Abstract—The divergence in the computer architecture landscape has resulted in different architectures being considered mainstream at the same time. For application and algorithm developers, a dilemma arises when one must focus on using underlying architectural features to extract the best performance on each of these architectures, while writing portable code at the same time. We focus on this problem with graph analytics as our target application domain. In this paper, we present an abstraction-based methodology for performance-portable graph algorithm design on manycore architectures. We demonstrate our approach by systematically optimizing algorithms for the problems of breadth-first search, color propagation, and strongly connected components. We use Kokkos, a manycore library and programming model, for prototyping our algorithms. Our portable implementation of the strongly connected components algorithm on the NVIDIA Tesla K40M is up to 3.25× faster than a state-of-the-art parallel CPU implementation on a dual-socket Sandy Bridge compute node.

Keywords—graph computations; BFS; color propagation; GPU; parallel performance; portability

I. INTRODUCTION

Our work attempts to answer the following questions in the context of graph computations and manycore processors:

(a) Can we identify frequently-used optimization strategies from the large and growing collection of tuned parallel graph computation implementations (e.g., [2], [5], [9], [23], [37]), and create a structured methodology for designing new parallel algorithms? If one were to build a new framework for high-performance domain-specific graph computations, what would be the key optimization strategies to consider, and best practices to follow?

(b) In addition to parallel-for, data-parallel scans, reductions, and sorting methods, what are some common abstractions used to design parallel graph algorithms?

(c) Is it possible to develop performance-portable implementations of graph algorithms using advanced parallel programming libraries and frameworks with the optimizations and abstractions identified above?

We begin by observing that several recent graph algorithms and their efficient implementations follow the loop nest structure shown in Algorithm 1. The first point to note in Algorithm 1 is that the listing uses only simple, array-based data structures. Current state-of-the-art parallel implementations for several graph problems use array-based stacks, queues, and priority queues, as these structures are more amenable to applying data-parallel operations such as scans and reductions. Implementations of this general template differ in terms of graph representation, data structure access patterns, number of iterations of the outer loop, graph topology-based heuristics to reduce total work, synchronization overhead, etc. Intra- and inter-iteration dependencies hinder automatic compiler-based loop transformations such as unrolling, coalescing, collapse, and fusion.

Consider level-synchronous parallelizations of Breadth-First Search (BFS). niter, the number of outer-loop iterations, is bounded by the graph diameter. The arrays \( S_i \) correspond to the vertices in the current frontier, and the adjacencies of these vertices can be visited in parallel. Arrays of size \( n \) \((A_i)\) are used to store parent information, whether a vertex has been previously visited or not, and distance from the source vertex. Finally, there is a barrier synchronization before every iteration of the outer loop. For low-diameter (diameter is \( O(\log n) \)) graphs, the overhead of barrier synchronization is insignificant in comparison to the work performed in the inner loops. The arrays \( S_i \) store

**Algorithm 1** A template followed by several serial and parallel graph algorithms operating on a sparse graph \( G(V,E) \). \( m = |E|, n = |V|, \) and \( m = O(n \log n) \).

Initialize temp/result arrays \( A_i[1..n], 1 \leq t \leq l \).

\[ l = O(1) \]

Initialize \( S_i[1..n] \).

\[ \text{for } i = 1 \text{ to } \text{niter} \text{ do} \]

\[ \text{Initialize } S_{i+1}[1..n], \]

\[ \sum_i |S_i| = O(m) \]

\[ \text{for } j = 1 \text{ to } |S_i| \text{ do} \]

\[ u \leftarrow S_i[j] \]

\[ \text{Read/update } A_i[u], 1 \leq t \leq l. \]

\[ \text{for } k = 1 \text{ to } |E[u]| \text{ do} \]

\[ v \leftarrow E[u][k] \]

\[ \text{Read/update } A_i[v]. \]

\[ \text{Read/update } S_{i+1}. \]

\[ \text{Read/update } A_i[u]. \]
vertices in an arbitrary order for BFS. For other algorithms, such as ones for single-source shortest paths (SSSP), the ordering of vertices in $S_i$ may be important to reduce the outer-loop iteration count. The direction-optimizing heuristic in a recent BFS algorithm [3] and the push-pull optimization in a recent parallel SSSP algorithm [6] can be viewed as work-reducing heuristics to switch between alternate representations of $S_i$ in the inner loop.

The label propagation community detection heuristic [29] and the PULP graph partitioning strategy [34] also fit within this general template. They differ in the access patterns of the temporary arrays and the result arrays in the inner loops. Algorithms that are similar to level-synchronous BFS, such as betweenness centrality [22], approximate diameter [8], strongly connected components [33], and biconnected components [32] also follow a similar structure. In fact, several PRAM graph algorithms can be viewed as instances of this template, and they would result in polylogarithmic parallel time algorithms (assuming low-diameter graphs and/or a $O(\log n)$ bound for the outer-loop iterations). This template is not restricted to shared-memory algorithms. Distributed-memory approaches for K-core decomposition also use a label propagation-like strategy [25], and implementations differ in the number of outer-loop iterations. Finally, opensource software packages providing state-of-the-art parallel implementations of graph algorithms, such as the Parallel Boost graph library [15], MTGL [4], Galois [19], SNAP [21], PowerGraph [14], Ligra [31], NetworKit [35], etc., include several programs that fall under the Algorithm 1 template.

So we hypothesize that the first step towards creating an efficient parallel implementation of a graph algorithm would be to recast it such that it fits the general template shown in Algorithm 1. The focus of the current work is efficient graph analytics on manycore platforms such as NVIDIA and AMD GPUs and the Intel Xeon Phi MIC coprocessor. We do not want to reinvent the wheel for data-parallel subroutine implementations and parallel-for support. Hence, we use an emerging node-level library and programming model called Kokkos [11], that lets us write code that is portable to GPUs, Intel Xeon Phi, as well as Intel and AMD x86 CPUs. In Section II, we discuss key Kokkos features that enable us to quickly develop and compare alternate implementations.

As the next step, we develop several manycore implementations for the graph problems of BFS, color propagation, and strongly-connected components (SCC), expressing them in the template shown in Algorithm 1. The inner loop nests of BFS and color propagation have several differences, and so we explore both of these problems. The general manycore SCC algorithm is based on our prior multicore SCC approach called Multistep [33]. Use of Kokkos lets us develop several alternatives for each problem and conduct a methodical evaluation of optimizations.

In Section III, we present the third step, the key optimizations that are critical to manycore performance and portability. We primarily evaluate several tuned loop transformation strategies for the inner loop nest, and we customize these strategies for our use case of small-world graph analytics. These loop transformations, in essence, improve load balance and reduce irregular memory accesses. Our proposed strategies are similar to a compiler-based loop collapse [1], [30]. However, compilers cannot automatically do this because of loop-carried dependencies. To the best of our knowledge, this is the first work to explore the loop transformations and manycore optimizations for SCC and color propagation problems. We arrive at this portable and performant manycore implementation of SCC using our algorithmic template, a Kokkos-based implementation, and architecture-aware optimizations.

The main observations from our empirical performance evaluation (see Section IV) are as follows:

- Our new loop collapse strategy, termed Local Manhattan Collapse, is very effective on GPUs and consistently results in the highest-performing variant for several problems.
- A GPU SCC implementation using the Local Manhattan Collapse strategy demonstrates up to a 3.25× speedup relative to a state-of-the-art parallel GPU implementation running on a dual-socket computer node.
- We find our GPU BFS implementation averages 1.74 GTEPS across a suite of 12 test graphs, comparable to the current state-of-the-art, without any BFS-specific tuning.

II. PORTABLE GRAPH ALGORITHMS FOR MANYCORE

A. The Kokkos Programming Model

The Kokkos library [11] was originally developed as a back-end for providing portable performance for scientific computing frameworks, but has since been extended to a more general-purpose library for parallel execution. The two primary capabilities of Kokkos include polymorphic multidimensional arrays optimized for varying data access patterns/layouts in different architectures and thread parallel execution that allows for fine-grained data parallelism on manycore devices.

The parallel execution model follows a dispatch model, where a single master CPU thread divides some $N$ units of work to be processed on GPU. Each unit of work is executed by a single thread or thread team. On GPU, a thread team is comprised of multiple warps each executing on the same multiprocessor. This team of threads operates in a data parallel SIMT fashion, and is able to “communicate” via shared memory. In addition to optimizing the data layout in different devices, Kokkos also provides us the option to use the hierarchy of memory in manycore devices, such as thread block shared memory and texture cache. We use these features of Kokkos for the appropriate data structures.
We also use the fast atomic operations and “thread team level” scan and reduction operations to synchronize between different threads in a thread block. One of the key design decisions we made, using low-level simple array-based data structures, helps us when using Kokkos, as the layout of these simple arrays is then optimized by Kokkos in different devices (CPUs, GPUs and Phis), different types of memory (e.g., shared memory), and different access patterns (e.g., coalesced access). Use of custom data structures that are optimized for any single architecture would have prevented Kokkos level optimizations, in turn affecting portability.

B. BFS

BFS is one of the most widely used basic graph subroutines, appearing in a vast number of more complex graph analytics. The goal of BFS is commonly to determine from a given root vertex either reachability status, distance, or analytics. The goal of BFS is commonly to determine from tines, appearing in a vast number of more complex graph subroutines implemented in our framework to perform graph SCC decomposition via the Multistep procedure (we refer the reader to [33] for a more detailed description). Once again, outside of a few changes to initializations and the very innermost loops, few alterations need to occur to the original BFS and color propagation codes for the SCC problem.

C. Color Propagation

Color propagation is an iterative procedure that is useful for many different graph connectivity problems [27], [31]–[33]. An overview of the general algorithm is given by Algorithm 2. Note how it also follows the Algorithm 1 template, where \( S_i \) is our current queue and the result array \( A \) can be considered as the current color assignments for all vertices in the graph. We initialize \( A \) to be unique vertex identifiers and \( S_1 \) as all vertices in the graph. We then examine all edges, and when there exists a source vertex that has a higher color than one of its neighbors, that vertex propagates its color to the neighbor. The next work set \( S_{i+1} \) is comprised of vertices that have had their color altered. This process continues iteratively until no further propagations occur. As with BFS, we implement color propagation in a straightforward manner within our general framework.

Algorithm 2 Color Propagation pseudocode.

\[
\begin{align*}
A[1..n] &\leftarrow 1..n & \text{\( \triangleright \) Set } A[i] = i, 1 \leq i \leq n \\
S_1[1..n] &\leftarrow 1..n & \text{\( \triangleright \) Set } S_1[i] = i, 1 \leq i \leq n \\
i &\leftarrow 1 \\
\text{while } S_i \neq \emptyset \text{ do} & \\
\quad S_{i+1} &\leftarrow \emptyset & \\
\quad \text{for } j \text{ = 1 to } |S_i| \text{ do} &  \\
\quad \quad u &\leftarrow S_j & \\
\quad \quad \text{for } k \text{ = 1 to } |E[u]| \text{ do} &  \\
\quad \quad \quad v &\leftarrow E[u][k] & \\
\quad \quad \quad \text{if } A[u] > A[v] \text{ then} &  \\
\quad \quad \quad \quad A[v] &\leftarrow A[u] & \\
\quad \quad \quad \text{Add } v \text{ to } S_{i+1} &  \\
\quad i &\leftarrow i + 1 & \\
\end{align*}
\]

D. Strongly Connected Components

The problem of computing strongly connected components (SCCs) in large directed small-world graphs is a common analytic for social networks [24] and a preprocessing step in scientific computing (among other uses) [28]. Using either BFS or coloring, straightforward parallel strongly connected component decomposition algorithms can be implemented [27], [36]. Combining both subroutines into an efficient Multistep procedure can result in considerable speedup for small-world graphs [33]. We use the BFS and color propagation subroutines implemented in our framework to perform graph SCC decomposition via the Multistep procedure (we refer the reader to [33] for a more detailed description). Once again, outside of a few changes to initializations and the very innermost loops, few alterations need to occur to the original BFS and color propagation codes for the SCC problem.

III. OPTIMIZATION METHODOLOGIES

In this section, we describe the optimization techniques used to achieve scalable performance on manycore architectures. These techniques are applicable to any algorithm that fits the template described in Algorithm 1. Furthermore, the optimizations are general enough for architectures that share similar characteristics, such as a very high core count, hierarchical memories, and small amounts of memory per thread. These characteristics of present day GPUs is expected to hold or become increasingly important in future manycore architectures. As a result, the optimizations described here are critical for scalable algorithms on modern and future systems.

A. Thread Teams, Local Synchronization, Shared and Global Memory

Current GPUs are organized as a number of streaming multiprocessors (for instance, 16 in a NVIDIA Maxwell GM204, each with a number of smaller cores (e.g., 128 CUDA cores in GM204). The number of threads that can be scheduled in a single streaming multiprocessor of a GPU can be up to 2048. The number of warps per streaming multiprocessor is 48-64, and the number of thread blocks is 8-16,
depending on the microarchitecture. A similar hierarchy is also seen in the multiple hyperthreads per core of a Xeon Phi coprocessors, along with NUMA effects due to placement of cores near different memory regions. This results in multiple levels of parallelism for which algorithm developers need to design. The programming model in Kokkos abstracts this to a thread team, where a thread team corresponds to a thread block on GPU. In order to effectively utilize a streaming multiprocessor (SM), it is crucial to be able to schedule multiple thread blocks in each SM. Within each thread block, there is enough concurrency for thousands of threads, so that multiple warps can be kept busy at the same time. All of our algorithms use the thread teams concept to synchronize locally and utilize shared memory to communicate within a team when necessary.

The number of thread blocks that can be scheduled concurrently in a single SM is determined by the amount of shared memory used by each thread block (or a thread team). As all the threads in a team use the shared memory to synchronize among themselves before synchronizing to the global memory, the amount of shared memory used is an important resource. Increasing its size would reduce the number of writes to global memory by doing more local synchronizations, but it would also decrease the number of concurrent thread blocks that can be scheduled. Our approach balances shared memory usage with the parallelism available within each thread block. Finally, it is important that reads and writes to global memory are coalesced. Essentially, we want reads and writes for a single warp to be performed at neighboring global memory addresses to reduce total memory bandwidth, improve cache utilization, and ensure that threads in the warp are not idle waiting for the memory request of a single thread.

B. Hierarchical Exploration to Improve SM Utilization

A common optimization technique for algorithms dealing with irregular graph structure is fine-grained warp utilization by limiting the serial of our algorithm, we remove a chunk of vertices \( V_T \) from the input work set \( S_i \) and pass it to a Kokkos thread team \( T \). For good team-level work balance and multiprocessor utilization, the size of \( V_T \) is usually within a small factor of the size of \( T \). The threads in each team work to process their input set, placing the high-degree vertices they encounter into a team-owned queue (when the degree is greater than the size of the thread team \(|T|\)) or warp-owned queue (when the degree is smaller than the size of a thread team, but larger than the size of a warp \(|W|\)). Smaller vertices get placed into a small thread-owned buffer \( Q_t \) for later serial expansion. The vertices in the team-level queue \( Q_T \) are collectively expanded by all threads, with potential updates to \( S_i+1 \) kept in team-level shared memory. Once the \( Q_T \) queue is exhausted, the warp queue \( Q_W \) is examined. Each warp removes a vertex from the queue, and cooperatively expands its adjacencies. Finally, each individual thread serially expands the vertices in its buffer. Once all work is exhausted, the team collectively pushes their updates to the next iteration’s work set \( S_{i+1} \).

We use team-level scans and reductions whenever possible to minimize global synchronizations.

The primary benefit to this type of approach is that it allows fine-grained warp utilization by limiting the serial

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Algorithm 3 Hierarchical Expansion.

Initialize \( A \) and \( S_1 \).

for \( i = 1 \) to \( nIter \) do

Initialize \( S_{i+1}[1...n] \).

for all Thread Teams do \( \triangleright \) Team-level parallelism

Retrieve subset \( V_T \) from \( S_i \).

for \( j = 1 \) to \(|V_T|\) do \( \triangleright \) Thread-level parallelism

\( v \leftarrow V_T[j] \).

if \( |E[v]| > |T| \) then

Add \( v \) to \( Q_T \). \( \triangleright \) Team-shared Queue

else if \( |E[v]| > |W| \) then

Add \( v \) to \( Q_W \). \( \triangleright \) Warp-shared Queue

else

Add \( v \) to \( Q_t \) \( \triangleright \) Thread-owned Queue

Team-level synchronization

for \( j = 1 \) to \(|Q_T|\) do \( \triangleright \) Warp-level parallelism

\( u \leftarrow E[v][k] \). \( \triangleright \) Thread-level parallelism

Read/update \( A[u] \).

for \( k = 1 \) to \(|E[v]|\) do \( \triangleright \) Serial expansion by thread

\( u \leftarrow E[v][k] \). \( \triangleright \) Thread-level parallelism

Read/update \( A[u] \).

Team-level synchronization

Update \( S_{i+1} \).
expansion of high degree vertices by a single thread or warp. This leads to better load balance at the thread and warp level. This approach also allows the use of shared memory to create new queues for the next iteration of an irregular graph problem, reducing the number of global synchronizations required. However, as vertices are assigned to a single team statically, there can still be some imbalance at the highest level. All teams might finish their work long before the team owning a highly-skewed vertex completes, delaying the start of the next iterations and vastly under-utilizing available processing resources.

C. Loop Collapse for Better Load Balance

As shown by the template in Algorithm 1, many graph algorithms follow the pattern of two nested loops, where the outer loop is over the vertices and the nested inner loop is over the edges. Often, these two loops are not perfectly nested, as the vertex contents of the outer loop determine the start and end indices for the edges examined of the inner loop. There might be additional operations within the outer loop, such as changing the properties of vertices, adding vertices to the next queue, etc. While perfectly nested loops are great candidates for compiler-based optimizations, loops containing these other operations cannot be automatically optimized by compilers.

The importance of collapsing these loops increases when both loops are parallelizable and when the work in different outer loop iterations is heavily unbalanced. In graph analytic algorithms on graphs with skewed degree distributions, when the work in the outer loop varies based on the degree of the vertices, collapsing the inner loop is critical. When there are few threads, like in the CPU, a simple dynamic scheduling runtime can alleviate the problem [33]. However, it is hard to scale this approach to the thousands of threads in manycore devices. In our framework, we do the optimization a compiler might do, and collapse the two loops over vertices and edges into a single loop over all possible edges.

We employ the Manhattan Collapse [30], where a prefix-sum operation, easily parallelizable on GPUs with a scan-based procedure, is used to compute the bounds of each outer loop iteration. With the results of the prefix sums, a binary search is then used to compute the indices of the original inner-loop and outer-loop within the collapsed loop (references as HIGHESTLESSTHAN in Algorithms 4 and 5). The overhead associated with reverse-engineering the vertex information is offset by the good load balance achieved by each thread. This general approach has been explored before by Merrill et al. for GPUs in the context of BFS [23] and by Davidson et al. for SSSP [9]. As with the work of Davidson et al., we consider two forms of the Manhattan Collapse, implementing it at both the global and local level.

1) Local Manhattan Collapse: For our local implementation, we do not require any additional global storage, apart from the queues and work arrays updated in the algorithm. An overview of this approach is given by Algorithm 4. We statically partition our work set $S_i$ on a per-vertex basis and pass each partition $V_T$ to our thread teams. The thread team computes prefix sums $P$ over $V_T$ based on out-degree. $P$ is stored in shared memory. The final prefix sum in $P$ is the sum of edges for $V_T$, and therefore proportional to the total work that the team needs to do. We can then equally distribute this work among all the threads in the team. To get a specific edge based on a per-thread work assignment $j$, the source vertex is determined by examining the prefix sums array, and finding the index $k$ that corresponds to a value in $P$ greater than or equal to $j$, and less than the value at the next highest index. The specific out-edge $u$ from the source vertex can be found by using the difference between the work assignment and the value at the found index in the prefix sums. With the $(u, v)$ pair, the thread can now perform its assigned work.

Algorithm 4 Local Manhattan loop collapse.

```
Initialize $A_t$ and $S_1$
for $i = 1$ to niter do
    Initialize $S_{i+1}[1..n]$
    for all Thread Teams do ▷ Team-level parallelism
        Retrieve subset $V_T$ from $S_i$
        $P ← \text{PrefixSums}(V_T)$
        $\text{Max} ← \text{max}(P)$
    for $j = 1$ to $\text{Max}$ do ▷ Thread-level parallelism
        $k ← \text{HIGHESTLESSTHAN}(P, j)$
        $u ← V_T[k]$
        $v ← E[u][j - P[k]]$
        Read/update $A_t[u]$ and $A_t[v]$
    Team-level synchronization
    Update $S_{i+1}$
```

The primary benefit of the Local Manhattan Collapse is that it leads to full warp and thread utilization of processor resources. When the cost of looking up a work assignment is low compared to the work that needs to be done, this approach is highly beneficial. As with Hierarchical Exploration, a major drawback to doing the Local Manhattan Collapse is that a vertex is still assigned to a single team, which might lead to work imbalances for highly skewed graphs.

2) Global Manhattan Collapse: To alleviate any potential work imbalance issues, we implemented a fully-partitioned approach, where the prefix sums for the current iteration are computed on the previous iteration as updates were pushed to the next-iteration work set $S_{i+1}$. By doing this, we can statically distribute an equal number of edges to each team instead of vertices. As can be seen in Algorithm 5, the approach closely follows our local method. The primary difference lies in the prefix sum arrays, $P_i$ and $P_{i+1}$, which must be globally stored and synchronously updated. To minimize data transfer requirements, each thread team can determine its start and end offsets in $P_i$ and do a single transfer of
the needed portion to shared memory. Additionally, pushes to $P_{t+1}$ and $S_{t+1}$ can also be coalesced, with only a single atomic update required per team. Because each team needs to determine the offset to start writing to $P_{t+1}$, as well as the current running sum, we package both these values into a single atomically-updated 64-bit long int and perform an atomic fetch-and-add on a current global value.

Ideally, the Global Manhattan Collapse should offer the best work partitioning among thread teams and fastest execution times for a given algorithm. However, as we will show in our results, there are other key factors that hurt the performance of the Global Manhattan Collapse relative to the Local method. For simple graph algorithms with minimal work per edge, the cost of reading and writing to an additional global array is relatively high. Amortizing this startup and end cost by increasing work per team is not necessarily a good solution, as we would ideally like to have an as-large-as-is-practical number of teams to hide the memory access latencies inherent to the rest of the implemented algorithm. Further, on graphs with a relatively consistent degree, or a modest number of outliers, this method offers no additional benefit in terms of equal per-thread work distribution relative to the local collapse. Finally, the maximal degree of many real-world graphs is bounded by $O(\sqrt{n})$. As long as the maximal degree is less than $O(n)$ and there are relatively few outliers, the level of fine-grained global work distribution offered by the global collapse is likely not necessary.

IV. PERFORMANCE ANALYSIS AND DISCUSSION

A. Experimental Setup

We evaluate our algorithms on single nodes of three clusters, the Shannon and Compton systems at Sandia and the NSF Blue Waters system at the NCSA. A Shannon node has two Intel Xeon E5-2670 Sandy Bridge-EP processors with 128 GB main memory and an NVIDIA Tesla K40M GPU. The K40M GPU has 12 GB DDR5 memory, 2880 cores, and a peak memory bandwidth of 288 GB/s. Each GPU-enabled compute node of Blue Waters has one AMD 6276 Interlagos processor with 32 GB main memory and an NVIDIA Tesla K20X GPU. The K20X GPU has 6 GB DDR5 memory, 2688 cores, and a peak memory bandwidth of 250 GB/s. For GPU parallelism, Kokkos uses the CUDA programming model. We use Compton nodes for running our Kokkos and OpenMP implementations on its Intel Xeon processors and Intel Xeon Phi MIC coprocessors (Kokkos also utilizes OpenMP for multicore parallelism). Compton nodes are identical to Shannon nodes but only have 64 GB memory and house MICs containing 57 cores at 1.1 GHz with 6 GB memory. In all cases, the version of Kokkos used in our evaluation came from release 11.10.1 of Trilinos, we used icc and nvcc with the -O3 optimization option for compilation along with -arch=sm_35 when compiling for GPU.

We used several real small-world directed graphs that range in size from 5.1 million to 936 million edges for testing. These are listed in Table I. The graphs are from the SNAP database [21], the Koblenz Network Collection [20], and the University of Florida Sparse Matrix Collection [10]. We selected these graphs to represent a wide mix of graph sizes and topologies. Graph topology also has a strong influence on the performance of BFS and color propagation, while the number of total and nontrivial SCCs, as well as the size of the largest SCC, play an important role in determining performance of the SCC algorithm.

We report BFS and color propagation performance in terms of the Giga Traversed Edges per Second (GTEPS) metric, which normalizes running time to the total number of edges accessed (in billions). Note that our input graphs are directed and most of them have a large SCC. For each BFS execution, we track the total number of edges visited. Similarly, we count the number of vertex color and edge updates to determine overall performance for color propagation. We also run multiple iterations of both algorithms on all the target systems to reduce any variation in running time. In order to be consistent with BFS and color propagation results, we normalize SCC performance also by the number of edges and report an overall GEPS (Giga Edges per second) rate for each graph.

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<th>m</th>
<th>Degree (S)CCs</th>
<th># of SCCs</th>
<th># of Nontrivial SCCs</th>
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</tbody>
</table>

Table I: INFORMATION ABOUT TEST NETWORKS. COLUMNS ARE # VERTICES, # EDGES, AVERAGE AND MAX. DEGREE, # OF SCCS, # NUMBER OF NONTRIVIAL SCCS, AND SIZE OF THE LARGEST SCC.
For the Kokkos GPU approach, we fix thread queue sizes at 16, work chunks at 256 vertices per thread team for Hierarchical and Local Manhattan, and work chunks at 2048 per thread team for Baseline (vertex chunks) and Global Manhattan (edge chunks). For Xeon Phi and CPU, we used larger queues of size 1024 and work chunks of 2048. These values were selected for exhibiting the fastest performance across a range of values on our test suite. We will fully explore the performance impact of these algorithmic parameters in future work.

B. BFS Performance

Figure 1 gives the performance rates of the Kokkos-based BFS implementations on the Tesla K40M GPU. The baseline rate in the figure corresponds to performance with a vertex-based partitioning of the frontier array among thread teams. It is not a trivial implementation and our speedup numbers are conservative in that sense. We see consistent and significant speedups with three loop collapse strategies (H: hierarchical, MG: Manhattan Collapse using global memory, ML: Manhattan Collapse using GPU shared memory). Using H, MG, and ML, the speedups (geometric mean) over baseline are 1.82×, 1.82×, 3.85×, respectively, for the twelve graphs considered. The graphs are ordered in the figure in increasing order of average vertex degree, from DBpedia (3.9) to HV15R (140). Apart from a couple of anomalies, there is a reasonable correlation between average vertex degree in the graph and the BFS performance of the best variant (ML). Prior GPU graph algorithms work [17] has also made similar observations. However, one striking aspect is that the tuned variant can be more than an order-of-magnitude faster than the baseline, as we note for the Flickr graph. This is likely due to the large skewed degree distribution of Flickr, which severely limits the performance of the baseline approach. Other work has also noted the importance of parallel work balance with this particular graph [9].

Next, we summarize the impact of other optimization strategies discussed in Section III. Figure 2 gives BFS performance of the baseline and the ML variant again. In addition, we add optimizations in a structured manner to the code, starting with Baseline and finally getting to tuned ML (indicated by M+C+S+L in the figure). The intermediate steps are untuned Manhattan Collapse (indicated by Baseline+M), Manhattan Collapse with memory coalescing (M+C), Manhattan Collapse with memory coalescing and utilizing team-based scan procedures (M+C+S). The final step is the usage of temporary shared memory arrays for each thread team. It is interesting to note that Manhattan Collapse by itself does not provide much performance improvement. It is only after a methodical restructuring of the code, including optimizations such as coalescing and use of optimized scan primitives, that we are able to get the full benefit of the loop collapse optimization.

C. Color Propagation Performance

Figure 3 shows the performance of the loop collapse strategies on color propagation. Unlike BFS, the global Manhattan Collapse strategy does not consistently improve performance over the baseline. For a majority of the graphs, it is actually slower than baseline. Using H and M, though, the speedups (geometric mean) over baseline are 1.72× and 3.10×, respectively, for the twelve graphs considered. Performance of the best variant (ML) seems to well-correlated with average graph degree, with the exception of the synthetic RMAT2M and GNP2M graphs. We see the highest overall speedup over baseline (nearly 4.5×) with the Flickr graph. MG performs poorly on several instances due to the nature of iterative color propagation, which tends to have a long tail containing lots of low degree vertices. This effect is especially pronounced on the web graphs (uk, IndoChina), which tend to have long strings of singly-connected vertices. This hurts the performance of MG relative to ML in two ways. Firstly, the low average degree increases the amount of total transfer per team to and from the global prefix sum array. Secondly, the consistently low vertex degrees offer no benefit with regards to work partitioning among teams relative to the other approaches.
D. SCC Evaluation and Performance Portability

We finally evaluate performance of various SCC implementations. Recall that SCC algorithms use both BFS-like and color propagation-like loop nests, in addition to other graph topology-related work reduction heuristics [18], [33]. Our baseline Kokkos implementation for SCC is based on our prior Multistep [33] multicore parallel algorithm. Note that the BFS subroutine in Multistep is direction-optimizing, similar to [3], [7]. Thus we have also been able to express a fast heuristic work-reduction strategy in our Kokkos framework. We further improved the baseline approach using the ML and MG loop collapse strategies, and other GPU-specific optimizations. Figure 4 provides a cross-platform comparison of the various approaches on our test suite. Performance rates are indicated in terms of billions of edges per second. Our prior CPU Multistep implementation uses OpenMP, and can be compiled and run on x86 systems as well as Intel’s Xeon Phi coprocessors. We thus report these results on the Sandy Bridge-EP host processor and the Xeon Phi coprocessor, indicated as SNB and KNC in the figure. The OpenMP Multistep implementation is labeled OMP. The Kokkos baseline approach runs on all four platforms, and it is labeled as previous (B). Because Kokkos uses a single thread per team for the Xeon Phis and CPU, we only report performance for the MG variant of loop collapse on these systems, as the ML variant would default into an inefficient (B). We don’t consider hierarchical exploration due to the consistently superior performance of MG in previous experiments. We only include ML as a comparator for systems that can’t utilize MG. Any missing data points in both Figure 4 and Table II are due to memory limitations on the GPUs and Xeon Phi.

Consider the SNB column of the figure first. We observe that the OMP multistep performance varies between 0.1 to 5 GEPS, a nearly 50× variation. The anomalously-high performance on RMAT2M and GNP2M is due to the fact these synthetic graphs are relatively easy instances for the Multistep algorithm (there is only a single non-trivial SCC, so color propagation is never run). The graphs are ordered from top to bottom by average vertex degree. While OMP tends to do better than the Kokkos baseline on smaller graphs, for four of the twelve graphs, including the larger uk web crawls, the Kokkos baseline is in fact faster than the state-of-the-art OpenMP-based Multistep. MG, the algorithmic variant designed for GPUs, did not do as well as the baseline in SNB.

The Xeon Phi performance results are quite interesting,
The Kokkos baseline variant now consistently outperforms Multistep OMP. In comparison to parallel SNB performance, the absolute performance results on KNC are lower. However, note that these results were obtained with little or no parameter tuning for KNC. Besides three instances, MG again lags behind baseline. Thus we can conclude that the loop collapse strategies designed specifically for GPUs may not really lead to portable performance on KNC, without additional tuning.

The GPU SCC performance results are as expected. Notably, we could easily combine the Kokkos BFS and coloring implementations to create this SCC algorithm, and overall performance is quite favorable in comparison to the best parallel CPU implementation.

The original Multistep algorithm compiled with OpenMP and running on CPU shows the most consistent performance, followed by the GPU Kokkos ML algorithm running on GPU. Exploring cross-architectural and cross-implementation performance on each graph instance, we note different reasons for why a particular implementation is faster or slower. Multistep was designed to run on CPU with low diameter graphs and, as such, tends to dominate performance-wise on the smaller graphs, where there is lower available work parallelism, the graphs are less skewed, and the problems are generally easier to solve. This is apparent on the two simplest instances, the GNP and RMAT graphs. The GPU ML code arguably shows increasing relative performance with increasing problem difficulty, which is exemplified by DBpedia, the most skewed graph with the largest number of nontrivial SCCs. The additional parallelism for GPU ML across the adjacencies of the largest outliers in DBpedia makes a large relative impact. MG for both CPU and GPU does not show as good performance for DBpedia because, while there is improved parallelism across the largest adjacencies, DBpedia also has a very long tail of low degree vertices. This makes color propagation run very slow with MG due to all of the additional read and writes to the global prefix sums array.

In Table II, we list the SCC GEPS rate of the best-performing variant on each platform. The geometric mean...
of GEPS rates on each platform for SCC are also listed. Overall, K20X is \(1.67 \times\) faster than KNC for SCC. For BFS and Coloring, we compare performance of the best variant on K40M to the best-performing one on K20X. We observe that K40M is overall \(1.23 \times\) faster for BFS and \(1.09 \times\) faster for color propagation.

Although we note the present performance benefit of running on the multicore system relative to our manycore implementations, we make a few final points. Current trends in HPC indicate increased parallelism to become more prevalent, with an explicit distinction between a host CPU and coprocessor/GPU becoming blurred. This will make any present performance gaps between multicore and manycore codes less relevant. Additionally, more complex memory hierarchies and wider variation in architecture designs will make it a challenge to write algorithms that are efficient on future architectures. Hence, as we motivated previously, developing manycore techniques that exploit wide parallelism and varied memory hierarchies through reliance on a general framework and back-end for architecture-optimized parallelization will ease the burden on algorithm designers to design portable code for future systems.

### E. Comparisons to Prior Work

To the best of our knowledge, this is the first work on Kokkos-based graph computations targeting GPUs and Xeon Phi accelerators. For SCC, we performed direct comparisons with our prior OpenMP based Multistep method, as discussed in the previous subsection. In terms of mean performance rates, we believe that 0.43 GEPS for SCC, using a high-level framework such as Kokkos, is significant. Our mean BFS performance rate on the K40M is 1.74 GTEPS across all the test networks considered, and the best rate is 2.82 GTEPS for the RMAT2M network. Recently, Nguyen et al. [26] compare performance of several parallel graph analysis frameworks (Ligra, Galois, PowerGraph, GraphChi, and variants) for various graph analytics routines on a 40-core Intel Westmere-EX system. The best BFS performance reported, with Galois on the twitter40 graph, corresponds to a GTEPS rate of 2.1. Merrill et al. [23] report up to 3.3 GTEPS on the RMAT2M network for an optimized CUDA BFS implementation. For tuned CUDA-based SSSP approaches, Davidson et al. [9] report a peak performance rate of 0.35 GTEPS on an RMAT network and an NVIDIA GTX 680 (GK104). Thus, we believe that our approaches are competitive with the current state-of-the-art on multicore and manycore platforms.

### V. CONCLUSIONS

We used an algorithmic template that is common to lot of graph algorithms to express algorithms for strongly connected components, breadth first search and color propagation. This algorithmic template was used for portable manycore implementations using the Kokkos library and then optimized for architecture specific features like teams of threads and algorithmic features like loop-collapsing. We gave credence to the the efficacy of our approach by demonstrating the performance of a strongly connected components algorithm that is up to \(3.25 \times\) faster than a parallel CPU implementation.

We conclude with some commentary on questions posed in Section I. To answer questions (a) and (b), we advocate using simple array-based data structures and an iterative loop nest to perform graph computations, as shown in Algorithm 1. This simplifies transitioning from serial to multicore to manycore algorithms. The Local Manhattan Collapse optimization proved to be the biggest contributor to performance improvement over a baseline version. Given that most current and emerging real-world networks have skewed degree distributions, this would be the primary optimization strategy for graph analytics. The algorithms we studied in this paper use the abstraction: “given a large unordered set of vertices, how do we efficiently read and update attributes of the vertices and their adjacencies?” Using Kokkos, we see promising results for performance portability. The performance of our Baseline SCC algorithm on Xeon Phi is \(1.97 \times\) faster than an OpenMP-based implementation. Further, the multicore CPU algorithm based on Kokkos is only 30% slower than a hand-tuned OpenMP code. So we conclude as an answer to question (c), yes, performance-portable graph algorithms are possible using the previously identified libraries and optimization techniques.

We intend to apply this methodology to other graph analytics in future work. There are several research efforts on using both the host and the accelerator for graph analytic workloads [12], [13], [17], and this is another avenue for future work.

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