Predicting Application Performance using Supervised Learning on Communication Features

Nikhil Jain†, Abhinav Bhatele†, Michael P. Robson†, Todd Gamblin†, Laxmikant V. Kale†

†Department of Computer Science, University of Illinois at Urbana-Champaign, Urbana, IL
†Center for Applied Scientific Computing, Lawrence Livermore National Laboratory, Livermore, CA
†{nikhil, mprobson, kale}@illinois.edu, †{bhatele, tgamblin}@llnl.gov

ABSTRACT

Task mapping on torus networks has traditionally focused on either reducing the maximum dilation or average number of hops per byte for messages in an application. These metrics make simplified assumptions about the cause of network congestion, and do not provide accurate correlation with execution time. Hence, these metrics cannot be used to reasonably predict or compare application performance for different mappings. In this paper, we attempt to model the performance of an application using communication data, such as the communication graph and network hardware counters. We use supervised learning algorithms, such as randomized decision trees, to correlate performance with prior and new metrics. We propose new hybrid metrics that provide high correlation with application performance, and may be useful for accurate performance prediction. For three different communication patterns and a production application, we demonstrate a very strong correlation between the proposed metrics and the execution time of these codes.

Categories and Subject Descriptors

C.2.1 [Computer-Communication Networks]: Network Architecture and Design—Network communications; C.4 [Performance of Systems]: Measurement techniques, Modeling techniques

General Terms

Measurement, Performance

Keywords

prediction, modeling, supervised learning, torus networks, contention, task mapping

1. INTRODUCTION

Intelligent mapping of application tasks on nodes of a torus partition can significantly improve the communication performance of the code [5, 7, 10]. The process of finding the best mapping for an application may require executing a large number of runs at different scales (number of processors). This can consume a significant amount of resources including both man hours and machine allocation. Hence, it is desirable to predict the communication performance of an application on a system without performing real runs, given a communication graph and the mapping of tasks to nodes on the machine. The central question we want to answer is whether a mapping improves performance over the default or another mapping. For this goal, predicting the correct rank correlation of different mappings based on their performance is sufficient. Predicting the absolute performance correctly is a secondary consideration.

Traditionally, task mapping on torus networks has focused on either reducing the maximum dilation or average number of hops per byte for messages in an application. These metrics make simplified assumptions about the cause of network congestion and do not provide accurate correlation with execution time. Mapping algorithms [4, 7, 15, 16] are generally designed to minimize these metrics which might be sub-optimal. Detailed network simulations provide another tool for predicting application performance but the simulation models also require a deep understanding of congestion issues and their root causes.

Understanding network congestion requires a study of message flow on the network. We present a detailed description of the life cycle of a message on the five-dimensional torus interconnect of Blue Gene/Q. From the instant that a message is placed in the injection queue on the source node to when it is received at the destination, we examine the various resources for which messages can contend, and thus potentially suffer from delays. This information is extremely valuable in writing network simulators, deciding which hardware counters to probe for contention, and developing new metrics that might provide a better correlation with application performance.

The focus of this paper is on the use of supervised learning algorithms, such as forests of randomized decision trees, to correlate individual metrics and their combinations with application performance. In addition to using prior metrics for predicting performance, we present several new metrics. Some of these can be obtained by analyzing the communication graph, network graph and the mappings, and others can be obtained through an experimental or simulated application run. For the results in this paper, the latter are obtained using real experiments. As part of future work, we plan to develop simulation techniques to obtain them offline. Maximum dilation is an example of an analytically derived
metric. In contrast, maximum bytes on a link is an experimentally obtained metric. In addition to these new metrics, we also use derived metrics that use information only from some outliers (nodes or links).

We present performance predictions using the randomized forests ensemble method for three different communication kernels: a two-dimensional five-point halo exchange, a three-dimensional 15-point halo exchange, and an all-to-all benchmark over sub-communicators. We show almost perfect correlation for runs on 16,384 and 65,536 cores of Blue Gene/Q. We also show predictions for a production application, pF3D, and for combining samples from different benchmarks into a single training set and test set.

In Section 2, we describe the common metrics used in literature and motivate the need for more precise metrics. Sources of contention on torus networks, methodology for collecting hardware counters data, and the proposed metrics are discussed in Section 3. The benchmarks and supervised learning techniques used in the paper and the measures of prediction success are described in Section 4. In Sections 5, 6, 7, we present results using prior metrics, new metrics and their combinations. We conclude our work in Section 8.

2. BACKGROUND AND MOTIVATION

Several metrics have been proposed in the literature to evaluate task mappings offline. Let us assume a guest graph, $G = (V_g, E_g)$ (communication graph between tasks) and a host graph, $H = (V_h, E_h)$ (network topology of the parallel machine). $M$ defines a mapping of the guest graph on the host graph ($G$ on $H$). The earliest metric that was used to compare the effectiveness of task mappings is dilation [3, 12]. Dilation for a mapping $M$ can be defined as,

$$dilation(M) = \max_{e_i \in E_g} d_i(M)$$

(1)

where $d_i$ is the dilation of the edge $e_i$ for a mapping $M$. Dilation of an edge $e_i$ is the number of hops between the end-points of the edge when mapped to the host graph. This metric aims at minimizing the length of the longest wire in a circuit [3]. We will refer to this as maximum dilation to avoid any confusion. We can also calculate the average dilation per edge for a mapping as,

$$\text{average dilation-per-edge}(M) = \frac{\sum_{e_i \in E_g} d_i(M)}{|E_g|}$$

(2)

Hoefler and Snir overload dilation to describe the “expected” dilation for an edge and “average” dilation for a mapping [11]. Their definition of expected dilation for an edge can be reduced to equation 1 above by assuming that messages are only routed on shortest paths, which is true for the IBM Blue Gene and Cray XT/XE family (if all nodes are in a healthy state). The average dilation metric, coined by Hoefler and Snir, is a weighted dilation and has been previously referred to as the hop-bytes metric by Sadayappan [9] in 1988 and Agarwal in 2006 [2]. Hop-bytes is the weighted sum of the edge dilations where the weights are the message sizes. Hop-bytes can be calculated by the equation,

$$\text{hop-bytes}(M) = \sum_{e_i \in E_g} d_i(M) \times w_i$$

(3)

where $d_i$ is the dilation of edge $e_i$, and $w_i$ is the weight (message size in bytes) of edge $e_i$.

Hop-bytes gives an indication of the overall communication traffic being injected on to the network. We can derive two metrics based on hop-bytes: the average number of hops traveled by each byte on the network,

$$\text{average hops-per-byte}(M) = \frac{\sum_{e_i \in E_g} d_i(M) \times w_i}{\sum_{e_i \in E_g} w_i}$$

(4)

and the average number of bytes that pass through a hardware link,

$$\text{average bytes-per-link}(M) = \frac{\sum_{e_i \in E_g} d_i(M) \times w_i}{|E_h|}$$

(5)

The former gives an indication of how far each byte has to travel on average. The latter gives an indication of the average load or congestion on a hardware link on the network. They are derived metrics (from hop-bytes) and all three are practically equivalent when used for prediction. In the rest of the paper, we use average bytes-per-link.

Another metric that indicates congestion on network links is the maximum number of bytes that pass through any link on the network,

$$\text{maximum bytes}(M) = \max_{l_i \in E_h} \left( \sum_{e_j \in E_g \mid e_j \rightarrow l_i} w_j \right)$$

(6)

where $e_j \rightarrow l_i$ represents that edge $e_j$ in the guest graph goes through edge (link) $l_i$ in the host graph (network). Hoefler and Snir use a second metric in their paper [11], worst-
3. CONTENTION ON TORUS NETWORKS

Although the coefficient of determination values ($R^2$, metric used for prediction success) are high, there is a significant variation in the y-values for different points with the same x-value. For example, in the maximum bytes plot (right), for $x = 6e9$, there are mappings with performance varying from 20 to 50 ms. These variations make predicting performance using simple models with a reasonably high accuracy ($\pm 5\%$ error) difficult. This motivated us to find new metrics and ways to improve the correlation between metrics and application performance.

3.1 Message flow and resource contention

Figure 2 presents the life cycle of a message on BG/Q. The tasks on a node send data onto the network through the Messaging Unit (MU) on the node. Injection memory FIFOs (imFifo) is the data structure used to transfer information between the tasks and the MU. To initiate a message send, a task puts a descriptor of the message in one of its imFitos. Selection of which imFitio to inject a descriptor in is typically based on the difference in coordinates of source and destination. The MU processes the descriptors in the imFitos, packetizes the message data it reads from the memory (packet size up to 512 B), and injects them into the injection network FIFOs. The descriptor of the message contains information of the binding that is used by the MU to inject into the appropriate injection network FIFO. In the default setting, there is a one-to-one mapping between imFitos and injection network FIFOs. This may lead to contention for injection network FIFOs if the distribution of source-destination pairs is such that a particular network injection FIFOs receives more traffic than others.

From the injection network FIFOs, packets are sent over the network based on the routing strategy and the destination. On the network, contention for hardware links is the most common source of performance degradation. When a packet injected on a link reaches an immediate neighbor, the network device decides the next link the packet needs to be forwarded to. If the next link is occupied, the packets are stored in buffers mapped to the incoming link. In the event of heavy contention for links, these buffers may get filled quickly, and prevent the use of the incoming link for data transfer. When packets eventually reach their destination, they are copied by the MU from the reception network FIFOs to either the reception memory FIFOs or the application memory. Limited memory bandwidth may prevent the MU from copying the data, and hence reception injection FIFOs and the buffers attached to the corresponding links may get filled. This may lead to blocking of the links for further communication.

3.2 Collecting hardware counters data

We use two methods to collect information that can indicate resource contention as described in Section 3.1:

**Blue Gene/Q Counters:** The Hardware Performance Monitoring API (BGPM) provides a simple interface for the BG/Q Universal Performance Counter (UPC) hardware. The UPC hardware programs and counts performance events from multiple hardware units on a BG/Q node. Using BGPM to control the Network Unit of the UPC hardware, we implemented a PMPI-based profiling tool that records the following information:

- Sent chunks: count of 32-byte chunks sent on a link.
- Counters used for collecting this information:
  - PEVT_NW_USER_PP_SENT: user-level point-to-point 32-byte packet chunks sent (includes chunks in transit).
  - PEVT_NW_USER_ESC_PP_SENT: user-level deterministic point-to-point 32-byte packet chunks sent (includes
4. EXPERIMENTAL SETUP

In this section, we describe the machines and benchmarks used for the experiments, and the scikit package which provides a Python interface for using several machine learning algorithms.

4.1 Machines

All the experiments were performed on two Blue Gene/Q systems – Mira and Vulcan. Mira is installed at the Argonne National Laboratory and Vulcan is hosted at the Lawrence Livermore National Laboratory. Table 2 presents the possible sizes of the torus dimensions (A, B, C, D and E) when partitions of 1024 and 4096 nodes are requested on BG/Q.

<table>
<thead>
<tr>
<th>Nodes</th>
<th>A</th>
<th>B</th>
<th>C</th>
<th>D</th>
<th>E</th>
</tr>
</thead>
<tbody>
<tr>
<td>1024</td>
<td>4</td>
<td>4</td>
<td>4</td>
<td>8</td>
<td>2</td>
</tr>
<tr>
<td>4096</td>
<td>4</td>
<td>4</td>
<td>8</td>
<td>16</td>
<td>2</td>
</tr>
<tr>
<td>4096</td>
<td>4</td>
<td>8</td>
<td>4</td>
<td>16</td>
<td>2</td>
</tr>
</tbody>
</table>

Table 2: Dimensions of the allocated job partitions on BG/Q.

4.2 Communication kernels

In our experiments, we use three benchmarks that exhibit communication patterns that are commonly seen in HPC applications. By running the benchmarks for four different message sizes, 8 bytes, 512 bytes, 16 KB and 4 MB, we are able to cover a broad range of communication scenarios typical in large scale scientific applications. In addition, these message sizes also cover different MPI protocols (short, eager, rendezvous), and routing protocols (deterministic and adaptive). All the benchmarks have been implemented in MPI and are described below.

4.2.1 Five-point 2D halo exchange

2D Halo implements the communication pattern of a two-dimensional Jacobi relaxation, which is a common kernel in HPC applications. MPI ranks are arranged in a 2D grid and in each iteration, every MPI rank exchanges one message each with its four nearest neighbors. Performance of this benchmark is measured by taking the average execution time over 100 iterations.

4.2.2 15-point 3D halo exchange

Our second benchmark, 3D Halo, decomposes the MPI ranks into a three-dimensional (3D) grid. In each iteration, every MPI rank communicates with fourteen neighbors (six faces and eight corners) that constitute a 15-point stencil (including itself). 3D Halo differs from 2D Halo in two important aspects: it puts more communication load on the network, and it decomposes MPI ranks into a higher dimension grid. We use the average execution time for 100 iterations to measure the performance.

4.2.3 All-to-all over sub-communicators

The Sub A2A benchmark is a departure from both 2D Halo and 3D Halo. It performs simultaneous all-to-alls within sub-communicators, and hence is more communication intensive. MPI ranks are decomposed into a 3D grid with sub-communicators of size 64 created along one of its dimensions. Sub A2A has been modeled on codes that perform FFTs, in which every rank shares data with all other ranks in the same Cartesian sub-grid. The average execution time for 50 iterations is used as the benchmark performance.

4.2.4 Input Data

We use a Python mapping tool, Rubik [6, 13] to generate 84 different task mappings for each of the benchmarks.

---

Table 1: Prior and new metrics that indicate contention for network resources.

<table>
<thead>
<tr>
<th>Indicator</th>
<th>Source</th>
<th>Derived from</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bytes on links</td>
<td>Counters</td>
<td>Sent chunks</td>
</tr>
<tr>
<td>Buffer length</td>
<td>Counters</td>
<td># Packets in buffers</td>
</tr>
<tr>
<td>Delay per link</td>
<td>Counters</td>
<td># Packets in buffers divided by received packets</td>
</tr>
<tr>
<td>Dilation</td>
<td>Analytical</td>
<td>Shortest-path routing between source and destination</td>
</tr>
<tr>
<td>FIFO length</td>
<td>Analytical</td>
<td>Based on PAMI source</td>
</tr>
</tbody>
</table>

---
and system sizes. Hardware counters data from real runs and data generated by the analytical program for the three benchmarks, four different message sizes, two partition sizes and 84 different task mappings, was used as an input to the machine learning program.

4.3 Prediction using ensemble methods

We employ supervised learning techniques used in statistics and machine learning to predict the performance (execution time) of an application for different mappings using metrics described in Section 3.3. The learning algorithm infers a model or function by working with a training set that consists of samples (mappings), and one or more input features (raw and/or derived such as average bytes) per sample. Each sample has a desired output value (execution time), also known as the target. We then use the trained algorithm to predict the output for a testing set (new mappings for which we wish to predict execution time). The values of the features and the target are normalized to obtain the best results.

We tested several supervised learning techniques ranging from statistical techniques, such as linear regression, to machine learning methods, such as support vector machines and decision trees. The scikit-learn package provides several of these algorithms/estimators in Python [14]. The two steps, as described previously, are to first fit the estimator and then to use it to predict unseen samples. For the benchmarks presented in this paper, ensemble learning provided the best fit. Ensemble methods use predictions from several, often weaker models, to produce a stronger model or estimator that gives better results than the individual models.

Random forests are a type of ensemble learning method developed by Leo Breiman in 2001 [8]. The idea is to build several decision trees and to use averaging methods on these independently built models. In a decision tree, the goal is to create a model that predicts the value of a target variable by learning simple decision rules inferred from the data features. These trees are simple to understand and to interpret as they can be visualised easily. However, decision tree based learners create biased trees if some patterns dominate. Random forests attempts to avoid such a bias by adding randomness in the selection of condition when splitting a node during the creation of a decision tree. Instead of choosing the best split among all the features, the split that is picked is one among a random subset of the features. Further, the averaging performed on the prediction from independently built decision trees leads to a reduction in the variance and hence an overall better model. We use the ExtraTreesRegressor class in scikit 0.13.1.

Figure 3 (left) shows an example decision tree that was used in one of our experiments. This tree was produced by training performed using two features, maximum bytes and average bytes, to predict the target, the execution time. The tree shows that at each level, based on the conditions derived from the training set on values of maximum bytes and average bytes, the prediction is guided to reach a leaf node, which determines the target value. Figure 3 (right) presents the overlay of a number of such decision trees over a 2D space spanned by the same two input features. Each color in the figure is a leaf region of one of the decision trees in the random forest generated by fitting the training set. As expected, regions from different decision trees overlap, and cover the entire space. The white circles are the test set being predicted - new mappings with known maximum bytes and average bytes. To predict, all the leaf regions, to which a test case belong, are computed. This provides a set of local predictions for the test case, which are combined to provide a unique target value.

4.4 Metrics for prediction success

The goodness or success of the prediction function (also referred to as the score) can be evaluated using different metrics depending on the definition of success. Our main goal is to compare the performance of two mappings and deter-
5. PERFORMANCE PREDICTION OF COMMUNICATION KERNELS

In this section, we present results on the prediction of execution times of several communication kernels (Section 4.2) for different task mappings.

5.1 Performance variation with mapping

Figure 4 presents the execution times for the three benchmarks for four message sizes – 8 bytes, 512 bytes, 16 KB and 4 MB. These sizes represent the amount of data exchanged between a pair of MPI processes in each iteration. For example, for 2D Halo, this number is the size of a message sent by an MPI process to each of its four neighbors. For a particular message size, a point on the plot represents the execution time (on the y-axis) for a mapping (on the x-axis). For 2D Halo, Figure 4(a) shows that for small messages such as 8 and 512 bytes, mapping has an insignificant impact. As the message size is increased to 16 KB, in addition to an increase in the runtime, we observe up to a 7x difference in performance for the best mapping in comparison to the worst mapping (note the logarithmic scale on the y-axis). Similar variation is seen as we further increase the message size to 4 MB. For a more communication intensive benchmark, 3D Halo, we find that mapping impacts performance even for 512-byte messages (Figure 4(b)). As we further increase the communication in Sub A2A, the effect of task mapping is also seen for the 8-byte messages as shown in Figure 4(c).

In the following sections, we do not present results for the cases where the performance variations from mapping are statistically insignificant: 8- and 512-byte results in case of 2D Halo and 8-byte results in case of 3D Halo.

5.2 Prior features

We begin with showing prediction results using prior metrics/features (described in Section 2) and quantify the goodness of the fit or prediction using rank correlation coefficient (RCC) and \( R^2 \) (Section 4.4). Figure 5 (top) presents the RCC values for predictions based on prior features (maximum dilation, average bytes per link and maximum bytes on a link). In most cases, we find that the highest value for RCC is 0.91, i.e., the pairwise ordering of 91% of mapping pairs was predicted correctly. For a testing set of 28 samples, an RCC of 0.91 implies incorrect prediction of the pairwise ordering of 38 mapping pairs. A notable exception is the 512-byte case for 3D Halo where the RCC is 0.96. In
contrast, for 16 KB message size, the highest RCC is only 0.86.
In the case of 2D Halo and 3D Halo, prediction using maximum bytes on a link has the highest RCC while prediction using maximum dilation performs very poorly with an RCC close to 0.60. However, for Sub A2A, prediction using average bytes per link is better than prediction using maximum bytes on a link for small to medium message sizes (by 4-5%).
The metric for absolute performance correlation, $R^2$, is also shown in Figure 5. For all benchmarks and message sizes, maximum bytes on a link performs the best with a score of up to 0.95 for 3D Halo and Sub A2A. These results substantiate the use of maximum bytes and average bytes as simple metrics that are roughly correlated with performance.

5.3 New features

We propose new metrics/features based on the buffer length, delay and FIFO lengths (see Table 1) and derive others by extracting counters and analytical data for outlier nodes and links:

**Average Outliers (AO)** We define a node or link as an average outlier if an associated value for it is greater than the average value of the entire data set. Selection of data points based on the average value helps eliminate low values that can skew derived features and hide information that may be useful.

**Top Outliers (TO)** Similar to the average outlier, we can define a node or link to be a top outlier if an associated value for it is within 5% of the maximum value across the entire data set.

We can use these two outlier selection criteria to define metrics that represent the features extracted from outliers. Among a large set of features that we explored using prior/new metrics in combination with known/new derivation methods, we focus on prediction using the following features that had the highest RCC: average buffer length ($\text{avg buffer}$), average buffer length of TO ($\text{avg buffer TO}$), sum of maximum dilation for AO ($\text{sum dilation AO}$), average bytes per link for AO ($\text{avg bytes AO}$), and the average bytes per link for TO ($\text{avg bytes TO}$).

The most important point to note as we transition from Figure 5 to Figure 6 is the general increase in RCC. For Sub A2A in particular, we observe that RCC is consistently 0.95. The previous poor predictions in the case of 16 KB message size for 3D Halo improve from 0.86 to 0.90 (RCC value). For the low traffic 2D Halo, new network-related features such as those based on the buffer length exhibit low correlation. As traffic on the network is increased (larger messages sizes) in 3D Halo and Sub A2A, the RCC of these new network-related features increases, and occasionally surpasses the RCC of other features.

We note that the $R^2$ value is consistently high only for the $\text{avg bytes TO}$ feature. For a number of features, the $R^2$ values are either low or zero. There are two reasons that can explain the low $R^2$ values: 1) the features did not correlate well with the performance, e.g. $\text{avg buffer}$ for 2D Halo and 3D Halo, or 2) the predicted performance followed the same trend as the observed performance but was offset by a factor, e.g., $\text{avg buffer}$ for large messages in 3D Halo.

5.4 Hybrid features

The previous sections have shown that up to 94% prediction accuracy can be achieved using a single feature. Depending on the benchmark and the message size, the feature that provides the best prediction may vary. This motivated us to use several features together to improve the correlation and enable better prediction.

In order to derive hybrid features that improve RCC, we performed a comprehensive search by combining the features that had high RCC values. In addition, the combinations of good features were also augmented with features that exhibited low to moderate correlation with performance. We made two important discoveries with these experiments: 1) combining multiple good features may result in a lower accuracy of prediction, and 2) the addition of features that had limited success on their own to good features can boost prediction accuracy significantly.

<table>
<thead>
<tr>
<th>Hybrid</th>
<th>Features combined</th>
</tr>
</thead>
<tbody>
<tr>
<td>H1</td>
<td>avg bytes, max bytes, max FIFO</td>
</tr>
<tr>
<td>H2</td>
<td>avg bytes, max bytes, sum dilation AO, max FIFO</td>
</tr>
<tr>
<td>H3</td>
<td>avg bytes, max bytes, avg buffer, max FIFO</td>
</tr>
<tr>
<td>H4</td>
<td>avg bytes, max bytes, avg buffer TO</td>
</tr>
<tr>
<td>H5</td>
<td>avg bytes TO, avg buffer TO, avg delay AO, sum hops AO, max FIFO</td>
</tr>
<tr>
<td>H6</td>
<td>avg bytes TO, avg buffer AO, avg delay TO, avg delay AO, sum hops AO, max FIFO</td>
</tr>
</tbody>
</table>

Table 3: List of hybrid features that achieve strong correlations.

Figure 7 presents results for the hybrid features that consistently provided high prediction success with different benchmarks and message sizes. Table 3 lists the features that were combined to create these hybrid features. In our experiments, we found that combining the two most commonly used features, max bytes and avg bytes improves the prediction accuracy in all cases. The gain in RCC was highest for the 4 MB message size, where RCC increased from
Figure 6: Prediction success based on new features on 16,384 cores of BG/Q. We observe a general increase in RCC, but $R^2$ values are low in most cases resulting in empty columns.

Figure 7: Prediction success based on hybrid features from Table 3 on 16,384 cores of BG/Q. We obtain RCC and $R^2$ values exceeding 0.99 for 3D Halo and Sub A2A. Prediction success improves significantly for 2D Halo also.

0.91 (individual best) to 0.94 for all benchmarks. The addition of max FIFO, which did not show a high RCC score as a stand alone feature, further increased the prediction accuracy to 0.96. We denote this set as H1.

To H1, we added another low performing feature, avg buffer to obtain H3. This improved the RCC further in all cases with the RCC score now in the range of 0.98 – 0.99 for 3D Halo and Sub A2A (Figure 7). Replacing avg buffer is this set with avg buffer TO improved the RCC for Sub A2A, but reduced the RCC for 2D Halo. Using a number of such combinations, we consistently achieved RCC up to 0.995 for Sub A2A. Given a testing set of size 28, this implies that the pairwise order of only 2 pairs was mispredicted.

Prediction using hybrid features also results in high $R^2$ values as shown in Figure 7. For 2D Halo, the scores go up from 0.95 and 0.93 to 0.975 and 0.955 for the 16 KB and 4 MB message sizes respectively. For the more communication intensive benchmarks, we obtained $R^2$ values as high as 0.99 in general. Hence, the use of hybrid features not only predicts the correct pairwise ordering of mapping pairs but also does so with high accuracy in predicting their absolute performance.

5.5 Results on 65,536 cores

Figure 8 shows the prediction success for the three benchmarks on 65,536 cores of BG/Q. From all the previously pre-
sented features (prior, new and hybrid), we selected the ones with the highest RCC scores for 16,384 cores, and present only those in this figure. Similar to results on 16,384 cores, we obtain significant improvements in the prediction scores using hybrid features in comparison to individual features such as max bytes and avg bytes TO. For Sub A2A, RCC improved by 14% from 0.86 to 0.98, with a RCC value of 1.00 for both 512 bytes and 4 MB message sizes. For 2D Halo and 3D Halo, an improvement of up to 8% was observed in the prediction success. Similar trends were observed for $R^2$ values.

Figure 9 presents the scatter-plot of predicted performance for the three benchmarks for the 4 MB message size. On the x-axis are the task mappings sorted by observed performance, while the y-axis is the predicted performance. The feature set $H_5$: avg bytes TO, avg buffer TO, avg delay AO, sum hops AO, max FIFO was used for these predictions. It is evident from the figure that an almost perfect performance based ordering can be achieved using prediction for all three benchmarks, as expected based on the high RCC values. In addition, as shown in Figure 9, absolute performance values can also be predicted accurately using the proposed hybrid metric. In particular, for Sub A2A with large communication volume, the predicted value curve completely overlaps with the observed value curve. These results suggest that the same set of features correlate with the performance irrespective of the system size being used.

6. COMBINING ALL TRAINING SETS

In the previous section, we presented high scores for predicting performance of the three benchmarks using hybrid metrics. For the prediction of individual benchmarks, the training and testing sets were generated using 84 different mappings of the same benchmark for a particular message size on a fixed core count. In this section, we relax these requirements, and explore the space where the training and testing sets are a mix of different benchmarks, message sizes and core counts.

6.1 Combining samples from different benchmarks

We first explore the use of training and testing sets that
are a combination of all three benchmarks with 16 KB and 4 MB message sizes. It is to be noted that the training and testing sets are now six times the size of individual sets (336 vs. 56 for the training set and 168 vs. 28 for the testing set). Figure 10 presents the prediction success for this experiment using prior, new and hybrid features with best correlation.

High RCC values, such as 0.97 for avg bytes, suggests that the combination of training sets results in a better prediction than the individual cases. Absolute number of mispredictions, presented in Figure 11, is close to the sum of mispredictions for individual cases (that are combined to form these sets) for a given metric. This suggests that the presented technique was successful in classifying the sample data from different kinds of communication patterns and message sizes, and in making good predictions using them.

These results indicate that if a large database consisting of different communication patterns and message sizes is created, predicting performance of different classes of applications (possibly with unknown communication structure) may be feasible. We leave an in-depth study of this aspect for future work.

6.2 Predicting performance on 65,536 cores using 16,384-core samples

We also experimented with predicting the performance on 65,536 cores using the data from runs on 16,384 cores as the training set. The testing set consists of all the benchmarks with message sizes as 16 KB and 4 MB on 65,536 cores, while the training set is the same benchmarks run on 16,384 cores. As shown in Figure 12, we obtained high RCC values for many features. The maximum RCC value of 0.975 was observed using the hybrid feature set H3: avg bytes, max bytes, avg buffer, max FIFO. In terms of absolute number of pairs with incorrect predicted ordering, ordering of ~ 3,200 pairs were mispredicted among a full set of 1,26,756 pairs.

We find these results to be very encouraging since a strong correlation for predicting performance on large node counts using data from smaller jobs may provide a scalable method for performance prediction. Using smaller systems to predict performance at scale has several advantages. First, generating data sets is more feasible in this regime because it consumes less resources. Second, manually generating various mappings for large systems is impractical, but using prediction based on runs on smaller node counts, a large number of mappings can be explored with low overhead.

7. RESULTS WITH PF3D

PF3D [17] is a multi-physics code used for studying laser plasma-interactions in the National Ignition Facility (NIF) experiments at LLNL. PF3D is a communication-heavy a-
plication and has been shown to benefit significantly from task mapping on Blue Gene/P [6]. This is the first attempt at mapping pF3D on Blue Gene/Q.

pF3D simulations consist of three distinct phases: wave propagation and coupling, advecting light, and solving the hydrodynamic equations. The MPI processes are arranged in a 3D Cartesian grid and all of the MPI communication is performed along different directions ($X, Y, Z$) of this grid. Wave propagation and coupling consists of two-dimensional (2D) Fast Fourier Transforms (FFTs) in $XY$-planes; the 2D FFTs are performed via two non-overlapping 1D FFTs along the $X$ and $Y$ directions using MPI_Alltoall. The advection phase involves planar exchange of data with neighbors in the $Z$-direction performed using MPI_Send and MPI_Recv. Finally, the hydrodynamic phase consists of near-neighbor data exchange in the positive and negative $X$, $Y$ and $Z$ directions. The FFT phase and the planar exchange in $Z$ account for most of the time spent in communication in pF3D. The logical 3D grid of processes used for pF3D in this paper was $16 \times 8 \times 128$.

In Figure 13, we present RCC scores for predicting the performance of pF3D on 16,384 cores of BG/Q. While avg bytes has a very low RCC, max bytes correctly predicts the pairwise ordering for 91% of the task mapping pairs. Interestingly, sum of dilations for messages that belong to the average outlier set exhibits a high RCC of 0.94. Similar to the benchmarks, the hybrid features show strong correlation with performance, and have RCCs exceeding 0.96 for this production application. The highest correlation achieved is for the hybrid set $H6$: avg bytes TO, avg buffer AO, avg delay TO, avg delay AO, sum hops AO, max FIFO that has an RCC of 0.995. The $R^2$ values are significantly lower, in contrast with the benchmarks. For max bytes, the $R^2$ value is only 0.76 which increases dramatically to 0.931 for the hybrid set $H6$.

Figure 14 summarizes the prediction results for pF3D on 16,384 cores using a scatter-plot of observed and predicted values. While the ordering of mapping is correctly predicted, the absolute values are significantly different for many mappings. As we scaled up to 65,536 cores, we found more irregularities between absolute values of predicted and observed performance. Moreover, the RCC values for known metrics were found to be as low as 0.52 on 65,536 cores. Using the new and hybrid metrics, the RCC values increased to 0.75 (45% improvement). These RCC values are significantly smaller in comparison to other results presented in this paper. As part of future work, we plan to research further on possible cause of such a divergence.

8. CONCLUSION

Significant time and effort wasted in real runs to evaluate the performance of different task mappings suggests the use of simulation or metrics to predict application performance offline. Metrics used previously in literature fall short in providing strong correlation with execution time. In this paper, we demonstrate the use of machine learning techniques to obtain high correlation between new metrics and performance of parallel applications for different task mappings.

In addition to prior metrics, such as maximum bytes, we have proposed new metrics, such as buffer length and messages in injection FIFOs, to include the effects of contention for network resources other than links. Using a combination of these metrics, which includes average and maximum bytes on links, maximum messages contending for an injection FIFO, and the average number of packets in buffers, we show rank correlation coefficients of up to 0.99, i.e. for only 15% of pairs the pairwise ordering is predicted incorrectly. This signifies an improvement of 14% in the prediction success for an all-to-all over sub-communicators benchmark and 8% for 2D and 3D halo exchange. In addition, prediction using such hybrid metrics also shows high $R^2$ scores which indicates good prediction in terms of absolute values. Identifying these metrics is a step towards accurate offline analysis of task mappings to find the best performing mapping. The next step is to develop techniques to compute metrics that
are obtained via real runs in this paper, using simulation and/or analysis.

We also successfully attempted combining of training and testing sets from different benchmarks and still retained high prediction accuracy. This suggests that if a large database consisting of different communication patterns and message sizes is created, predicting performance of different classes of applications (possibly with unknown communication structure) may be feasible. More importantly, we show that using training sets from small core counts, we can predict performance at a larger count with an RCC value of 0.975. This may provide a scalable method for performance prediction at large scales and for future machines without having to perform detailed network simulations. Finally, we have demonstrated that supervised learning and ensemble methods can be used to predict performance not only for simple communication kernels but also for complex production applications with several diverse communication phases.

Acknowledgments

This work was performed under the auspices of the U.S. Department of Energy by Lawrence Livermore National Laboratory under Contract DE-AC52-07NA27344. This work was funded by the Laboratory Directed Research and Development Program at LLNL under project tracking code 13-ERD-055 (LLNL-CONF-635857).

This research used computer time on Livermore Computing’s high performance computing resources, provided under the Laboratory Directed Research and Development Program. This research also used resources of the Argonne Leadership Computing Facility at Argonne National Laboratory, which is supported by the Office of Science of the U.S. Department of Energy under contract DE-AC02-06CH11357.

The authors thank W. Scott Futral (LLNL) and Kalyan Kumar (ANL) for their help with getting the jobs completed on time on Vulcan and Mira respectively. We also thank Dong Chen, Phil Heidelberger, Sameer Kumar, Jeff Parker and Bob Walkup from IBM for the numerous e-mail exchanges and discussions to improve our understanding of the Blue Gene networks.

9. REFERENCES


