L3 cache. All generated codes, implementations of different approaches, and library drivers are compiled with GCC v.7.2.0 compiler and with the -O3 flag. Matrices are first reordered with METIS [24] to improve parallelism.

We compare sparse fusion with two unfused implementations where each kernel is optimized separately: I. ParSy applies LBC to DAGs that have edges. For parallel loops, the method runs all iterations in parallel. LBC is developed for L-factors [11] or chordal DAGs. Thus, we make DAGs chordal before using LBC. II. MKL uses Intel MKL [56] routines with MKL 2019.3.199 and calls them separately for each kernel.

Sparse fusion is also compared to three fused approaches all of which take as input the joint DAG; the joint DAG is created from combining the DAGs of the input kernels using the inter-DAG dependency matrix $F$. We then implement three approaches to build the fused schedule from the joint DAG: I. Fused wavefront traverses the joint DAG in topological order and builds a list of wavefronts that represent vertices of both DAGs that can run in parallel. II. Fused LBC applies the LBC
Table 2. The list of kernel combinations. CD: loops with carried dependencies, SpIC0: Sparse Incomplete Cholesky with zero fill-in, SpILU0: Sparse Incomplete LU with zero fill-in, DSCAL: scaling rows and columns of a sparse matrix.

| ID | Kernel combination | Operations | Dependency DAGs | Reuse Ratio |
|----|-------------------|------------|-----------------|-------------|
| 1  | SpTRSV CSR - SpTRSV CSR | $x = L^{-1}y, z = L^{-1}x$ | CD - CD | $\max(2n+\text{size}, 2m) \geq 1$ |
| 2  | SpMV CSR - SpTRSV CSR | $y = Ax, z = L^{-1}y$ | Parallel - CD | $\max(2m, \text{size}, 2n) < 1$ |
| 3  | DSCAL CSR - SpILU CSR | $LU \approx \text{DAD}^T$ | Parallel - CD | $\max(2\text{size}, \text{size}, 2m) \geq 1$ |
| 4  | SpTRSV CSR - SpMV CSC | $y = L^{-1}x, z = Ay$ | CD - Parallel | $\max(2\text{size}, \text{size}, 2n) < 1$ |
| 5  | SpIC0 CSC - SpTRSV CSC | $LL^T \approx A, y = L^{-1}x$ | CD - CD | $\max(\text{size}, \text{size}, 2n) \geq 1$ |
| 6  | SpILU0 CSR - SpTRSV CSR | $LU \approx A, y = L^{-1}x$ | CD - CD | $\max(\text{size}, \text{size}, 2n) \geq 1$ |
| 7  | DSCAL CSC - SpIC0 CSC | $LL^T \approx \text{DAD}^T$ | Parallel - CD | $\max(\text{size}, \text{size}, 2n) \geq 1$ |

Table 3. The achieved GFLOP/s for the baseline code for the kernel combinations in Table 2 and for matrices in Table 1.

| Matrix ID | Kernel Combination ID |
|----------|-----------------------|
| 1        | 1 1.52 1.54 0.45 1.55 0.61 0.43 0.61 |
| 2        | 2 1.5 1.54 0.45 1.54 0.61 0.45 0.61 |
| 3        | 3 1.4 1.45 0.47 1.45 0.48 0.50 0.47 |
| 4        | 4 1.47 1.48 0.72 1.49 0.50 0.77 0.47 |
| 5        | 5 1.42 1.47 0.45 1.47 0.51 0.46 0.49 |
| 6        | 6 0.91 1.14 0.17 1.14 0.33 0.18 0.32 |
| 7        | 7 1.47 1.50 0.73 1.49 0.49 0.77 0.48 |
| 8        | 8 1.41 1.70 0.89 1.70 0.44 0.76 0.42 |

algorithm to the joint DAG and creates a set of s-partitions each composed of independent w-partitions. Then the s-partitions are executed sequentially and w-partitions inside an s-partition are executed in parallel. LBC is taken from ParSy and its parameters are tuned for best performance. The joint DAG is first made chordal and then passed to LBC. III. Fused DAGP applies the DAGP partitioning algorithm to the joint DAG and then executes all independent partitions that are in the same wavefront in parallel. DAGP is used with METIS for its initial partitioning, with one run (runs=1) and the remaining parameters are set to default.

The list of sparse kernel combinations investigated are in Table 2. To demonstrate sparse fusion’s capabilities, the sparse kernels are selected with different combinations of storage formats, i.e. CSR and compressed sparse column (CSC) storage, different combinations of parallel loops and loops with carried dependencies, and a variety of memory access pattern behaviour. For example, combinations of SpTRSV, $Lx = b$ and SpMV are main bottlenecks in conjugate gradient methods [4, 60], GMRES [9], Gauss-Seidel [41]. Pre-conditioned Krylov methods [17] and Newton solvers [45] frequently use kernel combinations 3, 5, 6, 7. The s-step Krylov solvers [6] and s-step optimization methods used in machine learning [45] provide even more opportunities to interleave iterations. Thus, they use these kernel combinations significantly more than their classic formulations.

Sparse Fusion’s Performance. Figure 6 shows the performance of the fused code from sparse fusion, the unfused implementation from ParSy and MKL, and the fused wavefront, fused LBC, and fused DAGP implementations. All execution times are normalized over a baseline. The baseline is obtained by running each kernel individually with a sequential implementation. The floating point operations per second (FLOP/s) for each implementation can be obtained by multiplying the baseline FLOP/s from Table 3 with the speedups in Figure 6. The sparse fusion’s fused code is on average 1.6× faster than ParSy’s executor code and 5.1× faster than MKL across all kernel combinations. Even though sparse fusion is on average 11.5× faster than MKL for ILU0-TRSV, since ILU0 only has a sequential implementation in MKL, the speedup of this kernel combination is excluded from the average speedups. The fused code from sparse fusion is on average 2.5×, 5.1×, and 7.2× faster than in order fused wavefront, fused LBC, and fused DAGP. Obtained speedups of sparse fusion over ParSy (the fastest unfused implementation) for SpILU0-SpTRSV and SpIC0-SpTRSV is lower than other kernel combinations. Because SpIC0 and SpILU0 have a high execution time, when combined with others sparse kernels with a noticeably lower execution time, the realized speedup from fusion will not be significant.

Locality in Sparse Fusion. Figure 7 shows the efficiency of the two packing strategies to improve locality. The effect of the packing strategy is shown for kernel combinations with a reuse ratio smaller and larger than one as shown in Table 2. Kernel combinations 1, 3, 5, 6, and 7 share the sparse matrix $L$ and thus have a reuse ratio larger than one while combination 2 and 4 only share vector $y$ leading to a reuse ratio lower than one. Figure 7 shows the range of speedup over all matrices for the selected packing strategy versus the other other packing method for each combination. As shown, the selected packing strategy in sparse fusion improves the performance in 88% of kernel combinations and matrices and provides 1-3.9× improvement in both categories.

Figure 8 shows the average memory access latency [18] of sparse fusion, the fastest unfused implementation (ParSy),
and the fastest fused partitioning-based implementation (Fused LBC) for all kernel combinations normalized over the ParSy average memory access latency (shown for matrix bone010 as example, other matrices exhibit similar behavior). The average memory access latency is used as a proxy for locality and is computed using the number of accesses to L1, LLC, and TLB measured with PAPI performance counters [52].

For kernels 1, 3, 5, 6, and 7 where the reuse ratio is larger than one, the memory access latency of ParSy is on average 1.3× larger than that of sparse fusion. Because of their high reuse ratio, these kernels benefit from optimizing locality between kernels made possible via interleaved packing. ParSy optimizes locality in each kernel individually. When applied to the joint DAG, LBC can potentially improve the temporal locality between kernels and thus there is only a small gap between the memory access latency of sparse fusion and that of fused LBC. For kernels 2 and 4 where the reuse ratio is smaller than one, the gap between the memory access latency of sparse fusion and fused LBC is larger than the gap between the memory access latency of sparse fusion and ParSy. Sparse fusion and ParSy both improve data locality within each kernel for these kernel combinations.

**Load Balance and Synchronization in Sparse Fusion.**

Figure 8 shows the OpenMP potential gain [44] of sparse fusion, ParSy, and Fused LBC for all kernel combinations normalized over ParSy’s potential gain (shown for matrix bone010 as example, but all other matrices in Table 1 follow similar behavior.) The OpenMP potential gain is a metric in Vtune [63] that shows the total parallelism overhead, e.g. wait-time due to load imbalance and synchronization overhead, divided by the number of threads. This metric is used to measure the load imbalance and synchronization overhead in ParSy, fused LBC, and sparse fusion.

Kernel combinations 2 and 4 have slack vertices that provide opportunities to balance workloads. For example, for matrices shown in Table 1, between 35-76% vertices can be slacked thus the potential gain balance of ParSy is 1.6× larger than sparse fusion and 2.4× lower than fused LBC. ParSy can only improve load balance using the workloads of an individual kernel. As shown in Figure 1, for the kernel combination 5, the joint DAG has a small number of parallel iterations.
in final wavefronts that makes the final s-partitions of the LBC fused implementation imbalanced (a similar trend exists for kernel combination 6). For these kernel combinations, the code from sparse fusion has on average 33% fewer synchronization barriers compared to ParSy due to merging. For kernel combinations 1, 2, 3, 4, and 7 the potential gain in sparse fusion is 1.3× less than that of ParSy. Merging in sparse fusion reduces the number of synchronizations in the fused code on average 50% compared to that of ParSy.

**Inspector Time.** Figure 9 shows the number of times that the executor should run to amortize the cost of inspection for implementations that have an inspector. For space only combinations 1, 3, 4, and 5 are shown, others follow the same trend. The number of executor runs (NER) that amortize the cost of inspector for an implementation is calculated using

\[
\text{Inspection Time} = \text{Baseline Time} - \text{Executor Time}
\]

The baseline time is obtained by running each kernel individually with a sequential implementation, the inspector and executor times belong to the specific implementation. The fused LBC implementation has a NER of 3.1-745. The high inspection time is because of the high cost of converting the joint DAG into a chordal DAG, typically consuming 64% of its inspection time. The NER of the fused DAGP implementation is either negative or higher than 80. The fused wavefront implementation sometimes has a negative NER because the executor time is slower than the baseline time. As shown, sparse fusion and fused wavefront have the lowest NER amongst all implementations. Sparse fusion’s low inspection time is due to pairing strategies that enable partitioning one DAG at a time. Kernel combinations such as, SpiC0-TRSV and SpiLU0-TRSV only need one iteration to amortize the inspection time and SpTRSV-SpMV, SpTRSV-SptRSV, and SpMV-SpTRSV need between 11-50 iterations. Sparse kernel combinations are commonly used in iterative solvers in scientific applications. Even with preconditioning, these solvers typically converge to an accurate solution after ten of thousands of iterations [4, 25, 36], hence amortizing the overhead of inspection.

**6 Conclusion**

We present sparse fusion and demonstrate how it improves parallelism, load balance, and data locality in sparse matrix combinations compared to when sparse kernels are optimized separately. Sparse fusion inspects the DAGs of the input sparse kernels and uses the MSP algorithm to balance the workload between wavefronts and determine whether to optimize data locality for within or between the kernels. Sparse fusion’s generated code outperforms state-of-the-art implementations for sparse matrix optimizations. In future work, we plan to investigate strategies that select the most profitable loops to be fused to support the fusion of more than two loops.

**Acknowledgments**

This work was supported in part by NSERC Discovery Grants (RGPIN-06516, DGECR00303), the Canada Research Chairs...
program, and U.S. NSF awards NSF CCF-1814888, NSF CCF-1657175; used the Extreme Science and Engineering Discovery Environment (XSEDE) [Towns et al. 2014] which is supported by NSF grant number ACI-1548562; and was enabled in part by Compute Canada and SciNet 1.

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