

Fast and high quality topology-aware task mapping

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Abstract

Considering the large number of processors and the size of the inter-connection networks on exascale-capable supercomputers, mapping concurrently executable and communicating tasks of an application is a complex problem that needs to be dealt with care. For parallel applications, the communication overhead can be a significant bottleneck on scalability. Topology-aware task-mapping methods that map the tasks to the processors (i.e., cores) by exploiting the underlying network information are very effective to avoid, or at worst bend, this limitation. We propose novel, efficient, and effective task mapping algorithms employing a graph model. The experiments show that the methods are faster than the existing approaches proposed for the same task, and on 4096 processors, the algorithms improve the communication hops and link contentions by 16% and 32%, respectively, on the average. In addition, they improve the average execution time of a parallel SpMV kernel and a communication-only application by 9% and 14%, respectively.

Keywords: Task mapping, communication graph, partitioning

1 Introduction

For parallel computing, task mapping has a significant impact on the performance, especially when supercomputers with hundreds of thousands or millions of processors shoulder the execution. It is usually the case that the communication pattern between the tasks has already been designed to minimize possible performance bottlenecks, such as high number of messages or communication volume, via tools such as graph and hypergraph partitioners, e.g., [PR96b, SS11, Kar11, ÇA99, DKUÇ14]. However, this effort alone is not sufficient, since the mapping-based metrics such as the maximum link congestion and the total number of hops the messages travel in the network can also be significant bottlenecks on the performance. This is especially true for today’s supercomputers with large-diameter interconnection networks, and concurrent and non-uniform job submissions yielding sparse and wide-spread processor allocations for parallel applications. There exist various studies in the literature which analyze the impact of task-to-processor mappings on the parallel performance [AYN⁺12, KKS06, BKK09, BNFC⁺12] and report significant speedups, e.g., 1.64X [GDS⁺06], just with an improved mapping.

There are two main research directions for mapping. The first one focuses on *block-based processor allocations*, e.g., the ones on IBM BlueGene [BKK09, GDS⁺06, ACG⁺04, YCM06]. The second direction focuses on *sparse allocations* in which the allocated processors are not contiguous within the network [DRL⁺14, HS11]. This direction is more general in the sense that sparse allocations have been used in various parallel systems, and the mapping algorithms based on this model can also be used for the block-based model. In this work, we follow the second direction.

The problem of finding an optimal task-to-processor mapping is NP-complete [HS11], and various heuristics have been proposed [BGKC10, BM91, CLZC11, CA95]. Many of these heuristics use graphs and related combinatorial structures to model the task interactions as well as the network topologies. For example, the open-source mapping library LibTopoMap [HS11] uses a task graph and network topology information. The task graph is first partitioned into the number of allocated nodes and various graph-based algorithms are used to map the tasks to the allocated processors. Other libraries such as JOSTLE [WC01] and Scotch [PR96b] also exist. These two libraries apply simultaneous partitioning and mapping of the task and topology graphs.

A good mapping algorithm must be able to provide high-quality task-to-processor mappings. In addition, it needs to be efficient in order not to intervene the supercomputer’s performance. In this work, we follow these two important criteria and propose novel, very efficient, refinement-based mapping algorithms and show that they can produce high-quality mappings w.r.t. the topology-related metrics such as the average link congestion and the total hops the messages take within the network. We compared the performance of the proposed algorithms with that of LibTopoMap and Scotch. The experiments on a supercomputer with a 3D torus network and 4096 processors show that the algorithms can improve the *weighted hop* and *maximum congestion* metrics (that

will be described below) by 16% and 32% on the average, respectively, compared to the default mapping. These metric improvements yield a 43% performance improvement on one case for a synthetic, communication-only application (Figure 4b–PATO) and a 23% improvement on the performance of a Sparse-Matrix Vector Multiplication (SpMV) kernel (Figure 5–METIS). Overall, with 4096 and 8192 processors, they improve the performance of a parallel SpMV kernel and a communication-only application by 9% and 14%, respectively (Table 1).

The organization of the paper is as follows. In Section 2, the background for the topology-aware task mapping is given. We also summarize the related work and the target architecture in this section. In Section 3, we present three mapping algorithms minimizing various metrics. Section 4 presents the experimental results, and Section 5 concludes the paper.

2 Background

Let $G_t = (V_t, E_t)$ be a directed MPI task graph, where V_t is the task set, and E_t is the edge set modeling task-to-task communications, i.e., $(t_1, t_2) \in E_t$ if and only if $t_1 \in V_t$ sends a message to $t_2 \in V_t$. Let $G_m = (V_m, E_m)$ be the network topology graph where V_m is the set of computing nodes equipped with many processors/cores, and E_m is the edge set modeling the physical communication links between the nodes, i.e., $(m_1, m_2) \in E_m$ if and only if $m_1 \in V_m$ and $m_2 \in V_m$ have a link in between. Let $V_a \subseteq V_m$ be the set of computing nodes reserved for the application. The topology-aware mapping problem can be defined as finding a mapping function ($\Gamma : V_t \rightarrow V_a$) that minimizes the parallel execution time. For a task set $S \subseteq V_t$, we use $\Gamma[S]$ to denote the node set to which the tasks in S are mapped to.

There are two well-received metrics to model the network communication overhead. The *total hop count* (TH), which is the total length of paths taken by communication packets, is a latency-based metric. The *maximum message congestion* (MMC), which is the maximum number of messages sent across a link. We assume that the messages are not split and sent through only a single path via static routing.

Let Γ be a given mapping where $m_1 = \Gamma(t_1)$ if and only if t_1 is assigned to one of node m_1 's processors. Then

$$\text{TH}(\Gamma) = \sum_{(t_1, t_2) \in E_t} \text{dilation}(t_1, t_2),$$

where $\text{dilation}(t_1, t_2) = \text{SPL}(\Gamma(t_1), \Gamma(t_2), G_m)$ which is the shortest-path length between $\Gamma(t_1) \in V_m$ and $\Gamma(t_2) \in V_m$.

Let the congestion on a network link $e \in E_m$ be

$$\text{Congestion}(e) = \sum_{(t_1, t_2) \in E_t} \text{inSP}(e, \Gamma(t_1), \Gamma(t_2), G_m), \quad (1)$$

where, $\text{inSP} = 1$ if and only if e is on the shortest path between $\Gamma(t_1)$ and $\Gamma(t_2)$, and 0 otherwise. Therefore, a link's congestion is equal to the number of

messages that passes through it. Hence, the maximum message congestion is

$$\text{MMC}(\Gamma) = \max_{e \in E_m} \{\text{Congestion}(e)\}.$$

The above metrics assume unit communication costs and link capacities. In order to handle heterogeneous costs and bandwidths, the task and topology graph models are extended as follows. Each edge $e' \in E_t$ in the task graph is associated with a cost $c(e')$ that corresponds to the communication volume sent/received between the tasks. Similarly, each link $e \in E_m$ in the topology graph is associated with a communication capacity $bw(e)$ that corresponds to the link bandwidth. Moreover, for further heterogeneity support, each node $m \in V_m$ is associated with a computation capacity $w(m)$ that corresponds to the number of available (allocated) processors in m . All the nodes that are not in V_a have zero capacity. Based on these attributes on the edges, the *weighted hop* (WH) metric, i.e., the total number of hops taken by each packet, is

$$\text{WH}(\Gamma) = \sum_{(t_1, t_2) \in E_t} \text{dilation}(t_1, t_2) c(t_1, t_2).$$

We define the *volume congestion* (VC) of a link $e \in E_m$ as

$$\text{VC}(e) = \frac{\sum_{(t_1, t_2) \in E_t} \text{inSP}(e, \Gamma(t_1), \Gamma(t_2), G_m) c(t_1, t_2)}{bw(\Gamma(t_1), \Gamma(t_2))}$$

which is the ratio between the overall volume of the communication passing from e and its bandwidth. Then, the *max volume congestion* (MC) metric is the congestion of the bottleneck link in the network with the maximum VC value.

A communication-bounded parallel application can be either latency-bounded or bandwidth-bounded depending on the communication pattern. For example, applications with frequent communication steps and small messages are likely to be latency-bounded, while those with larger messages are likely to be bandwidth-bounded. For latency-bounded applications, TH and WH metrics can better correlate with the communication overhead. On the contrary, MMC and MC are better choices for the bandwidth-bounded applications. Depending on the communication pattern, it might be better to minimize MC (MMC) in the expense of an increase on WH (TH), or vice versa. However, it is indeed difficult to find a consensus between these metrics. Therefore, here we define *average message congestion* (AMC) and *average congestion* (AC) metrics to account for both number of hops and congestion. Let E_m^t be the set of the links that are used throughout the execution of the parallel application. The message and volume based average link congestions over the used links can be defined as

$$\text{AMC}(\Gamma) = \frac{\sum_{e \in E_m} \text{Congestion}(e)}{|E_m^t|},$$

$$\text{AC}(\Gamma) = \frac{\sum_{e \in E_m} \text{VC}(e)}{|E_m^t|}$$

Since $\text{TH} = \sum_{e \in E_m} \text{Congestion}(e)$, AMC is the ratio of TH to the total number of links used during the application. Similarly, AC is related to the ratio of WH to the number of links used during the execution (they are equal only when the communication links have unit bandwidths).

The communication during the execution is a real-time process that can easily be affected by many outside factors (e.g., network traffic and overhead from competing jobs). Hence, the theoretical metrics given above can only approximate the actual overhead. In Section 4.5, we will discuss and evaluate the interplay of these metrics on the performance.

2.1 Related Work

Existing mapping algorithms can be divided into two classes: *single-phase* and *two-phase* methods. The algorithms in the former class perform simultaneous partitioning and mapping using the task and topology graphs, whereas those in the latter partition the task graph in the first phase and map the parts to the processors in the second one.

Pellegrini and Roman [PR96a] proposed a single-phase recursive-bipartitioning algorithm for various topologies. Wallshaw and Cross [WC01] proposed a multi-level partitioning algorithm which performs mapping in the initial partitioning and refinement phases.

The two-phase mapping methods make an abstraction of the partitioning phase and work with a pre-partitioned task set. These studies can be divided into two based on the task-dependency model and network topology. The first set adapts geometric task-interaction models, e.g., [BGL⁺12, BJI⁺14] on IBM’s BlueGene systems with block-based node allocations and on sparse allocations [DRL⁺14]. Still, most of the work focuses on the connectivity-based models, specifically graph models. This problem is shown to be NP-complete [HS11, Bok81], and many heuristics exist in the literature, e.g., [BGKC10, BM91, CLZC11, CA95]. In this work, we investigate a two-phase graph-based approach.

2.2 The architecture

Six of today’s ten fastest supercomputers have torus networks (one 3D, four 5D, one 6D torus in Top500, June 2014). In this work, our target architecture is NERSC’s Hopper supercomputer with Cray XE6’s 3D torus. The network has Gemini routers directly connected to 2 computing nodes. Each router is associated with a coordinate on x , y , and z dimensions, and connected to all six neighbor routers. The torus network provides wrap-arounds, and the messages between the nodes are statically routed following the shortest paths. The network links have different bandwidth values on various dimensions.

On Cray systems, the scheduler allocates a non-contiguous set of nodes for each job [ATW⁺11]. Although it attempts to assign nearby nodes, no locality guarantee is provided. The topology information, e.g., the routers’ coordinates and connections, as well as link bandwidths, can be captured using system

calls (`rtr`), and a static topology graph can be obtained. Then during runtime, each MPI process can obtain its node id, and the vertices in the topology graph can be associated with the computational units.

In Hopper, the network latencies for the nearest and farthest node pairs are $1.27\mu\text{s}$ and $3.88\mu\text{s}$, respectively. The link bandwidths vary from 4.68 to 9.38 GB/sec. Ideally, reducing the hop counts between the communicating tasks lowers the overhead. But when there are many communicating tasks, a link can be congested due to the communication pattern, which might cause communication stalls and harm the performance. Still, thanks to static routing, the congestion can be measured and optimized accurately.

3 Fast and High Quality Mapping Algorithms

We propose three mapping algorithms to minimize WH and MC. Here we will describe these algorithms. Their adaptation for TH and MMC is trivial. Among them, the ones that minimize WH can be applied to various topologies, whereas those minimizing MC require static routing.

3.1 A Greedy mapping algorithm

The first algorithm *Greedy Mapping* given in Algorithm 1 finds a mapping $\Gamma : V_t \rightarrow V_m$ to minimize WH. It uses the task graph G_t and the topology graph G_m . The algorithm is similar to *greedy graph growing*, and initially maps N_{BFS} seed task vertices to the nodes. It assumes a symmetric G_t while finding the neighbors of a given task since WH is an undirected metric, i.e., the number of hops between m_1 and m_2 is the same regardless of direction.

Throughout the algorithm, the total connectivity of each task to the mapped ones are stored in a heap *conn*. The algorithm first maps the task t_{MSRV} with the maximum send-receive communication volume to an arbitrary node. Until all tasks are mapped, the algorithm gets an unmapped task from *conn* after all the N_{BFS} seeds are mapped. Otherwise, the farthest task to the set of mapped tasks is found by a breadth-first search (BFS) on G_t where all the mapped tasks are assumed to be at level 0 of the BFS. Ties are broken in the favor of the task with a higher communication volume. If G_t is disconnected, a task with the maximum communication volume from one of the disconnected components is chosen. Once t_{best} is found, its best node is obtained by `GETBESTNODE`. If t_{best} is connected to none of the mapped tasks, `GETBESTNODE` performs a BFS on G_m to **return** one of the farthest allocated nodes to the set of the non-empty nodes, i.e., the ones with a mapped task. Otherwise, if t_{best} is connected to at least one of the mapped tasks, a BFS on G_m is performed from the nodes to whom one of the $neighbor(t_{best})$ is mapped (again assuming these nodes are at level 0). As an early exit mechanism, a BFS stops when the empty nodes (without a mapped task) are found at a BFS level. Then among these empty nodes, the one with the minimum WH overhead is returned. Therefore, the algorithm performs multiple BFS executions on G_t and G_m .

Algorithm 1: Greedy Mapping

Data: $G_t = (V_t, E_t), G_m = (V_m, E_m)$: task and topology graphs, N_{BFS} : # vertices to be initially mapped

```
connt ← 0 for each  $t \in V_t$  ▶ initialize the max-heap
 $\Gamma[t] \leftarrow -1$  for each  $t \in V_t$  ▶ initialize the mapping
▶ Find the task with MSRV
 $t_0 \leftarrow t_{MSRV}$ 
▶ Map  $t_0$  to an arbitrary node
 $\Gamma[t_0] \leftarrow m_0$ 
▶ Update connectivity for the tasks in  $neighbor(t_0)$ 
for each  $t_n$  in  $neighbor(t_0)$  do
  |  $conn.update(t_n, c(t_0, t_n))$ 
while there is an unmapped  $t$  do
  | if number of mapped tasks  $< N_{BFS}$  then
  | |  $t_{best} \leftarrow$  the farthest unmapped task ▶ found by BFS
  | else
  | |  $t_{best} \leftarrow conn.pop()$  ▶ the one with maximum conn.
1 |  $m_{best} \leftarrow GETBESTNODE(t_{best}, G_m, G_t, \Gamma, conn)$ 
  |  $\Gamma[t_{best}] \leftarrow m_{best}$ 
  | for each  $t_n$  in  $neighbor(t_{best})$  do
2 | |  $conn.update(t_n, c(t_{best}, t_n))$ 
```

For simplicity, the description above assumes one-to-one task-to-node mapping, i.e., $|V_t| = |V_a|$. In reality, each node has multiple processors, so multiple tasks can be assigned to a single node. These cases can be addressed by using the computation loads and capacities, and modifying GETBESTNODE so that it returns only a node with some free capacity. Another common solution is using traditional graph partitioning as a preprocessing step to reduce the number of tasks to the number of the allocated nodes while minimizing the edge-cut [HS11]. We follow this approach and use METIS [Kar11] to partition G_t into $|V_a|$ nodes, where the target part weights are the number of available processors on each node in V_a . Since graph partitioning algorithms do not always obtain a perfect balance, as a post processing, we fix the balance with a small sacrifice on the edge-cut metric via a single Fiduccia-Mattheyses (FM) iteration [FM82]. When the number of processors in the nodes are not uniform, we map the groups of tasks with different weights at the beginning of the greedy mapping since their nodes are almost decided due their uniqueness.

In the algorithm, N_{BFS} controls the number of initial seed mappings. A large N_{BFS} distributes the loosely connected components of the task graph to the nodes that are farther from each other. However, this will not work well for the task graphs with a low diameter. In our implementation, we use $N_{BFS} \in \{0, 1\}$ to generate two different mappings and return the one with the lower WH.

The complexity of the algorithm is dominated by the operations at lines 1

and 2. Each update of the heap (line 2) takes $\mathcal{O}(\log |V_t|)$ time, and this line is executed at most $|E_t|$ times, yielding $\mathcal{O}(|E_t| \log |V_t|)$. The BFS operation in GETBESTNODE has $\mathcal{O}(|E_m|)$ cost, yielding an overall complexity of $\mathcal{O}(|V_t||E_m|)$. For a task t_{best} and a candidate node at the last level of the BFS performed in GETBESTNODE, the cost of computing the change on WH is proportional to the number of edges of t_{best} (the hop count between two arbitrary nodes can be found in $\mathcal{O}(1)$, since G_m 's are regular graphs). Since there are at most $|V_a|=|V_t|$ candidate nodes and $|E_t|$ edges, the complexity of this part throughout the algorithm is $\mathcal{O}(|V_t||E_t|)$. Therefore, the complexity of Algorithm 1 is $\mathcal{O}(|V_t|(|E_m| + |E_t|))$ —in practice it runs faster thanks to the early exits in GETBESTNODE BFSs.

3.2 A Refinement algorithm for the weighted hop

Algorithm 1 is our main algorithm and the other two will refine its mapping. Even after its execution, it is possible to improve WH via further refinement. We have implemented a Kernighan-Lin [KL70] type algorithm which uses “task swaps” to refine WH (Algorithm 2). It gets a Γ , G_t , and G_m as input and modifies Γ to lower the WH metric. Similar to greedy mapping, for simplicity, Algorithm 2 assumes that G_t is symmetric and Γ is a one-to-one mapping between the tasks and nodes.

Algorithm 2: WH Refinement

Data: $G_t = (V_t, E_t)$, $G_m = (V_m, E_m)$, Γ , Δ

- ▶ compute the current WH for Γ
- WH \leftarrow CALCULATEWEIGHTEDHOPS(G_t , G_m , Γ)
- while** WH is improved **do**
 - ▶ compute WH incurred by each task
 - ▶ place the tasks in a max-heap *whHeap*
 - 1 **for** t in V_t **do**
 - $wh_t \leftarrow$ TASKWHOPS(t , G_t , G_m , Γ)
 - whHeap.insert*(t , wh_t)
 - 2 **while** *whHeap* is not empty **do**
 - 3 $t_{wh} \leftarrow$ *whHeap.pop*()
 - for** the first Δ nodes $m \in V_a$ visited in the order of the BFS from $\Gamma[\text{neighbor}(t_{wh})]$ **do**
 - $t \leftarrow$ the task mapped to m
 - 4 **if** swapping t_{wh} and t improves WH **then**
 - $\Gamma[t] \leftarrow \Gamma[t_{wh}]$
 - $\Gamma[t_{wh}] \leftarrow m$
 - 5 Update *whHeap* for neighbors of t_{wh}
 - 6 Update *whHeap* for neighbors of t
 - break**

The algorithm selects a pair of task vertices and swaps them to improve WH. The first task t_{wh} is chosen using a max-heap, *whHeap*, which initially

organizes the tasks w.r.t. the WH amount they incur computed by a function TASKWHOPS function (line 1). Hence, t_{wh} is the task individually responsible for the largest WH. Choosing the second task for the swap operation is more complicated; a naive approach that considers to swap t_{wh} with all the other tasks requires $\mathcal{O}(|V_t|^2)$ comparisons. In order to avoid this cost, we have implemented a BFS-based task-selection algorithm. A simple observation is that to reduce WH, t_{wh} needs to move closer to its neighbor tasks. Therefore, we perform a BFS on G_m starting from the nodes which have a neighbor of t_{wh} , i.e., the nodes in $\Gamma[neighbor(t_{wh})]$ (these are the level 0 nodes of BFS). Whenever a V_a node with a the task t is found, the WH value after the potential $\Gamma[t_{wh}] \leftrightarrow \Gamma[t]$ swap operation is computed. The actual swap is performed as soon as this computation reveals an improvement on WH. Since the likelihood of a WH improvement decreases when we go deeper on the BFS tree, we use an early exit mechanism to avoid a full BFS traversal of G_m . Here we give an example to clarify the statement, assume $c(e) = 1 \forall e \in E_t$. If the maximum hop count between $\Gamma[t_{wh}]$ and $\Gamma[neighbor(t_{wh})]$ is d then when t_{wh} is moved to a node after the BFS level d , WH incurred by t_{wh} cannot be improved. Furthermore, when we go deeper in the BFS, t_{wh} 's incurred WH value will increase. Even in this case, the overall WH may still be improved due to the reduction of WH incurred by the second task t . However, this is less likely to happen considering $\Gamma[t]$ is handpicked for t_{wh} but $\Gamma[t_{wh}]$ is only a random node for t . The early exit mechanism reduces the number of considered swap operations that are unlikely to improve WH. In Algorithm 2, a parameter Δ is used as an upper bound on this number. If Δ operations are checked for t_{wh} , the algorithm continues with the next *whHeap* vertex. A refinement pass is completed when *whHeap* is empty, and the next pass is performed only if there is an improvement in the previous pass.

The complexity of the loop at line 1 is $\mathcal{O}(|V_t| \log |V_t| + |E_t|)$. The loop at line 2 iterates $|V_t|$ times. The complexity of the BFS operation at line 3 is $\mathcal{O}(|E_m|)$ and it is also performed $|V_t|$ times. The complexity of the swap operation and the calculation of new WH at line 4 is proportional to the total number of edges of t_{wh} and t for each candidate node. Since there are at most Δ candidate nodes for each BFS, the complexity of line 4 for each pass becomes $\mathcal{O}(\Delta |E_t|)$. Lines 5 and 6 are executed at most once for each vertex and during a single pass, their total cost is $\mathcal{O}(|E_t| \log |V_t|)$. Therefore, the overall complexity of the algorithm becomes, $\mathcal{O}(|V_t| \log |V_t| + |E_t| + |E_m| |V_t| + \Delta |E_t| + |E_t| \log |V_t|)$. The most dominant factor is the complexity of the BFS operations which is $\mathcal{O}(|E_m| |V_t|)$. Fortunately, the practical execution time is very low, since we stop after $\Delta=8$ swap candidates. We experimented with other exit mechanisms based on the maximum BFS level instead of the number of swap operations, and the preliminary experiments favored the approach described above. Furthermore, we observed that most of the improvement in WH is obtained after only a few passes. Hence, in order to be more efficient, we perform a pass only if WH is improved more than 0.5% in the previous one.

Similar to that of Algorithm 1, the description above assumes a one-to-one task-to-node mapping and performs the refinement on the node level, i.e., by swapping the vertices representing a group of tasks. With slight modifications, it

can perform the refinement on the finer level task vertices or in a multilevel fashion from coarser to finer levels. In our experiments we choose to perform only on the coarser task graphs we obtained after METIS, since with WH-improving swap operations on the finer level, the total internode communication volume can also increase and the performance may decrease. Although this increase can also be tracked during the refinement, we do not want to sacrifice from the efficiency.

3.3 A Refinement algorithm for the maximum congestion

Although Algorithms 1 and 2 significantly improve WH, they can negatively affect MC or MMC, and this can degrade the performance especially for the bandwidth-bounded applications. Therefore, we propose another refinement algorithm (Algorithm 3), which improves the MC metric with minimal WH damage (adapting this algorithm to refine MMC is trivial). The algorithm can accurately model and minimize MC for the interconnection networks with static routing. We will discuss the required enhancements for the dynamic-routing networks.

Algorithm 3: MC Refinement

Data: $G_t = (V_t, E_t), G_m = (V_m, E_m), \Gamma, \Delta$
▶ calculate initial max and average congestions
 $MC, AC \leftarrow calculateCongestion(G_t, G_m, \Gamma)$
▶ initialize the link congestion heap
▶ store the tasks whose messages goes through links
 $congHeap, commTasks \leftarrow INITCONG(G_t, G_m, \Gamma)$
while MC or AC is improved **do**
1 $e_{mc} \leftarrow congHeap.pop()$
2 **for** $t_{mc} \in commTasks[e_{mc}]$ **do**
3 **for** the first Δ nodes $m \in V_a$ visited in the order of the BFS from
 $\Gamma[neighbor(t_{mc})]$ **do**
 $t \leftarrow$ the task mapped to m
 if swapping t_{mc} and t improves MC or AC **then**
 $\Gamma[t] \leftarrow \Gamma[t_{wh}]$
 $\Gamma[t_{wh}] \leftarrow m$
 Update $congHeap$ for t_{mc} and t edges
 Update $commTasks$ for t_{mc} and t edges
 goto line 1

The algorithm gets a Γ, G_m , and G_t and modifies Γ to find mapping with a better congestion. First, it computes the initial congestion of Γ , and initializes the $congHeap$ using an INITCONG function. This max-heap stores the topology graph edges w.r.t. their congestion values. The algorithm also initializes $commTasks$, that is used to query the tasks whose messages go through link e , i.e., $commTasks[e]$. Since, a message can go at most D (network diameter) hops, the maximum size of $commTasks$ becomes $|E_t|D$, which is manageable

since D is not a large number.

After the initialization, the algorithm finds the most congested link e_{mc} . Then for each $t_{mc} \in commTasks[e_{mc}]$, the node to which t_{mc} will be moved is sought via BFS traversals on G_m starting from the nodes $\Gamma[neighbor(t_{mc})]$. The second task t to swap is chosen from the tasks of the V_a nodes traversed during the BFSs. This BFS order is important to have a minimal damage on WH. For each such candidate node, a virtual swap operation is performed to compute new MC and AC values. As soon as an improvement is detected, the actual swap operation is performed, and the execution continues with the next congested link. Whenever a vertex is moved, updates on the *congHeap* and *commTasks* are performed for all the incoming and outgoing edges of t_{mc} and t in E_t . If there is no improvement after $\Delta=8$ trials, the early exit mechanism terminates the inner **for** loop. Then, the next task in *commTasks*[e_{mc}] is chosen and the search restarts. If no improvement is found for the most congested link the algorithm stops. This algorithm can be applied both to coarser and finer task graphs. However, we only apply it on the coarser graph due to the reasons explained before.

With static routing, a message route can be found in $\mathcal{O}(D)$ time. The congestions of all the links can be calculated in $\mathcal{O}(D|E_t|)$ time, and the cost of initializing *congHeap* is $\mathcal{O}(|E_m| \log |E_m|)$. A task insertion to a *commTasks* set (implemented as a red-black binary tree using `std::set` in C++) can be done in $\mathcal{O}(\log |V_t|)$. Since each message (an edge in E_t) can pass through at most D links, the complexity of *commTasks*'s initialization is $\mathcal{O}(D|E_t| \log |V_t|)$. Therefore the initialization phase has a complexity of $\mathcal{O}(D|E_t| \log |V_t| + |E_m| \log |E_m|)$. A refinement pass starts at line 2. The main **for** loop iterates at most $|V_t|$ times and the complexity of a BFS at line 3 is $\mathcal{O}(|E_m|)$. Hence, the overall BFS complexity in a pass is $\mathcal{O}(|V_t||E_m|)$. For each candidate swap operation, we compute MC and AC by temporarily updating *congHeap* where an update costs $\log |E_m|$ and a candidate swap requires at most D updates for each of the t_{mc} and t edges. Since we consider at most Δ swap operations for a t_{mc} which can be any vertex in V_t , the cost of MC and AC computation is $\mathcal{O}(E_t \Delta D \log |E_m|)$ for each e_{mc} . Once an improvement is found, the data structures *congHeap* and *commTasks* are updated and this happens only once per pass (a **while** loop iteration). Hence, the overall complexity of a pass is dominated by the BFS and MC-AC computations. Therefore the overall complexity of a pass becomes $\mathcal{O}((E_t \Delta D \log |E_m|) + (|V_t||E_m|))$.

Algorithm 3 accurately sees the maximum congestion on a static-routing network. For the networks with dynamic routing, an approximate refinement algorithm with a similar structure can be used. For example, the bandwidth on the Blue Gene/Q and Blue Gene/P can be maximized by placing the heavily communicating tasks to the diagonals of the torus [BGL⁺12, BJI⁺14].

4 Experiments

In order to evaluate the quality of the mapping algorithms, we conducted various experiments on two irregular applications, an SpMV kernel and a synthetically generated application. The proposed methods are implemented in the UMPA framework. The U_G and U_{WH} variants minimize WH using Algorithms 1 and 2, and U_{MC} and U_{MMC} minimizes MC and MMC, respectively using Algorithm 3 (we do not give the results for TH variant as they are very close to those of U_G and U_{WH}). These mapping methods are compared against the default MPI mapping (SMP-STYLE) in Hopper (DEF), the mapping provided by SCOTCH (SMAP, version 5.1.0 as the newer one does not support sparse allocations) [PR96b], and the ones provided by LibTopoMap (TMAP) [HS11].

We selected 25 matrices from University of Florida (UFL) sparse matrix collection, belonging to 9 different classes (the list is at the supplementary page: <http://web.cse.ohio-state.edu/~deveci/umpamap>). We used 7 graph and hypergraph partitioners to partition these matrices: SCOTCH [PR96b], KAFFPA (KAHIP) [SS11], METIS [Kar11], PATOH [ÇA99], and UMPA [DKUÇ14]. A summary of the partitioning results are given in Section 4.1. MPI task communication graphs corresponding to these partitions are created and mapped to real processor allocations in Hopper. The analysis of the metrics and algorithm efficiency is presented in Section 4.2. Section 4.3 analyzes the impact of the mapping algorithms on the communication time, whereas Section 4.4 evaluates the performance improvements for a Trilinos SpMV kernel [HBH⁺05]. We analyze the impact of the partitioning and mapping metrics on the parallel performance in Section 4.5.

4.1 Partitioning results

The matrices are first converted to a column-net hypergraph model, i.e., the rows represent the tasks with loads proportional to their number of non-zeros. The columns represent sets of data communications where each message has a unit communication costs. On these matrices we perform 1D row-wise partitioning for 1024, 2048, 4096, 8192 and 16384 parts (we only use 19 matrices for 16384 parts; balanced partitions were not feasible for the remaining 6). The graph partitioners, SCOTCH and KAFFPA, are run to minimize the edge-cut, and METIS and PATOH are run to minimize the total communication volume TV. Being a multi-objective partitioner, UMPA is used with different metrics: UMP_{MV} minimizing maximum send volume (MSV) and TV; UMP_{MM} minimizing maximum number of sent messages (MSM), total number of messages (TM) and TV; UMP_{TM} minimizing TM and TV; as their primary, secondary, and tertiary objectives, respectively [DKUÇ14]. All the partitioners are run with their default parameters.

Figure 1 shows the mean metric values normalized with that metric value of PATOH. Overall, all the tools obtain similar results, but edge-cut minimizing ones, SCOTCH and KAFFPA, obtain a slightly worse communication volume quality. For the MSV metric, UMP_{MV} has the best results, e.g., it obtains a

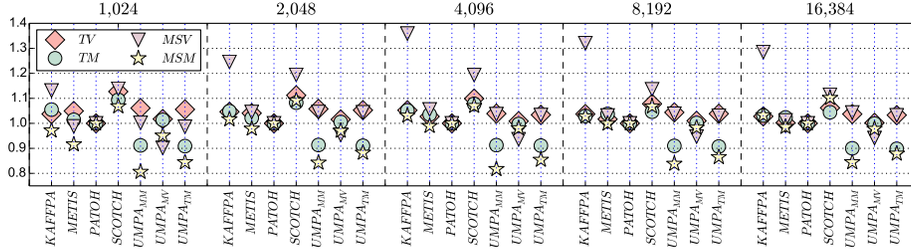


Figure 1: Geometric means of the partition metrics w.r.t PATOH for the corresponding part number.

5–10% better average MSV value w.r.t. PATOH which obtains the best results for TV. For the message metrics, UMPA_{MM} obtained a 16–19% better MSM value, and UMPA_{TM} obtained a 9–10% better TM value. These numbers are not given here to compare the partitioners since the experiment is not designed for that purpose. We want to better understand the impact of the partitioning and mapping metrics on the execution time.

4.2 Mappings on Hopper

Here we evaluate the mapping metric results on Hopper, which has a 3D Torus network and 24 processors per node. Even though the proposed algorithms do not have constraints on the number of processors, we tested them on numbers that are powers of two. Using all the processors in a node results in non-uniform processor allocations per node (since 24 does not divide 1024), in which case we experienced a few failing algorithms in LibTopoMap. Therefore, we used 16 processors per node (4 processors per NUMA domain).

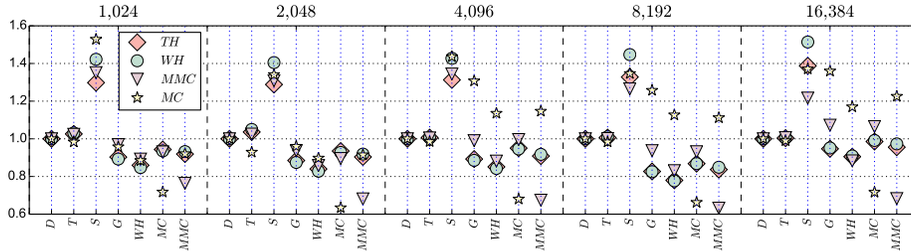


Figure 2: Mean metric values of the algorithms on $G_t^{\text{PATO}_H}$ graphs normalized w.r.t. those of DEF. The numbers at the top denote the number of the processors, and the letters at the bottom correspond to the mapping algorithms DEF, TMAP, SMAP, U_G , U_{WH} , U_{MC} , U_{MMC} , respectively.

We create directed task graphs by running all the partitioners on each matrix; for each graph and part number, we have 7 MPI task graphs. We will refer a task graph as G_t^X when the part vector obtained from the partitioner X is used to create it. The mapping algorithms are then used to map these graphs

to 5 different Hopper processor allocations. Figure 2 shows the average metric values of all mapping algorithms normalized to those of the default mapping on $G_t^{\text{PATO}^{\text{H}}}$ graphs. Almost all algorithms have their best WH and MC values on $G_t^{\text{PATO}^{\text{H}}}$, and best TH and MMC values on $G_t^{\text{UMPA}^{\text{TM}}}$. The results are expected for WH and TH, since WH is closely related to the communication volume, and TH is related to total number of messages. On the other hand, it is expected to have better MC and MMC values on the task graphs with lower MSV and MSM values, respectively. However, in our experiments, we see a better correlation of these metrics with TV and TM.

In Fig. 2, the DEF mapping obtains already good results on WH and TH. This is due to the part ID assignment in recursive-bisection-based partitioners and the placement mechanism in Hopper: the partitioner puts highly communicating tasks to the parts with closer IDs. On the machine side, Hopper places the consecutive MPI ranks within a single node, then it moves to the closer nodes using space filling curves. Therefore, highly communicating consecutive MPI ranks are placed fairly close to each other. However, there is still room for improvement when we exploit the actual task communication requirements. For example, U_G obtains 5–18% and 5–17% better values on WH and TH, respectively.

Metric improvements on more sparse allocations (with less number of processors) are higher: U_G significantly reduces WH and TH, and U_{WH} improves them by another 4–5%. Also the variants that improve the WH metric also improve MC and MMC. For example, U_G (U_{WH}) improves MC by 4% (10–12%) on 1024 and 2048 processors. However, when the number of processors is high, they increase the MC metric by 13–36% when the number of parts is high. Still, they reduce MMC, TH, and WH except U_G on 16,384 processors. Also, U_{MC} significantly reduces (27–37%) the MC metric for all cases and have 1–13% improvement on WH and TH. Similarly, U_{MMC} reduces MMC by 24–37% with small increases on TH and WH.

LibTopoMap provides 6 algorithms and the best one in our experiments employs recursive graph bi-partitioning. Here we only present the best variant’s performance (TMAP or T). The primary metric for LibTopoMap is MC. If TMAP’s MC value is not smaller than the DEF mapping, it returns the DEF mapping. Overall, TMAP improves MC by 1–7% with 1–5% increase on the other metrics. On the other hand, SMAP’s results are worse than DEF mappings for most of the cases.

Figure 3 presents the (geometric) mean mapping times of the algorithms. The times of the SMAP, U_G and U_{WH} are the lowest, and they are followed by U_{MC} and U_{MMC} . TMAP’s execution time is more than the other methods and it is 1.3–2.6 times slower than the slowest UMPA variant.

4.3 Communication-only experiments

In task mapping, the communication is usually modeled by assuming all the messages are transferred at once. However, this may not be the case in practice: load imbalance can delay some transfers, and applications might be using com-

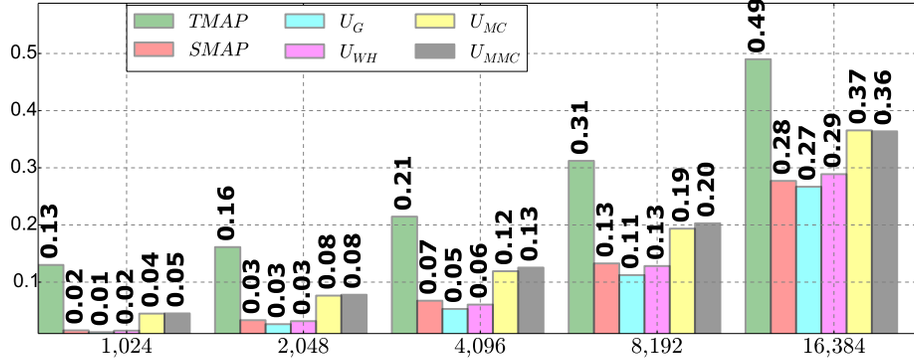
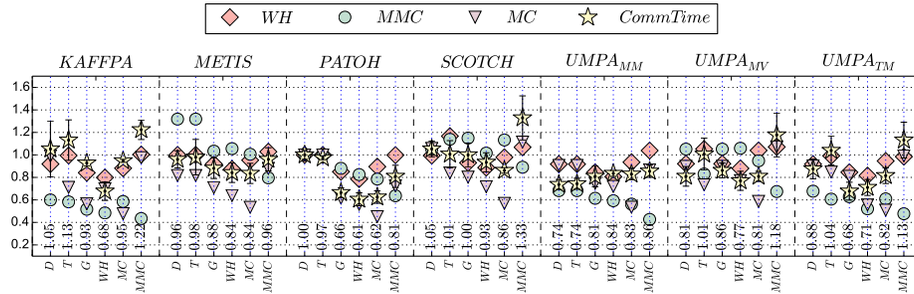
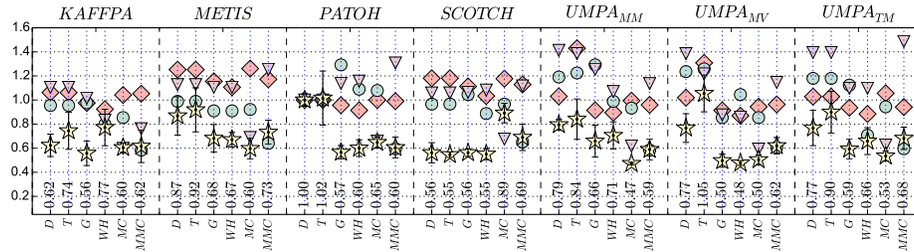


Figure 3: (Geometric) mean execution times of different mapping algorithms on PATOH partitions. The time of U_{WH} , U_{MC} , and U_{MMC} includes U_G time, as they run on top of it.



(a) cage15, #procs = 4096, scale = 4K



(b) rgg, #procs = 4096, scale = 256K

Figure 4: Average execution times and metrics for pure communication-based applications generated from `cage15` and `rgg`: the numbers at the bottom are the normalized execution times w.r.t. DEF mapping on G_t^{PATOH} . The partitioner names are given at the top, and the names at the bottom are the mapping algorithms, as given in Figure 2.

mon techniques such as communication-computation overlap to hide the latency. Hence, improvements due to mapping may not be visible on an application’s execution time. Here, to limit the impact of these factors, we generate irregular, communication-only applications based on the SpMV communication patterns of the two largest matrices in our dataset: `cage15` and `rgg_n.2.23_s0` (in short `rgg`). In this SpMV-like executions, no computation is performed and all the transfers are initialized at the same time where each processor exactly follows the pattern in the corresponding communication graph. To make the improvements more visible and reduce the noise, we scale the transfer volume, i.e., message sizes, by using the factors $4K$ and $256K$ for `cage15` and `rgg`, respectively. The experiment is performed with 4096 processors. Each mapping algorithm is run with the 7 communication graphs (one per partitioner), and for each mapping, the execution is repeated 5 times to reduce the noise on the time. Figure 4 shows the normalized mean execution times with standard deviations and the metric values normalized w.r.t. those of DEF mapping on G_t^{PATOH} . Although we run SMAP in this experiment, its communication time is worse than the others. For the sake of figure clarity, we exclude it from the figures. In addition, we do not report TH metric as it is highly correlated with WH metric. Results with 8192 processors and a different sparse allocation can be found at <http://web.cse.ohio-state.edu/~deveci/umpamap>.

Figure 4a shows the results for `cage15` communication graphs. The overall execution time correlates well with WH. In most cases, U_G and U_{WH} improve WH, MC, and the communication time w.r.t. DEF with a few exceptions. For example, on $G_t^{\text{UMPA}_{MM}}$, WH minimizing algorithms (Algorithms 1 and 2) improve all three metrics at the same time. However, the execution times with these mappings slightly increase. U_{WH} obtains much better WH, MC, and execution times compared to U_G . On G_t^{KAFFPA} , it improves WH in the expense of increasing MC but the execution time significantly reduces. Overall, U_G and U_{WH} improve the performance up to 34% and 39% w.r.t. the DEF. For all graphs, U_{MC} obtains the best MC values and it usually improves the performance w.r.t. DEF. Moreover, it obtains the best execution time on G_t^{SCOTCH} , which has the highest TV w.r.t other partitioners on this graph. Among the UMPA algorithms, U_{MMC} obtains the worst execution times although it always significantly reduces MMC. This is expected; since the message sizes are scaled, the executions have a high TV value and the volume-related metrics are likely to be the bottleneck rather than the message-related ones. TMAP can not improve the results of the DEF in some of cases, e.g., G_t^{METIS} , G_t^{PATOH} , and $G_t^{\text{UMPA}_{MM}}$, and returns the default mapping (the times vary 2–3% due to noise). Lastly, although DEF obtains the best mean execution time on UMPA graphs, overall, the best times are obtained on PATOH graphs with U_{WH} and U_{MC} with 39% and 38% improvement, respectively.

Figure 4b shows the results for `rgg` communication graphs. The proposed mapping algorithms improve the execution time for all the graphs except for G_t^{SCOTCH} . Similar to `cage15` experiments, the best performance is obtained by U_G , U_{MC} and U_{WH} . The best execution time is achieved by U_{MC} on $G_t^{\text{UMPA}_{MM}}$ with a 32% improvement w.r.t DEF mapping. TMAP obtains the same mappings

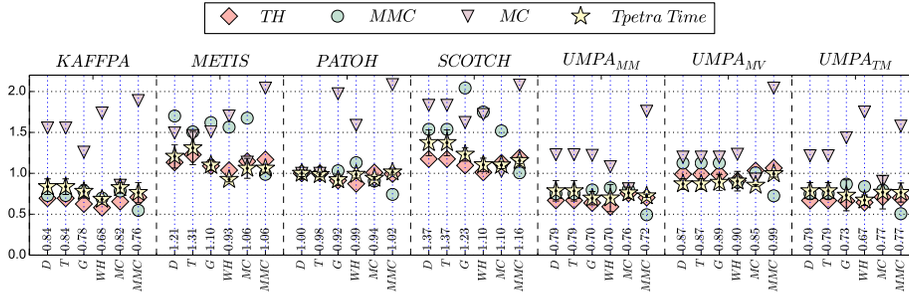


Figure 5: Trilinos SpMV results for `cage15` on 4096 processors. Each metric is normalized w.r.t that of DEF on G_t^{PATOH} .

with DEF on most of the graphs except $G_t^{\text{UMPA}_{\text{MM}}}$ and $G_t^{\text{UMPA}_{\text{MV}}}$. As the results for G_t^{PATOH} show, the proposed algorithms improves the performance 35–43% for `rgg` experiments.

The execution time is improved better with the algorithms minimizing WH and then MC. The improvements achieved by U_{MMC} is not as high as the others since for these “scaled” applications, the volume metrics are likely to be the bottleneck. In Section 4.5, we perform a regression analysis to better analyze the relation between the metrics and the execution time.

4.4 SpMV experiments

In this section, we study the impact of the proposed algorithms on the SpMV performance. We use `cage15` and perform SpMV using the Tpetra package of Trilinos with 500 and 1000 iterations. Figure 5 shows the performance results, where each metric and the overall execution time is normalized w.r.t. that of DEF on G_t^{PATOH} . The experiment is run for 4096 and 8192 parts on two different allocations. Only the results of a single allocation on 4096 processors is shown due to space limitations. The rest of results can be found at the supplementary page. The SpMV operation is repeated 5 times for each mapping and communication graph. We report the average of these 5 executions, and error bars represent the standard deviations. Unlike the previous experiment, TH is reported instead of WH, as its correlation with the total execution time is better.

In this setting, U_{WH} obtains the best performance; it decreases the overall execution time almost always (except for $G_t^{\text{UMPA}_{\text{MV}}}$) and up to 23% (for G_t^{METIS}) w.r.t DEF. U_{G} obtains a similar performance with slightly higher execution times. Although U_{MC} improves the performance for many cases w.r.t DEF, its performance is not as competitive as in the previous experiment since the message sizes are much smaller. Similar to the communication-only experiment, U_{MMC} obtains smaller improvements than the other UMPA variants. The overall performance of TMAP is very close to DEF, since it returns the DEF mapping for most of the cases.

Overall, TH highly correlates with the execution time. Moreover, this cor-

relation also holds among different communication graphs. In Section 4.2, we already observed that TH is much lower on the graphs with a lower TM. Improving the TH metric via both partitioning (with the objective TM) and mapping significantly reduces the parallel SpMV time. The best TM values for `cage15` have been found by KAFFPA, UMPA_{MM} and UMPA_{TM} (see the supplementary page for a `cage15`-only version of Fig. 1) and as Fig. 5 shows, these are the best partitioners for the default mapping. U_{WH} reduces the execution time by another 9–16% for the for these cases and obtains the best overall execution time for $G_t^{\text{UMPA}_{\text{TM}}}$. This is more than two times faster than the slowest variant, which is obtained by the DEF on G_t^{SCOTCH} . It also has 34% lower TM and 44% lower TH value. This shows the importance both the partitioning and mapping on SpMV performance.

4.5 Regression analysis

To analyze the performance improvements obtained for the communication-only applications and SpMV kernel w.r.t. the partitioning and mapping metrics, we use a linear regression analysis technique and solve a nonnegative least squares problem (NNLS). In NNLS, given a variable matrix \mathbf{V} and a vector \mathbf{t} , we want to find a dependency vector \mathbf{d} which minimizes $\|\mathbf{V}\mathbf{d} - \mathbf{t}\|$ s.t. $\mathbf{d} \geq 0$. In our case, \mathbf{V} has 14 columns: the partitioning metrics MSV, TV, MSM, TM; the mapping metrics WH, TH, MC, MMC, AC, AMC; inter-node communication volume (ICV), i.e., the total communication volume on the network excluding the intra-node communication (from TV); number of inter-node communication messages (ICM); the maximum receive volume of a node (MNRV); and the maximum number of messages received by a node (MNRM). A row \mathbf{t} of \mathbf{V} corresponds to an execution where the time is put to the corresponding entry of \mathbf{t} . To standardize each entry of \mathbf{V} and make them equally important, each column of \mathbf{V} is normalized by first subtracting the column mean from each column entry, and dividing them to the column standard deviation. We then use MATLABs *lsqnonneg* to solve NNLS. The coefficient \mathbf{d}_i of the output shows the dependency of the execution time to the i th metric.

We perform linear regression on the communication-only experiments' results with `cage15` graphs, 4096 processors and two sparse Hopper allocations. The analysis distinguished three metrics with non-zero coefficients. The metric with the highest coefficient is WH, followed by MSV and MC (0.023, 0.020 and 0.20), whereas the message-based metrics are found not to highly correlate with the performance. This is expected since the communication is scaled and the volume metrics' importance are increased. The results show that from the mapping perspective, WH and MC are the most important metrics for the applications with a high communication volume, whereas from the partitioning perspective, it is likely to be MSV.

We used the same experimental setting (`cage15`, 4096 processors, two allocations) for the SpMV kernel which is more latency bounded than the communication-only counterpart since there is no scaling on the communication volume. The metrics with non-zero coefficients are found to be AMC, ICV, MMC, TH, and

Table 1: Average improvements of the mapping algorithms on communication-only applications and SpMV kernel that runs for 500 and 1000 iterations for the first and second allocations, respectively.

	# procs	Rep.	DEF	TMAP	U_G	U_{WH}	U_{MC}	U_{MMC}
cage15 SpMV	4096	1	1.44 sec.	1.01	0.93	0.87	0.93	0.95
		2	2.77 sec.	1.00	0.91	0.89	0.94	0.95
	8192	1	1.25 sec.	0.99	0.98	0.96	0.99	1.01
		2	3.43 sec.	1.01	0.99	0.94	1.01	1.04
		Gmean	2.03 sec.	1.00	0.95	0.91	0.97	0.99
	cage15 Comm	4096	1	0.28 sec.	1.06	0.90	0.83	0.88
2			0.28 sec.	1.06	0.88	0.82	0.88	1.18
8192		1	0.19 sec.	1.01	1.02	0.89	1.01	1.16
		2	0.20 sec.	1.00	0.95	0.89	0.99	1.18
		Gmean	0.23 sec.	1.03	0.93	0.86	0.94	1.17
rgg Comm		4096	1	0.39 sec.	1.11	0.77	0.83	0.79
	2		0.33 sec.	0.98	0.96	0.77	0.84	0.87
		Gmean	0.36 sec.	1.05	0.86	0.80	0.81	0.86

MNRV (0.109, 0.070, 0.051, 0.050, 0.040). Since AMC better correlates with the performance compared to TH, it can be a good practice to utilize the already used links while reducing TH. One weakness of the regression analysis is that when highly correlating metrics are given in \mathbf{V} , the analysis may return a positive coefficient for only one of them. In our case, the importance of MNRM, ICM, and TM is hidden by the regression analysis. We also computed pairwise Pearson correlation of the metrics and observed a high correlation (≥ 0.92) of these metrics with AMC.

4.6 Summary

Table 1 presents a summary of the improvements achieved by the mapping algorithms in our experiments. For each allocation and part number, we calculate the geometric mean of the execution times obtained with the mapping methods on all graphs. The table shows the geometric mean of the execution times for DEF, and the normalized time for the other algorithms. The average for all allocations and part numbers are given at the bottom of the table. Overall, U_{WH} improves the cage15 SpMV kernel time by 4–13%, whereas the improvements for the communication-only cage15 and rgg applications are 14% and 20%, respectively, w.r.t. DEF.

5 Conclusion

We have proposed fast and high quality topology-aware task mapping methods that use graph models. We have compared the proposed methods with some other graph-based algorithms from the literature and with a default method used in Nersc’s Hopper Supercomputer. The experiments showed that on a set

of 25 matrices from the UFL collection, the proposed methods obtained high quality mappings in a very short time for the target system. The experiments with 4096 processors revealed significant improvements on the mapping metrics compared to the Hopper’s default mapping. These improvements yield a 43% performance improvement on one case for a communication-only application and a 23% improvement on the SpMV performance. Overall, with 4096 and 8192 processors, the proposed algorithms improve the performance of the SpMV kernel and the communication-only applications by 9% and 14%, respectively. We also evaluated the metrics according to their correlation with the performance. For the applications with a large communication volume, our analysis revealed that the weighted hop metric is the most dominant one, and for those with smaller messages, the average message congestion is a good metric that correlates with the performance.

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