© 2007 Abhinav Bhatele

APPLICATION–SPECIFIC TOPOLOGY–AWARE MAPPING AND LOAD BALANCING FOR THREE–DIMENSIONAL TORUS TOPOLOGIES

BY

ABHINAV BHATELE

B. Tech., Indian Institute of Technology, Kanpur, 2005

THESIS

Submitted in partial fulfillment of the requirements for the degree of Master of Science in Computer Science in the Graduate College of the University of Illinois at Urbana-Champaign, 2007

Urbana, Illinois

Adviser:

Professor Laxmikant V. Kalé

Abstract

The advent of supercomputers like Blue Gene/L, XT3, XT4 and Blue Gene/P has given rise to new challenges in parallel programming. Interconnect topology of machines is one such factor which has become crucial in optimizing the performance of applications on clusters with thousands of processors. If the topology of an interconnect is clearly defined, it is possible to minimize the communication volume and balance it evenly across processors. It can be minimized by co-locating communicating entities on nearby objects. This can be done at two stages in the program: 1. during program start-up where we define the initial mapping for migratable entities and static mapping for non-migratable entities, and 2. during load balancing when migratable entities are moved across processors to keep the load evenly balanced.

In this thesis, we examine how topological considerations in mapping and load balancing algorithms can help communication and make applications faster. To present concrete results, several applications written in CHARM++, 7-point 3dimensional Stencil, NAMD and LEANCP are considered. Results on 7-point Stencil are presented as a proof of principle because it is a relatively simple to analyze and map. NAMD and LEANCP on the other hand are production codes with thousands of users. NAMD is a classical molecular dynamics application and heavily depends on load balancing for optimal performance. Load balancers deployed in NAMD have been modified to place communicating objects in proximity to one another by considering the communication in multicasts. LEANCP is a quantum chemistry application based on the Car-Parrinello *ab-initio* Molecular Dynamics (CPAIMD) method. In LEANCP, the default mapping of its several object arrays to processors by the CHARM++ runtime has been modified to optimize communication. Results of improvements from these strategies are demonstrated for all these applications on IBM's Blue Gene/L and Cray's XT3. To my aunt, mother and girlfriend. Girija, Vimlesh and Shambhavi.

Acknowledgements

There are many people whom I need to thank for their help and support with this thesis. The order in which they are mentioned is immaterial and I apologize if I have missed out some names. I wish to thank my advisor Professor Laxmikant V. Kalé for showing the torch and helping me move forward inch by inch. He has always given me an ear, no matter how busy he has been. He has been my inspiration to continue for a PhD after my Master's degree.

My colleagues in the Parallel Programming Laboratory (PPL) have made my stay in US easier and a lot of fun. I thank Eric Bohm for helping me set my foot in research and for the constant technical help with LEANCP. Thanks to Gengbin Zheng for his help with various CHARM++, load balancing and NAMD problems. I also want to thank my colleagues Chee Wai Lee, Nilesh Choudhury and Sayantan Chakravorty for their help with various things.

Special thanks to Glenn Martyna and Sameer Kumar at IBM for their mentoring and support at my summer internship and otherwise. Thanks to Jim Phillips for help with understanding NAMD code. Many thanks to Fred Mintzer and Glenn Martyna at IBM and Shawn Brown and Chad Vizino at PSC for runs on Blue Gene/L and BigBen respectively. Thanks to Chao Huang at PPL for the Blue Gene/L topology interface and to Deborah Weisser at Cray Inc. for help with the XT3 topology interface. Again, it would have been difficult without Shawn and Chad's help to get the topology interface running on XT3.

Thanks to my friends for their support. Heartfelt thanks to all the wonderful people in my family who have always shown confidence in me. My mother, father and uncle for seeing a dream for me; My Maa, for her endless love and blessings which have brought me this far; and my brothers for their support. And finally, I thank my girlfriend who has been patient with me whenever I told her I was busy with my thesis. She has been a constant source of encouragement, motivation and inspiration.

Table of Contents

\mathbf{Li}	st of	Tables	x				
\mathbf{Li}	st of	Figures	x				
\mathbf{Li}	List of Abbreviations						
\mathbf{Li}	st of	Symbols	i				
1	Intr	oduction	1				
	1.1	Parallel Machines and Topologies	1				
	1.2	Effect of Topology on Communication	3				
	1.3	CHARM++ and its Applications	4				
2	Pro	cessor and Communication Graphs	6				
	2.1	Processor Topologies	6				
	2.2	Communication Scenarios	8				
		2.2.1 Regular and Irregular Communication	8				
		2.2.2 Point-to-Point, Multicasts and Reductions	9				
	2.3	Previous Work	0				
3	Obj	ects in Charm $++$	2				
	3.1	Chares and Arrays	3				
	3.2	Dynamic Mapping Framework	3				
	3.3	Topology Interface in CHARM++	4				
		3.3.1 Cray XT3	4				
	3.4	3D Jacobi	5				
		3.4.1 Topology Aware Mapping	5				
		3.4.2 Evaluation of Mapping Strategies	7				
4	Lea	nCP and Regular Communication	9				
	4.1	Parallel Implementation	9				
	4.2	Communication in LEANCP	2				
	4.3	Topology Aware Mapping	3				
		4.3.1 Mapping GSpace and RealSpace Arrays	4				
		4.3.2 Mapping of Density Objects	5				
		4.3.3 Mapping PairCalculator Arrays	5				

5	NA	MD and Multicasts	7
	5.1	Parallel Implementation	27
	5.2	Communication in NAMD	28
		5.2.1 Topological mapping of patches	29
	5.3	Load Balancing	80
		5.3.1 Topology-aware decisions	32
6	Res	ults	4
	6.1	3D Jacobi	35
		6.1.1 Performance on Blue Gene/L	35
		6.1.2 Performance on Cray XT3 [']	88
	6.2	LEANCP	39
	6.3	NAMD	12
7	Fut	ure Work and Conclusion	4
7	Fut 7.1	ure Work and Conclusion 4 Generalized Topology Sensitive Mapping 4	4 14
7	Fut 7.1 7.2	ure Work and Conclusion 4 Generalized Topology Sensitive Mapping 4 Load Balancers for Section Multicasts 4	4 14 15
7	Futo 7.1 7.2 7.3	ure Work and Conclusion 4 Generalized Topology Sensitive Mapping 4 Load Balancers for Section Multicasts 4 Summary 4	4 14 15 15
7	Fut: 7.1 7.2 7.3	ure Work and Conclusion 4 Generalized Topology Sensitive Mapping 4 Load Balancers for Section Multicasts 4 Summary 4	4 14 15 15
7 Re	Fut 7.1 7.2 7.3	ure Work and Conclusion 4 Generalized Topology Sensitive Mapping 4 Load Balancers for Section Multicasts 4 Summary 4 nces 4	4 14 15 15
7 Re Aj	Fut 7.1 7.2 7.3 eferen	ure Work and Conclusion 4 Generalized Topology Sensitive Mapping 4 Load Balancers for Section Multicasts 4 Summary 4 nces 4 dix A 5	4 44 45 45 7
7 Re Aj	Fut 7.1 7.2 7.3 eferen	ure Work and Conclusion 4 Generalized Topology Sensitive Mapping 4 Load Balancers for Section Multicasts 4 Summary 4 nces 4 dix A 5 dix B 5	4 45 45 7 7 3
7 Re Aj Aj	Fut 7.1 7.2 7.3 eferen open open	ure Work and Conclusion 4 Generalized Topology Sensitive Mapping 4 Load Balancers for Section Multicasts 4 Summary 4 nces 4 dix A 5 dix B 5 dix C 5	4 4 4 5 4 5 7 6 1 5 3 4

List of Tables

6.1	Performance of 3D Jacobi on Blue Gene/L (CO mode) for different	
	decompositions (Problem Size: fixed at $512 \times 512 \times 512$). The numbers	05
	are time in seconds for 1000 iterations.	35
6.2	Performance of 3D Jacobi on Blue Gene/L (CO mode) for different	
	decompositions (Problem Size: variable from $512 \times 512 \times 512$ to $1024 \times$	
	1024×1024). The numbers are time in seconds for 1000 iterations.	35
6.3	Performance and hop-counts for 3D Jacobi on Blue Gene/L (VN	
	mode) for block size $32 \times 32 \times 32$ (Problem Size: fixed at $512 \times$	
	$512\times1024).$ The numbers are time in seconds for 1000 iterations	37
6.4	Performance and hop-counts for 3D Jacobi on Cray XT3 for different	
	decompositions (Problem Size: fixed at $512 \times 512 \times 512$). N stands	
	for naïve and T stands for topology-aware mapping. The numbers	
	are time in seconds for 1000 iterations.	38
6.5	Performance and hop-counts for 3D Jacobi on Cray XT3 for different	
	decompositions (Problem Size: variable from $512 \times 512 \times 256$ to $1024 \times$	
	512×1024). N stands for naïve and T stands for topology-aware	
	mapping. The numbers are time in seconds for 1000 iterations.	38
6.6	Performance of LEANCP on Blue Gene/L (CO mode). The numbers	
0.0	represent time per step in seconds	40
67	Performance of LEANCP on Blue Gene/L (VN mode). The numbers	10
0.1	represent time per step in seconds	40
68	Reduction in hon bytes for 3D NAMD on Blue Cone/L (Bonch	40
0.8	$mark(Arg(A_1))$	49
6.0	mark: ApoA1)	4Z
0.9	Reduction in nop-bytes for 3D NAMD on Blue Gene/L (Benchmark: F_1 -	10
	A1 Pase	42

List of Figures

1.1	A three-dimensional torus network (All connections are not shown for the sake of simplicity)	2
$2.1 \\ 2.2$	(a) A 2-dimensional mesh network, (b) A 2-dimensional torus network A 4-dimensional hypercube network	7 7
$3.1 \\ 3.2$	Programmer's and System's view of the same program under execution Topology-aware mapping of 3D Jacobi's data array onto the 3D pro- cessor grid. Different colors (shades) signify which chares get mapped	12
3.3	to which processors	16 17
$4.1 \\ 4.2$	Flow of control in LEANCP	20 24
5.1 5.2 5.3	Communication between patches and other objects in NAMD Computes and patches mapped to the processor torus Choice of the best processor to place a compute. The best choice is on the extreme left where we get two proxies on one processor. The worst is on the right where we cannot find any proxy/patch on the processor	28 29 31
5.4	Topological search for a underloaded processor in the 3D torus	32
$ \begin{array}{r} 6.1 \\ 6.2 \\ 6.3 \\ 6.4 \end{array} $	Performance of 3D Jacobi on Blue Gene/L (CO mode) Hops for 3D Jacobi on Blue Gene/L (CO mode) Average Hops for LeanCP on Blue Gene/L for the naïve mapping Average Hops for LeanCP on Blue Gene/L for the topology mapping	36 37 41 41
8.1 8.2 8.3 8.4	Performance of 3D Jacobi on Blue Gene/L (VN mode).Hops for 3D Jacobi on Blue Gene/L (VN mode).Performance of 3D Jacobi on Cray XT3.Hops for 3D Jacobi on Cray XT3.	51 51 52 52
9.5	Performance of LeanCP using the WATER_32M_70Ry benchmark on Blue Gene/L	53
9.0	on Blue Gene/L	53

List of Abbreviations

- NAMD Nanoscale Molecular Dynamics
- CPAIMD Car-Parrinello *ab-initio* Molecular Dynamics
 - PPL Parallel Programming Laboratory
 - IBM International Business Machines
 - PSC Pittsburgh Supercomputing Center
 - 3D 3-dimensional
 - VP Virtual Processor
 - PE Physical Processor
 - 2D 2-dimensional
 - et al. et alii (and others)
 - RTS runtime system
 - 1D 1-dimensional
 - 6D 6-dimensional
 - API Application Programming Interface
 - a.k.a. also known as
 - FFT Fast Fourier Transform
 - 4D 4-dimensional
 - MD Molecular Dynamics
 - etc. et cetera
 - PME Particle Mesh Ewald
 - ORB Orthogonal Recursive Bisection

- BG/L Blue Gene/L
 - CO co-processor mode
 - VN virtual node mode
 - vs. versus
- ApoA1 Apolipoprotein-A1
- F_1 -ATPase Adenosine Triphosphate Synthase

List of Symbols

overlap matrix Psi
GSpace
number of states
number of planes in g-space
number of planes
number of density points in g-space
number of sub-planes
types of atoms
RealSpace
RhoG
RhoR
PairCalculator
Rydberg
Megahertz
Megabyte
Gigahertz
Gigabyte
Angstrom

1 Introduction

Computation and communication are the two aspects of a parallel program which decide its efficiency and performance. Computation needs to be divided evenly among processors to achieve perfect load balance. Communication on the other hand needs to be minimized across processors to ensure minimum overhead. These two tasks are not independent of each other and need to be performed together.

To minimize communication, we need to place communicating objects on the same physical processor. This might not be feasible always if we wish to avoid overloading a particular processor with computational work. In such cases, communicating objects should be placed on processors which are close in terms of the number of network links a message has to travel to go from one to the other. This might not be possible for irregular or flat topologies but is possible for clearly defined three-dimensional (3D) topologies. Let us look at a few of them and analyze how information about such topologies can be utilized to our benefit during a program run.

1.1 Parallel Machines and Topologies

Most supercomputers today have their processors connected through a multi-dimensional interconnect network. The network can be a 3D mesh, a 3D torus or a butterfly or fat-tree network or something different. New supercomputers might have complex topologies which are a hybrid of simple ones. We need to understand the topology of the machine to map objects onto physical processors effectively.



Figure 1.1: A three-dimensional torus network (All connections are not shown for the sake of simplicity)

For example, in a 3D mesh network, processors are connected to six other processors and two arbitrary processors communicate using messages which pass through multiple processors. A 3D torus (Figure 1.1) is similar to a 3D mesh with the difference that the processors at the ends of the mesh are connected to each other and are just one hop away. As seen in the figure, the processor in blue has six neighbors at distance one (colored in green). In a mesh, this processor would only have three neighbors at distance one. These extra connections from end to end reduce the bisection bandwidth of the network by half. IBM's Blue Gene/L, Blue Gene/P, Cray's XT3 and XT4 have a 3D torus interconnect. Processor topologies will be discussed in detail in Section 2.1.

The fastest and biggest supercomputers today are connected by three-dimensional torus interconnects. As we scale to millions of processors, newer and more radical topologies are sure to emerge. If the topology of an interconnect is clearly defined, it is possible to minimize communication volume and balance it evenly across processors for most applications. We will now try to motivate this thesis by explaining how topology can have a significant effect on the communication and hence parallel performance of a program.

1.2 Effect of Topology on Communication

Without a knowledge of the topology of a machine, we would place entities of a parallel program arbitrarily. However, this might lead to unoptimized communication among the entities. If we can co-locate communicating objects close to each other on nearby processors, we can minimize communication volume.

Communication volume is characterized by hop-bytes [1] which is in turn dependent on the hop-count. **Hop-count** is the number of hops (processors) through which a message has to jump to go from one processor to another. **Hop-bytes** are obtained by multiplying the hop-count for a message by its message size. The sum of hop-bytes for all messages for a program gives us its total communication volume. Ideally we want all communication to be local to each processor so that this volume is zero. However this is not possible in many scenarios (for example, if there is all-to-all communication). Hence, we aim at minimizing this volume by placing communicating objects on nearby processors.

To achieve the above-mentioned effect, we require two kinds of information during the actual program run: 1. communication graph of the parallel entities in a program and, 2. topology information about the processors being used on the particular machine. This bring us to the choice of a parallel language which helps us express both easily. The parallel language and framework used in this thesis is CHARM++. We now introduce CHARM++ and some applications written in CHARM++ which are used in this work for implementation of the ideas and algorithms, and for obtaining results.

1.3 Charm++ and its Applications

The CHARM++ [2, 3] parallel language and runtime system is based on an objectoriented parallel programming model. CHARM++ is message-driven and based on the idea of virtualization. Virtualization is the idea of decomposing a problem into a large number of small parts where this number is generally a lot more than the number of processors. Mapping these small parts (virtual processors or VPs) to physical processors (PEs) leads to an adaptive overlap of computation and communication. It also provides us with a framework for obtaining the communication properties of the application and flexibility of mapping VPs to PEs. This framework will be discussed in Chapter 3.

As a proof of principle, to demonstrate the effectiveness of topology mapping, we use a simple application written in CHARM++. It is a 7-point 3-dimensional stencil (referred to as 3D Jacobi henceforth in this thesis) which does regular communication with its neighbors. The two *real* applications which have benefitted from the research in this thesis are NAMD and LEANCP. NAMD [4, 5] is a classical molecular dynamics application which involves calculation of forces and velocities of atoms in a system. It has both migratable (objects that can be moved across processors) and non-migratable (objects which stay on a fixed processor during the entire run) objects. NAMD does an initial assignment of the non-migratable objects. The migratable objects are assigned by a load balancer to the processors which is critical for good performance. In this thesis, we look at the benefits of topological placement of these migratable objects close to their interacting non-migratable counterparts.

LEANCP [6, 7] is a quantum chemistry application also written in CHARM++. It is an implementation of the Car-Parrinello *ab initio* molecular dynamics (CPAIMD) algorithm. LEANCP has several arrays of objects which communicate in a disciplined fashion among their own members and with members of other arrays. This gives us a fairly regular communication graph which should be relatively easy to map on to a processor topology. But there are complex intertwined dependencies among these objects which make this problem difficult. We try to map communicating objects of the same array close to one another and also to other array objects which are involved.

Further chapters in this thesis are organized as follows: We begin with a description of processor topologies and role of communication in a program in Chapter 2. We discuss different communication scenarios and analyze which ones can be optimized in certain ways. We also list some of the previous work in this area. We then move on to describe the CHARM++framework in Chapter 3. Here we discuss CHARM++arrays and how they are mapped. We also discuss the topology interface which provides information about the machine and the allocated partition at runtime. 3D Jacobi is discussed in Section 3.4. Topology aware mapping for this application is also discussed there. Chapters 4 and 5 discuss the ideas implemented in LEANCP and NAMD respectively to improve their performance. Results which prove and exemplify the effectiveness of topology-awareness are presented in Chapter 6 and future work is discussed in Chapter 7.

2 Processor and Communication Graphs

To set the context for our work, we discuss the common processor and communication graphs which we come across nowadays. The most powerful supercomputers today are connected by a 3D torus interconnect network. However, parallel machines can have different kinds of interconnect topologies ranging from fat-tree to hypercube networks. Communication graphs can range from most general unstructured kind to very specific regular ones. It is important to understand different communication scenarios to do hop-count calculations and optimize the mapping algorithms.

2.1 Processor Topologies

Processors in a parallel machine are connected physically in some fashion. These connections decide the topology of the interconnect joining them. We are going to omit a discussion of networks like crossbar switching and bus networks which do not scale to machines with thousands of processors. Below is a description of the common topologies used in machines today:

n-Dimensional Mesh In a n-dimensional mesh network, each processor is connected to two other processors in each physical dimension. This gives a total of 2n connections for each processor. The most common case is n = 3 which is called a 3D mesh. If the size of the mesh in each dimension is N, the diameter of the mesh is $3 \times (N - 1)$. Figure 2.1 (a) shows a 2-dimensional (2D) mesh network.



Figure 2.1: (a) A 2-dimensional mesh network, (b) A 2-dimensional torus network



Figure 2.2: A 4-dimensional hypercube network

- **n-Dimensional Torus** A n-dimensional torus is a mesh with the processors on the end of each dimension connected together. This reduces the diameter of the network by half. Figure 2.1 (b) shows a 2D torus network.
- **n-Dimensional Hypercube** In a n-dimensional hypercube each of 2^n processors are connected to n other processors. A 4-dimensional hypercube can be seen in Figure 2.2. A (n+1) dimensional hypercube can be constructed by connecting the corresponding processors in a n dimensional hypercube.

Some of the biggest supercomputers today have a 3D mesh or torus interconnect network. It is also becoming common to have multiple processing elements (cores) per physical node of the torus giving a fourth dimension for mapping. Let us discuss the interconnects for the two machines used for obtaining results in this thesis: IBM's Blue Gene/L and Cray XT3. Blue Gene/L has a 3D torus interconnect [8]. Each midplane (512 nodes) of the machine is a complete torus of dimension $8 \times 8 \times 8$. Midplanes when joined together form bigger toruses. Each node has two processors giving rise to 1024 processors in a single midplane. Blue Gene/P has a similar network configuration but it has 4 processors per node. This gives 2,048 processors in a single midplane.

Cray XT3 processors are connected by a proprietary SeaStar 3D mesh interconnect [9]. It is also reconfigurable as a 3D torus interconnect but only when using the entire machine. Any smaller subset gives us a 3D mesh. It is also different in the sense that all dimensions of the torus are different and not even. The size of the torus on the Cray XT3 at PSC (called BigBen) is $11 \times 12 \times 16$ in the X, Y and Z dimensions.

2.2 Communication Scenarios

We now proceed to discuss different communication scenarios which occur in benchmarks and real-life applications and then explain the ones which we will concentrate upon in this thesis.

2.2.1 Regular and Irregular Communication

Based on whether each object interacts with a fixed number or variable number of objects, we can classify the cases into:

1. **Regular Communication:** In this case, each object communicates with a fixed number of objects. Let us consider the example of a 2D stencil where each element in the array is updated by using values from four of its neighbors. For this benchmark, it is known that at every step, each element will communicate with four other elements. Such cases are the most simple and

easy to pin down to physical processors in a topological fashion. The application LEANCP is also an example of the same category because each array element communicates with a fixed number of objects which can be identified by analyzing the algorithm.

2. Irregular Communication: It often happens that the application changes the properties of the communicating objects which in turn changes the objects they communicate with. For example, in a molecular dynamics application, atoms migrate to newer objects and hence the communicating neighbors of both the old and the new object change. Such a scenario is referred to as irregular communication.

2.2.2 Point-to-Point, Multicasts and Reductions

Based on the degree of communication which essentially refers to if data is being sent to/by multiple objects or data is being sent on a one-to-one basis, we can classify the scenarios into:

- 1. **Point-to-point Communication:** When objects communicate with each other by sending one-on-one messages, it is referred to as point-to-point communication. It is fairly straightforward to understand and map the communication graph in this case.
- 2. Multicasts: When an object sends the same data to different objects, then it is said to be performing a multicast. Broadcast is a special case of multicast where the message is sent to all the objects in a program. Information about multicasts helps in doing optimizations and minimizing communication. For example, in most cases a multicast can be done through a tree. This does not reduce the number of messages but reduces the load on a particular object and hence contention around a particular processor.

3. **Reductions:** In this case also multiple objects are involved but the phenomenon observed is just the opposite of what happens in a multicast. Several objects send data to one object which collects all this data and manipulates it further.

For the applications we are going to consider in this thesis, we will encounter all the scenarios mentioned above. 3D Jacobi is an example of simple, regular pointto-point communication. In LEANCP, communication is regular but we have pointto-point messages, multicasts and reductions. NAMD has irregular communication and also uses point-to-point messages, multicasts and reductions.

2.3 Previous Work

There has been substantial research on the general problem of mapping a communication graph to a processor graph. The problem has been shown to be NPcomplete [10, 11, 12] and so efforts have been made in two directions to conquer it: heuristic algorithms and physical optimization techniques. As early as 1981, Bokhari [10] came up with an algorithm of pairwise exchanges to reduce interprocessor communication. Results were shown for a specific array processor (the finite element machine). Lee and Aggarwal [11] came up with a greedy algorithm which used initial assignment and pairwise exchange. Results were presented for a 8-node and 16-node hypercube system graph.

Physical optimization techniques like simulated annealing and genetic algorithms have also been used. Bollinger and Midkiff [13] propose a two-phase annealing approach to schedule traffic along network links. Arunkumar and Chockalingam [14] propose a genetic approach which combines the benefits of global search algorithms with local search heuristics. They claim to improve upon the genetic algorithms in terms of the mappings produced and the time taken to obtain them which is generally large in such cases. An interesting idea to reduce the time taken is proposed by Mansour et. al [15] where graph contraction and then interpolation are used to find solutions for large graphs.

The problem has also been studied for specific topologies and/or task graphs which comes closer to the work discussed in this thesis. Ercal et. al [16] use a recursive task allocation scheme based on the Kernighan-Lin mincut bisection heuristic for a hypercube. Agarwal et. al [1, 17] try to solve the mapping problem for processes with persistent load patterns using load balancing for 3D torus-like topologies. Results are shown through simulation studies for two benchmarks, Jacobi and LeanMD [18]. They also show actual results for a 2D Jacobi-like benchmark on up to 1,000 processors of Blue Gene/L. Performance improvement is about two times compared to a random mapping. In this thesis, we present results for 3D Jacobi on up to 16,384 processors of Blue Gene/L and 2,048 processors of Cray XT3. Performance improvement is up to seven times compared to the default mapping for some runs. In addition, we also show results for two production codes, NAMD and LEANCP on the aforementioned machines.

In this light, it becomes important to also mention previous research done specifically for optimizing task-layout on these two machines. Bhanot et. al [19] propose a general method for optimizing task layout on the Blue Gene/L machine and present results for two applications, SAGE and UMT2000 on up to 2,048 nodes. Another interesting study by Weisser et. al [9] studies the performance impact of fragmentation of the Cray XT3 machine by the resource allocator.

3 Objects in Charm++

CHARM++ [2, 3, 20, 21] is an object-oriented parallel programming framework. It includes a programming language based on C++ and an adaptive runtime system. CHARM++ is based on the idea of virtualization. Virtualization is the idea of dividing the problem into multiple virtual processors (VPs) which are mapped to physical processors (PEs) by an intelligent runtime system. The number of VPs is typically much larger than the number of PEs (which makes the degree of virtualization greater than one).

In the CHARM++ programming model, decomposition of the problem is left to the programmer; mapping and communication are handled by the runtime system (RTS). The programmer decomposes the problems into objects and models communication between them as remote method invocations. Method invocation on objects on other processors is translated into messages by the RTS. Figure 3.1 shows the programmer's and system's view of a program being executed by the CHARM++ RTS. The user is concerned only with objects and dependencies between them. The RTS translates these dependencies into messages between objects which are on dif-



Figure 3.1: Programmer's and System's view of the same program under execution

ferent processors. When a remote method is invoked in CHARM++ (referred to as an "entry method"), the RTS sends a message to the concerned processor. Each processor has a queue into which this message gets queued and is executed when this message is processed by the scheduler. Thus, CHARM++ is an asynchronous message-driven message-passing model.

3.1 Chares and Arrays

The two most important kinds of objects in CHARM++ are called Chares and Chare Arrays. A *Chare* is the basic unit of computation in CHARM++. It can be created on any processor and accessed remotely (through entry methods). A *Chare Array* is a collection of chares indexed by some index type. They can be considered similar to any other arrays we come across in programming languages. We will refer to them as chare arrays, CHARM++ arrays or simply arrays interchangeably. Each element of a chare array is called an array element. CHARM++ supports 1-dimensional (1D) to 6-dimensional (6D) arrays.

3.2 Dynamic Mapping Framework

It is important to understand how chares and chare arrays are mapped by the RTS during a run. CHARM++ does a default mapping of chares and chare arrays to processors. For 1D arrays, it does a round-robin mapping while for 2D to 6D arrays, it calculates a hash function using the indices and takes the remainder of division with a big prime number (specifically, 1280107).

CHARM++ provides the user with the flexibility to decide his own mapping for an array of chares and pass it on to the RTS before array creation. The user needs to inherit from the CkArrayMap class and override the virtual function "procNum" which is called by the RTS when it is looking for the processor on which a particular object resides. This map can be passed to the RTS when calling the constructor for array creation.

3.3 Topology Interface in Charm++

Apart from the mapping interface, we also need information about the topology of the machine. We need information like the dimensions of the 3D mesh/torus, whether we have a torus (wraparound) in each dimension and functionality to get coordinates on the torus for a processor rank and vice-versa. This was implemented for Blue Gene/L for specific research in [22, 23], On Blue Gene/L, this information is available in a data structure called "BGLPersonality" and can be accessed using some system calls.

With multiple machines coming up with similar topologies, a need was felt to make this interface more user-friendly and to hide machine-specific details from the user. So, we implemented an API which provides the application with simple calls to obtain information about the topology of the machine without having to worry about the particular machine. This API now gives meaningful information on Blue Gene/L, Cray XT3 and Blue Gene/P. The implementation details about this interface can be found in Appendix C.

3.3.1 Cray XT3

Obtaining topology information is not straightforward on Cray XT3. There are no system calls which can provide information about the dimensions of the partition which has been allocated during a run. With help from technical support at Pittsburgh Supercomputing Center (PSC), we derived this information in several steps. Every node on the XT3 has a unique node ID. A static routing table is available on the machine which has the physical coordinates and neighbors for every node. The CHARM++ RTS reads this file during program start-up. To get the physical coordinates corresponding to a processor rank, we obtain the node ID for the rank through a system call and then get the coordinates from the routing table. Once we have the coordinates for all processors in an allocation, we can get the dimensions of the torus by looking at its diagonally opposite corners.

3.4 3D Jacobi

We begin with the first application which has been used in this thesis to demonstrate the benefit from topology-aware mapping. 3D Jacobi is an implementation of a 3dimensional 7-point stencil in CHARM++. We have a three dimensional array of doubles. All elements are initialized with non-zero values. In every iteration, each element of the array updates itself by computing the average of its six neighbors (two in each dimension) and itself.

To parallelize the computation using CHARM++, we create a 3D chare array. Each element of this chare array is responsible for the computation of some contiguous elements of the data array. The data array is divided into smaller 3D partitions and nearby elements on the chare array get nearby 3D boxes (Figure 3.2). These chares communicate with their neighbors to exchange the updated data on the boundaries. Let us now see the mapping of these chares onto processors.

3.4.1 Topology Aware Mapping

The CHARM++ runtime does the mapping of virtual processors to physical processors by default. Henceforth, we will refer to this mapping as the naïve or default mapping. The RTS has no knowledge of the topology of the machine by itself and so its mapping is totally oblivious to the communication in the application and the



Figure 3.2: Topology-aware mapping of 3D Jacobi's data array onto the 3D processor grid. Different colors (shades) signify which chares get mapped to which processors

topology of the machine.

3D Jacobi has a fairly simple communication pattern. Each chare object talks to its six neighbors (two in each dimension) to exchange the boundary elements. If we can place these objects on the same or nearby processors, we can minimize the distance travelled by each message. The idea is to divide the 3D chare array into equal-sized boxes and then map those to corresponding processors on the 3D torus (Figure 3.2). Let us take a concrete example: say, we have a data array of size 512³ and the size of the torus is 8³ (which gives us 512 processors). Let each chare get a partition of size 16³ from the data array which gives us $512^3/16^3 = 32^3$ chares. Now we need to map a 3D chare array of size 32^3 on a torus of dimensions 8³. Hence each processor gets $512^3/16^3 = 4^3 = 64$ chares. We start from one end of the chare array and place partitions of size 4^3 on each processor considering the topology of the machine. The mapping is similar to superimposing a 3D object over another. The different shades in the figure show how the data array gets mapped to the torus.



Figure 3.3: Hop-count measurement for 3D Jacobi running on Blue Gene/L (using one processor per node)

3.4.2 Evaluation of Mapping Strategies

Using the topology mapping we discussed above, running on 8,192 processors of Blue Gene/L, we get nearly 7.5 times improvement in performance compared to the naïve mapping. Also, the default mapping does not scale very well as we increase the processor count but topology-aware mapping scales beautifully. We will discuss more about the performance improvement in Section 6.1.

To measure the improvement analytically, we need to quantify the reduction in hop-bytes. Since the size of each message in this application is same, so we can look at the hop-counts and still get the same idea. For each run, we calculate the total hops travelled by all messages in one iteration. Figure 3.3 shows the hop-counts for running the application on 512 to 8,192 processors of the Blue Gene/L machine. We get a 8 to 9 times improvement in the hop-count in general and at 8,192 processors the hop-count is reduced by 14 times compared to the naive mapping. This can be attributed to the size of torus becoming large enough $(16 \times 32 \times 16)$ at this point that it makes a huge difference between a random mapping which throws objects all over the torus and a topology-aware mapping.

For the topology scheme used above, it is fairly easy to calculate the number of hops mathematically. Let us say, we have a chare array of size N^3 and processor torus of size P^3 . This gives a chare array of size $N^3/P^3 = C^3$ on each processor. Within this $C \times C \times C$ box of chares, the only chares that communicate externally are the ones on the surface (on the faces, edges and corners). Each processor sends one message per chare on the face, one more for the chares on the edges and a third one for the chares on the corners. Hence total number of messages per processor = (C-2) * (C-2) * 6 + (C-2) * 8 * 2 + 8 * 3a. For the example above, we have 32^3 chares on 8^3 processors which means 4^3 chares per processor. Thus each processor has a box of $4 \times 4 \times 4$ chares. Hence number of messages sent by each processor every iteration is $4 \times 6 + 2 \times 8 \times 2 + 8 \times 3 = 80$. This gives a total of $512 \times 80 = 40960$ hops per iteration.

4 LeanCP and Regular Communication

An accurate understanding of phenomena occurring at the quantum scale can be achieved by considering a model representing the electronic structure of the atoms involved. The Car-Parrinello *ab initio* Molecular Dynamics (CPAIMD) method [24, 25, 26, 27] is one such algorithm which has been widely used to study systems containing $10 - 10^3$ atoms. The implementation of CPAIMD in CHARM++ is called LEANCP [6, 7] (a.k.a. OpenAtom, which is the name it will bear after its production release). To achieve a fine-grained parallelization of CPAIMD, computation in LEANCP is divided into a large number of virtual processors which enables scaling to tens of thousands of processors. We will look at the parallel implementation of this technique, understand its computational phases and the communication involved and then analyze the benefit from topology-aware mapping of its objects.

4.1 Parallel Implementation

In an ab-initio approach, the system is driven by electrostatic interactions between the nuclei and electrons. Calculating the electrostatic energy involves computing several terms: (1) quantum mechanical kinetic energy of non-interacting electrons, (2) Coulomb interaction between electrons or the Hartree energy, (3) correction of the Hartree energy to account for the quantum nature of the electrons or the exchange-correlation energy, and (4) interaction of electrons with atoms in the system or the external energy. Hence, CPAIMD computations involve a large number of phases (Figure 4.1) with high interprocessor communication. These phases are



Figure 4.1: Flow of control in LEANCP

discretized into a large number of virtual processors which generate a lot of communication, but ensures efficient interleaving of work. The various phases are:

- Phase I: In this phase, the real-space representation of the electronic states is obtained from the g-space representation through a transpose based 3dimensional Fast Fourier Transform (FFT).
- **Phase II:** Electron density in real-space is obtained via reductions from the real-space state representation.
- **Phase III:** Fourier components of the density in g-space are created from the corresponding copy in real-space through a 3D FFT.
- Phase IV: Once we have the g-space copy, it is used to compute the "Hartree and external energies" via multiple 3D FFTs which can be performed independently.
- **Phase V:** The energies computed in the previous phase are reduced across all processors and send to the corresponding planes of the different states through

multicasts. This is exactly reverse of the procedure used to obtain the density in phase II.

- Phase VI: In this phase, the forces are obtained in g-space from real-space via a 3D FFT.
- Phase VII: For functional minimization, force regularization is done in this phase by computing the overlap matrix Lambda (Λ) and applying it. This involves several multicasts and reductions.
- Phase VIII: This phase is similar to Phase VII and involves computation of the overlap matrix Psi (Ψ) and its inverse square root (referred to as the S → T process) to obtain "reorthogonalized" states. This phase is called orthonormalization.
- Phase IX: The inverse square matrix from the previous phase is used in a "backward path" to compute the necessary modification to the input data. This again involves multicasts and reductions to obtain the input for phase I of the next iteration.
- Phase X: Since Phase V is a bottleneck, this phase is interleaved with it to perform the non-local energy computation. It involves computation of the kinetic energy of the electrons and computation of the non-local interaction between electrons and the atoms using the EES method [28].

For a detailed description of this algorithm please refer to [7]. We will now proceed to understand the communication involved in these phases through a description of the various chare arrays involved and dependencies among them.

4.2 Communication in LeanCP

The ten phases described in the previous section are parallelized by decomposing the physical system into 15 chare arrays of different dimensions (ranging between one and four). A description of these objects and communication between them follows:

- 1. **GSpace and RealSpace:** These represent the g-space and real-space representations of the electronic states. They are 2-dimensional arrays with states in one dimension and planes in the other. They are represented by G(s,p) $[n_s \times N_g]$ and R(s,p) $[n_s \times N]$ respectively. GSpace and RealSpace interact through transpose operations in Phase I and hence all planes of one state of GSpace interact with all planes of the same state of RealSpace. RealSpace also interacts with RhoR through reductions in Phase II.
- 2. **RhoG and RhoR:** They are the g-space and real-space representations of electron density and are 1-dimensional (1D) and 2-dimensional (2D) arrays respectively. They are represented as $G_{\rho}(p) [N_{g\rho}]$ and $R_{\rho}(p, p') [(N/N_y) \times N_y]$. RhoG is obtained from RhoR in Phase III through two transposes.
- 3. RhoGHart and RhoRHart: RhoR and RhoG are used to compute their Hartree and exchange energy counterparts through several 3D FFTs (in Phase IV). This involves transposes and point-to-point communication. RhoGHart and RhoRHart are 2D and 3D arrays represented by $G_{HE}(p, a)$ [$N_{gHE} \times n_{atom-type}$] and $R_{HE}(p, p', a)$ [(1.4N/N_y) × N_y × $n_{atom-type}$].
- 4. Particle and RealParticle: These two 2D arrays are the g-space and realspace representations of the non-local work and denoted as $G_{nl}(s,p)$ $[n_s \times N_g]$ and $R_{nl}(s,p)$ $[n_s \times 0.7N]$. Phase X for the non-local computation can be overlapped with Phases II-VI and involves communication for two 3D FFTs.
- 5. Ortho and CLA_Matrix: The 2D ortho array, O(s, s') does the postprocessing of the overlap matrices to obtain the T-matrix from the S-matrix. There are three 2D CLA_Matrix instances used in each of the steps of the inverse square method (for matrix multiplications) used during orthonormalization. In the process, these arrays communicate with the paircalculator chare arrays mentioned next.
- 6. PairCalculators: These 4-dimensional (4D) arrays are used in the force regularization and orthonormalization phases (VII and VIII). They communicate with the GSpace, CLA_Matrix and Ortho arrays through multicasts and reductions. They are represented as $P_c(s, s', p, p')$ of dimensions $N_s \times N_s \times N_g \times N'_g$. A particular state of the GSpace array interacts with all elements of the paircalculator array which have this state in one of its first two dimensions.
- 7. Structure Factor: This is a 3D array used when we do not use the EES method for the non-local computation.

4.3 Topology Aware Mapping

LEANCP provides us with a scenario where the load on each virtual processor is static (under the CPAIMD method) and the communication is regular and clearly understood. Hence, it should be fairly straightforward to intelligently map the arrays in this application to minimize inter-processor communication and keep load balanced. As discussed in Chapter 3, CHARM++ does a default mapping of objects (elements of chare arrays) to processors. Let us see how we can do better than the default mapping by using the communication and topology information at runtime. We begin with the two most important arrays in LEANCP which represent electronic states in g-space and real-space.



Figure 4.2: Mapping of different arrays to the 3D torus of the machine

4.3.1 Mapping GSpace and RealSpace Arrays

GSpace and RealSpace are 2D arrays with states in one-dimension and planes in the other. These arrays interact with each other through transpose operations where all planes of one state in GSpace, G(s, *) talk to all planes of the same state, R(s, *) in RealSpace (state-wise communication). GSpace also interacts with symmetric and asymmetric PairCalculators. Each plane of GSpace, G(*, p) interacts with the corresponding plane, P(*, *, p, *) of the PairCalculators (plane-wise communication). So, GSpace interacts state-wise with RealSpace and plane-wise with PairCalculators. If we place all the planes of GSpace together (on nearby processors), we favor one communication but disregard the other. Likewise, if we place all states of GSpace together, we favor the other communication. To strike a balance between the two extremes, we use a hybrid map, where we place a subset of planes and states on one

processor.

We start with laying out the GSpace array on the torus and then map other objects based on GSpace's mapping. The 3D torus is divided into rectangular boxes (which we will refer to as "prisms") such that we get prisms equal in number to the planes in GSpace. The longest dimension of the prism is chosen to be the same as one of the dimensions of the torus. Within each prism for a specific plane, the states in G(*, p) are laid out in increasing order along the long axis of the prism. Figure 4.2 shows the GSpace chares at the bottom being mapped along the long dimension of the torus in the center. Once GSpace is mapped, we place the RealSpace chares. We come up with prisms perpendicular to the GSpace prisms which are formed by including processors holding all planes for a particular state of GSpace, G(s, *). The corresponding states of RealSpace, R(s, *) are mapped on to these prisms.

4.3.2 Mapping of Density Objects

RhoR objects communicate with RealSpace plane wise and hence $R_{\rho}(p, *)$ have to be placed close to R(*, p). To achieve this, we start with the centroid of the prism used by R(*, p) and place RhoR chares in proximity to it. RhoG chares, $G_{\rho}(p)$ are mapped near RhoR chares, $R_{\rho}(p, *)$ but not on the same processors as RhoR to avoid bottlenecks.

4.3.3 Mapping PairCalculator Arrays

Since PairCalculator and GSpace chares interact plane-wise, the effort is to place G(*,p) and P(*,*,p,*) together. Chares with indices P(s1,s2,p,p') are placed around the centroid of G(s1,p), ..., G(s1+sgrain,p) and G(s2,p), ..., G(s2+sgrain,p). This minimizes the hop-count for orthonormalization input and output.

The mapping schemes discussed above substantially reduce the hop-count for different phases. They also restrict different communication patterns to specific prisms within the torus, thereby reducing contention and ensuring balanced communication through out the torus. State-wise and plane-wise communication is confined to different (orthogonal) prisms. This helps avoid scaling bottlenecks as we will see in Section 6.2.

5 NAMD and Multicasts

Understanding biomolecular systems has been aided by the development of molecular dynamics applications for simulation of biomolecular reactions. NAMD [4, 5, 29, 30] is a molecular dynamics code which is widely used for molecular dynamics simulations. It was written using CHARM++about ten years ago and has performed well on a variety of machines and molecular systems. With the emergence of large supercomputers, further optimizations have become necessary to scale NAMD to tens of thousands of processors. Since the molecular system is a simulation box with spatial coordinates for each atom, it should be possible to do a topological placement of the atoms to minimize communication. Lets us look at how this is done in NAMD.

5.1 Parallel Implementation

Classical Molecular Dynamics (MD) requires the calculation of forces due to bonds and non-bonded forces (which include electrostatic and Van der Waal's forces). For parallelization, NAMD does a hybrid of spatial and force decomposition to combine the advantages of both [31]. The simulation box is divided into patches each containing a few atoms. For every pair of interacting patches, we create a chare (called a compute) which is responsible for calculating the pairwise forces between the patches. There are different kinds of computes depending on the forces they calculate: bonded (angles, dihedrals, crossterms etc.) and non-bonded computes. The number of patches is considerably less than the number of computes. The



Figure 5.1: Communication between patches and other objects in NAMD

computational work associated with bonded computes is also less compared to nonbonded computes.

Mapping decisions are made in NAMD considering the facts mentioned above. Patches and bonded computes are non-migratable chares which means they do not move from their home processor once assigned. Non-bonded computes on the other hand are migratable and can be moved around during load balancing. NAMD benefits greatly from the adaptive overlap of communication and computation and the load balancing framework of CHARM++. We will analyze this in detail in the subsequent sections and see how topological considerations help at some places.

5.2 Communication in NAMD

The bonded forces and Van der Waals' component of the non-bonded forces are computed by the bonded and non-bonded computes respectively. Each compute is responsible for calculating the force between a pair of patches and hence receives a message each from them. The patches on the other hand have to send the same information to multiple computes, one each for the patches that they communicate with. Hence the patches multicast their data to their computes and receive forces



Figure 5.2: Computes and patches mapped to the processor torus

back via reductions (Figure 5.1). To avoid sending the same data to a processor twice (if a patch interacts with two computes on that processor), NAMD has the idea of "proxies". If a patch interacts with one or more computes on a processor, it has a proxy on that processor to which it sends the data. The proxy sends the data to the computes internally within the processor. The long-range electrostatic forces are computed using the Particle Mesh Ewald (PME) method. This requires the patches to send messages to the PME grid and receive the forces back. Once all forces are received, they are integrated and the atoms migrated to their new positions.

5.2.1 Topological mapping of patches

For large processor runs, the number of patches is considerably less than the number of processors. On a machine like Blue Gene/L, it is possible to place interacting patches and their computes close to one another. The patches are non-migratable and we need to decide an initial static mapping for them. Ideally, we would like to place the patches maximally away so that the simulation box occupies the whole torus and then place the computes between them.

A strategy to divide the simulation box and place it on the 3D torus of Blue Gene/L is discussed in [22, 23]. An orthogonal recursive bisection (ORB) is done on the 3D torus until we get partitions equal to the number of patches. Then we map the spatially divided simulation box onto the processor partitions. Figure 5.2 shows the mapping of patches, bonded and non-bonded computes on to the 3D torus. This scheme has now been made general using the topology interface in CHARM++ and can now be used on any machine with topology information.

Once the patches are mapped, the non-migratable *bonded* computes are placed with no topological consideration. Non-bonded computes are also placed likewise. But these create load imbalance and hence are reshuffled soon within a major load balancing step. We discuss the NAMD load balancers next.

5.3 Load Balancing

NAMD depends heavily on the load balancing framework provided by CHARM++ for good performance. Computational load in NAMD is persistent across iterations and hence, load information from previous iterations can be used in future iterations. Every few hundred or thousand iterations, a few iterations are instrumented and the load information from these steps is used during the load balancing step to unload the overloaded processors.

There are two major load balancers at work in NAMD. The first one is a comprehensive load balancer which considers every compute and processor and assigns a mapping for all the computes. This leads to movement of a lot of computes. This is called only once, right after start-up. The second one is a refinement load balancer



Figure 5.3: Choice of the best processor to place a compute. The best choice is on the extreme left where we get two proxies on one processor. The worst is on the right where we cannot find any proxy/patch on the processor.

which considers only overloaded processors and tries to move computes away from them. This is done every few hundred or thousand steps. Both load balancers use a greedy strategy for load balancing. They pick the heaviest compute (or a random compute on the heaviest processor) and try to place it on an underloaded processor. An underloaded processor is one whose load is below the average load multiplied by an overload factor (decided empirically).

In order to minimize communication and achieve load balance together, the load balancers try to minimize the addition of new proxies. Hence, considering all the eligible underloaded processors, they come up with a choice table (Figure 5.3). The best choice is to place the compute on a processor which has two proxies, one each for its two patches; otherwise on a processor with a patch and a proxy, or with both the patches; if such a processor does not exist, then on one with at least one proxy or one patch and so on ... If we cannot find a processor with any of these choices, then we place it on the least overloaded processor we can find. The following metrics help evaluate the performance of a load balancer in NAMD: (1) Maximum load on any processor should be close to the average load, (2) Maximum number of proxies for a patch should be reasonable, (3) Performance should be good, and (4) On a machine with topology information, we can also aim at reducing the number of hops which messages have to travel. That is the focus of our work in this thesis.



Figure 5.4: Topological search for a underloaded processor in the 3D torus

topology-awareness in them.

5.3.1 Topology-aware decisions

During load balancing, when we are trying to find a processor to place the compute on, we can consider the topology of the machine. This can help to reduce the distance between the patches and the proxies they communicate with. If we are unable to find a processor with even one proxy or patch to place the compute on, then we need to create new proxies. In such a case, it is best to create proxies on a processor such that the sum of the distance (in terms of hops) from its two patches is minimized.

To ensure this, we find the coordinates of the two processors which host the patches with which the given compute interacts. We then try to find a processor within the region enclosed by these two processors (see Figure 5.4) on the torus. For any point within this region (which we call the inner brick), the sum of distances from the two patches (at the corners) is same. Hence we try to find the least

overloaded processor within this brick.

If we fail, then we need to try the rest of the torus (called the outer brick). We spiral around the inner brick on the outside and try to find the first underloaded processor we can. As soon as we find one, we place the compute on it. We will see in Section 6.3, the benefit from this optimization in terms of reduction in hop-bytes and better performance.

6 Results

In this chapter, we discuss and analyze the results of the schemes developed in the previous chapters for topology sensitive mapping and load balancing for different applications. Results are presented for 3D Jacobi, LEANCP and NAMD on up to 16,384 processors of Blue Gene/L and 2,048 processors of Cray XT3.

A brief description of the two machines follows:

- IBM's Blue Gene/L: The Blue Gene/L machine at IBM T J Watson (referred to as the "Watson BG/L") has 20,480 nodes. Each node contains two 700 MHz PowerPC 440 cores and has 512 MB of memory shared between the two. The nodes on Blue Gene/L are connected into a 3D torus. We used the Watson BG/L for most of our runs. Blue Gene/L can be run in two modes: co-processor or CO mode where we just use one processor per node for the computation and virtual node or VN mode where we use both processors per node.
- Cray's XT3: BigBen at Pittsburgh Supercomputing Center (PSC) has 2,068 compute nodes each of which has two 2.6 GHz AMD Opteron processors. The two processors on a node share 2 GB of memory. The nodes are connected into a 3D torus by a custom SeaStar interconnect. We do not get a contiguous allocation of processors on XT3 if we use the default queue. We had to take help from the PSC staff to set up a reservation to get a mesh of 8 × 8 × 16 which is 1,024 nodes and 2,048 processors.

	$8 \times 8 \times 8$		16 imes16 imes16		32 imes 32 imes 32	
Processors	Naïve	Topology	Naïve	Topology	Naïve	Topology
512	128.93	67.00	44.22	28.78	33.55	20.60
1024	69.41	34.02	22.59	15.17	19.50	11.17
2048	35.59	19.03	13.22	8.09	10.86	5.92
4096	20.07	11.58	7.44	4.38	14.22	2.25
8192	13.99	6.95	7.52	2.39	9.95	1.33

Table 6.1: Performance of 3D Jacobi on Blue Gene/L (CO mode) for different decompositions (Problem Size: fixed at $512 \times 512 \times 512$). The numbers are time in seconds for 1000 iterations.

	$8 \times 8 \times 8$		16 imes16 imes16		32 imes 32 imes 32	
Processors	Naïve	Topology	Naïve	Topology	Naïve	Topology
512	128.77	66.77	44.38	28.85	33.56	20.59
1024	139.94	68.88	45.45	29.12	38.44	20.61
2048	143.69	73.01	46.40	29.67	33.98	20.68
4096	178.35	93.51	55.50	33.99	34.13	20.62
8192	239.25	150.35	120.02	48.08	113.88	21.10

Table 6.2: Performance of 3D Jacobi on Blue Gene/L (CO mode) for different decompositions (Problem Size: variable from $512 \times 512 \times 512$ to $1024 \times 1024 \times 1024$). The numbers are time in seconds for 1000 iterations.

6.1 3D Jacobi

Static topology mapping of 3D Jacobi performs very well because of the same number of dimensions of the data and the torus. Communication is regular and all messages are of equal size. Each chare has the same amount of computation and hence the problem is completely load balanced too.

6.1.1 Performance on Blue Gene/L

We tried various experiments with 3D Jacobi to see the effects of virtualization (number of chares), effect of naïve vs. topology-aware mapping and effect of CO vs. VN mode. Tables 6.1 and 6.2 present the performance on Blue Gene/L in CO mode for different problem sizes and decompositions. We tried three different sizes



Figure 6.1: Performance of 3D Jacobi on Blue Gene/L (CO mode)

of chares: each containing 8^3 or 16^3 or 32^3 elements of the data array. Table 6.1 shows the time taken for 1000 iterations of 3D Jacobi for a fixed problem size of 512^3 . Size of the chares decided the number of chares per processor and in total. Chare size of 8^3 gives us $8^3 = 512$ chares per processor for a 512-processor run which is on the higher side. Chare size of 32^3 gives us 8 chares per processor which is more reasonable. For the naïve mapping, the performance at the 8K point for 16^3 decomposition and 4K point for 32^3 decomposition is dismal because we are left with just one chare per processor which leaves no room for overlap between computation and communication. This leaves the chares waiting for messages to arrive. This is not a bottleneck in the case of topology mapping because the messages are only one hop away. Hence, we see almost perfect scaling for topology mapping (for 32^3 chare size) up to 8K processors (efficiency of 96.8 %).

Table 6.2 shows weak-scaling for the problem where we doubled the problem size (doubling one dimension of the data array each time) as we doubled the number of processors. Here we have the same number of chares across all processor runs for a given decomposition. The number of chares per processor for 8^3 , 16^3 and 32^3



Figure 6.2: Hops for 3D Jacobi on Blue Gene/L (CO mode)

	Perfe	ormance	Hop	-count
Processors	Naïve Topology		Naïve	Topology
512	98.96	44.95	204180	24576
1024	58.81	23.74	222916	26624
2048	46.43	13.75	274384	30720
4096	19.38	8.32	356600	38912
8192	32.95	4.79	668904	77824
16384	13.36	2.30	1118128	96256

Table 6.3: Performance and hop-counts for 3D Jacobi on Blue Gene/L (VN mode) for block size $32 \times 32 \times 32$ (Problem Size: fixed at $512 \times 512 \times 1024$). The numbers are time in seconds for 1000 iterations.

decompositions is 512, 64 and 8 respectively. The huge number of chares being mapped all over the machine for the naïve mapping accounts for the unreasonably high times for 8K runs. It can be observed that the highly effective topology mapping for the 32^3 chare size leads to similar times for all processor runs. Performance data and hop-counts for the 32^3 decomposition is plotted in Figures 6.1 and 6.2 respectively. Topology mapping leads to a reduction in hops by 8 to 14 times for different processor runs, as noted earlier.

Table 6.3 shows the performance of 3D Jacobi for the most reasonable chare size

	16 imes 16 imes 16					32 imes	32 imes 32	
	Time		Time Hop-count		Time		Hop-count	
Procs	N	Т	Ν	Т	Ν	Т	Ν	Т
256	34.88	22.63	802744	53248	21.06	16.65	102484	13312
512	18.14	12.50	1036388	69632	13.08	8.98	132584	17408
1024	9.13	7.20	1221812	86016	8.75	5.31	153600	21504
2048	5.40	5.65	1404928	118784	4.16	4.42	176196	29696

Table 6.4: Performance and hop-counts for 3D Jacobi on Cray XT3 for different decompositions (Problem Size: fixed at $512 \times 512 \times 512$). N stands for naïve and T stands for topology-aware mapping. The numbers are time in seconds for 1000 iterations.

	16 imes 16 imes 16					32 imes	32 imes 32	
	Time		Hop-count		Time		Hop-count	
Procs	N	Т	Ν	Т	Ν	Т	Ν	Т
256	17.44	11.99	403764	40960	14.39	8.51	52252	10240
512	18.14	12.50	1036388	69632	13.08	8.98	132584	17408
1024	18.87	13.04	2423272	143360	12.81	9.51	308756	35840
2048	20.79	15.00	5571796	294912	13.19	10.96	696924	73728

Table 6.5: Performance and hop-counts for 3D Jacobi on Cray XT3 for different decompositions (Problem Size: variable from $512 \times 512 \times 256$ to $1024 \times 512 \times 1024$). N stands for naïve and T stands for topology-aware mapping. The numbers are time in seconds for 1000 iterations.

of 32^3 on Blue Gene/L in VN mode. Here we use both processors on each node and use a fixed problem size of $512 \times 512 \times 1024$. Also included are hop-counts which validate the benefit achieved by topology mapping compared to naïve mapping. Plots showing the same data can be found in Appendix A.

6.1.2 Performance on Cray XT3

BigBen at PSC was used in dedicated mode for these results to get a contiguous allocation of processors. We did not use the whole machine and for any processor counts less than the full machine, the network of XT3 is not a complete torus in all directions. So the results are for a 3D topology with a torus in some dimensions and mesh in others. Experiments similar to Blue Gene/L were repeated on Cray XT3 to test our mapping schemes and the topology interface written in CHARM++ for XT3 (which was being tested for the first time). We did not do the 8³ chare size experiments because it creates a huge number of chares which is not the best choice for the degree of virtualization. Tables 6.4 and 6.5 show performance numbers for strong and weak scaling for 256 to 2,048 processors. The maximum improvement in performance is only 1.7 times compared to 7.5 times on Blue Gene/L. This might be attributed to the more efficient interconnect of XT3.

But the story of hop-counts is exactly similar to that of Blue Gene/L. We get a reduction of up to ten times in the hop-count. It is worth noting that in Table 6.4 at 2,048 processors, there is no improvement; in fact there is a slight slow down with topology mapping. Since it is difficult to get dedicated time on this machine, we have not been able to analyze this further. We do not see similar problems in Table 6.5, so a possibility is that this might be because of the few chares we have per processor on 2K processors. Appendix A shows the same data in plots.

6.2 LeanCP

We now showcase the benefits of topology mapping on the first real application used for our tests. Liquid water was used as a test case due to the importance of aqueous solutions in biophysics. The two water systems considered consist of 32 and 256 water molecules respectively. The g-space spherical cutoff for the two systems is 70 Ry which is why they are called WATER_32M_70Ry and WATER_256M_70Ry respectively.

Tables 6.6 and 6.7 compare the performance of naïve vs. topology mapping on Blue Gene/L for CO and VN modes respectively. For WATER_32M_70Ry, topology mapping helps overcome the communication bottlenecks at 2K processors in CO

	WATE	R_32M_70Ry	WATE	R_256M_70Ry
Processors	Naïve	Topology	Naïve	Topology
512	0.274	0.259	-	-
1024	0.189	0.150	19.10	16.4
2048	0.219	0.112	13.88	8.14
4096	0.167	0.082	9.13	4.83
8192	0.129	0.063	4.83	2.75
16384	-	-	3.40	1.71

Table 6.6: Performance of LEANCP on Blue Gene/L (CO mode). The numbers represent time per step in seconds.

	WATE	R_32M_70Ry	WATER_256M_70Ry		
Processors	Naïve	Topology	Naïve	Topology	
1024	0.207	0.174	-	-	
2048	0.296	0.130	18.54	18.14	
4096	0.189	0.082	10.61	5.10	
8192	0.189	0.067	7.27	3.48	

Table 6.7: Performance of LEANCP on Blue Gene/L (VN mode). The numbers represent time per step in seconds.

mode. Performance is consistently twice as good as the naïve mapping from 2K to 8K processors. In VN mode, performance at 8K processors is three times better. WATER_256M_70Ry also shows similar improvements in timings with performance being two times better for large processor runs. Plots showing the same data for the two systems can be found in Appendix B.

To validate the benefits of topology mapping for LEANCP, we use the performance analysis tool in CHARM++, called Projections [32]. This tool helps us visualize the average number of hops per entry method and per processor. Figures 6.3 and 6.4 show the average hop-bytes per processor for the naïve and topology mapping. Both runs used the WATER_32M_70Ry benchmark and 2,048 processors of Blue Gene/L in CO mode. As we can see, the average hop-count per processor for the topology case is 7 compared to 9 for the default case. Also, communication is much more uniform in the case of topology aware mapping which balances the



Figure 6.3: Average Hops for LeanCP on Blue Gene/L for the naïve mapping



Figure 6.4: Average Hops for LeanCP on Blue Gene/L for the topology mapping

	CO	Mode	VN	mode
Processors	Naïve	Topology	Naïve	Topology
512	202.39	150.10	-	-
1024	437.01	316.29	505.34	434.62
2048	776.79	514.13	813.06	512.25
4096	1672.93	1193.04	1435.57	1155.41
8192	-	_	2910.08	2290.12

Table 6.8: Reduction in hop-bytes for 3D NAMD on Blue Gene/L (Bench-mark:ApoA1)

	CO	Mode	VN	mode
Processors	Naïve	Topology	Naïve	Topology
512	374.94	374.62	-	-
1024	570.47	570.53	484.54	484.06
2048	1161.59	549.35	710.99	711.31
4096	3933.59	2274.28	1335.08	1363.08
8192	-	-	6544.91	3822.21

Table 6.9: Reduction in hop-bytes for 3D NAMD on Blue Gene/L (Benchmark: F_1 -ATPase)

communication load on each processor and on the network links.

6.3 NAMD

Performance of NAMD was measured for two molecular systems called Apolipoprotein-A1 (ApoA1) and Adenosine Triphosphate Synthase (F₁-ATPase) containing 92,224 and 327,506 atoms respectively. Both benchmarks use a 12 Å cutoff and 1 femtosecond time-step. The size of the simulation box in Å for the two systems is $108.86 \times 108.86 \times 77.76$ and $178.30 \times 131.54 \times 132.36$.

Table 6.8 shows the reduction in hop-bytes as a result of topology-aware mapping of patches and topology-aware load balancing of computes for ApoA1. The numbers given are hop-bytes in MB per iteration added across all processors for all the multicast messages. We get nearly 30% improvement (reduction in hop-bytes) at 4K processors in CO mode. Likewise at this point, we also get an improvement in time-step per iteration from 4.68 to 3.88 milliseconds (ms).

Similar numbers for the F_1 -ATPase system are presented in Table 6.9. There is a reduction in hop-bytes of more than 40% using the topology-aware load balancers at 4K processors. Better performance of the load balancers compared to ApoA1 might be because F_1 -ATPase has four times the number of patches and three times the number of computes compared to ApoA1. A better mapping for a larger system means more reduction in the hop-bytes. Using 8k processors, in VN mode, we get a performance improvement from 9.3 to 7.83 ms per step. For lower processor runs, the improvement is not as much which still needs to be investigated (This work is still in its early stages).

From the results presented in this chapter, it is quite obvious that topology awareness can yield enormous benefits to some applications. We just need to understand the communication patterns of the application and the topology of the machine being used. In the next chapter, we will look at generalizing some of the techniques to make them useful to other applications.

7 Future Work and Conclusion

In this thesis, we have focused on specific parallel applications and analyzed the benefit from topology-aware mapping. The approach has been to understand the parallel implementation and the communication characteristics of the application. Based on the communication graph, we have tried to place communicating objects on nearby processors on a 3D mesh or torus topology. The two different places where we have done this are: 1. defining a static mapping during start-up as in the case of 3D Jacobi and LEANCP, and 2. modifying the load balancers to take the topology of machine into account (as in NAMD). So, our effort has been concentrated on a few specific applications and a 3D mesh or torus topology. We shall now discuss future efforts to make these ideas more general so that they can be used by other applications on diverse topologies.

7.1 Generalized Topology Sensitive Mapping

We were quite successful in mapping different chare arrays in LEANCP by considering their communication dependencies. We wish to extend this idea to do this for any application with regular communication. Given the communication dependencies between chare arrays and the topology, the CHARM++ RTS should automatically do an intelligent mapping. This would remove the burden of mapping from the user and give optimized performance compared to a random mapping.

A more general idea is to successfully do this for any arbitrary communication graph and topology. This has been proved to be NP-complete by Bokhari [10]. Hence the effort would be towards development of heuristics which perform well for most scenarios. This would useful for problems with regular communication like Jacobi, Matrix multiplication and FFTs.

7.2 Load Balancers for Section Multicasts

Multicasts in NAMD are a very special case. All patches send the same data to their multicast targets and each multicast target (compute) receives data from only two patches. In general, a target can receive messages from any number of multicast roots.

We plan to build upon the algorithms developed in the NAMD load balancers to do this for a more general case. We wish to write a load balancer within the CHARM++ load balancing framework which considers generic multicasts and the topology of the machine. Since it would be within the CHARM++ framework, other applications will also be able to use it. This can be used by applications like cosmology (ChaNGa [33]) and meshing applications.

7.3 Summary

Work presented in this thesis demonstrates the benefit of topology-awareness in mapping of objects on to a parallel machine statically or during load balancing. We first describe the topology interface in CHARM++ which facilitates the mapping of objects onto 3D topologies. As a first example, we take a simple application, 3D 7-point stencil and prove that topology mapping does help in reducing contention during communication, thereby reducing the time required for the same. The number of hops travelled by messages is reduced by ten times on the average which gives us a performance boost of two to seven times in certain cases. The two real applications which benefit from the idea of topology-aware mapping are LEANCP and NAMD. LEANCP has regular communication between its numerous chare arrays and thus provides us with a structured well-defined example for topology-aware mapping. Since the application is heavily communication bound, it shows good improvements compared to the naïve mapping. NAMD's load balancers on the other hand depend on many factors for optimal performance. Hence, it becomes difficult to judge the real benefit of topology mapping. Yet NAMD also shows a reasonable improvement in terms of hop-bytes and some improvement in performance for higher processor counts. We hope to utilize the insights gained from the study of these applications to create a generalized automatic framework for topology-aware mapping. This would benefit many applications in the future.

and performance of the application.

References

- Tarun Agarwal, Amit Sharma, and Laxmikant V. Kalé. Topology-aware task mapping for reducing communication contention on large parallel machines. In Proceedings of IEEE International Parallel and Distributed Processing Symposium 2006, April 2006.
- [2] L. V. Kalé, B. Ramkumar, A. B. Sinha, and A. Gursoy. The CHARM Parallel Programming Language and System: Part I – Description of Language Features. *Parallel Programming Laboratory Technical Report #95-02*, 1994.
- [3] L. V. Kalé, B. Ramkumar, A. B. Sinha, and V. A. Saletore. The CHARM Parallel Programming Language and System: Part II – The Runtime system. *Parallel Programming Laboratory Technical Report #95-03*, 1994.
- [4] James C. Phillips, Rosemary Braun, Wei Wang, James Gumbart, Emad Tajkhorshid, Elizabeth Villa, Christophe Chipot, Robert D. Skeel, Laxmikant Kalé, and Klaus Schulten. Scalable molecular dynamics with NAMD. *Journal* of Computational Chemistry, 26(16):1781–1802, 2005.
- [5] James C. Phillips, Gengbin Zheng, Sameer Kumar, and Laxmikant V. Kalé. NAMD: Biomolecular simulation on thousands of processors. In *Proceedings* of the 2002 ACM/IEEE conference on Supercomputing, pages 1–18, Baltimore, MD, September 2002.
- [6] Ramkumar V. Vadali, Yan Shi, Sameer Kumar, L. V. Kale, Mark E. Tuckerman, and Glenn J. Martyna. Scalable fine-grained parallelization of planewave-based ab initio molecular dynamics for large supercomputers. *Journal of Comptational Chemistry*, 25(16):2006–2022, Oct. 2004.
- [7] Eric Bohm, Glenn J. Martyna, Abhinav Bhatele, Sameer Kumar, Laxmikant V. Kale, John A. Gunnels, and Mark E. Tuckerman. Fine Grained Parallelization of the Car-Parrinello ab initio MD Method on Blue Gene/L. *IBM Journal of Research and Development: Applications of Massively Parallel Systems (to appear)*, 52(1/2), 2007.
- [8] M.Blumrich, D.Chen, P.Coteus, A.Gara, M.Giampapa, P.Heidelberger, S.Singh, B.Steinmacher-Burow, T.Takken, and P.Vranas. Design and analysis of the blue gene/l torus interconnection network. *IBM Research Report*, December 2003.

- [9] Deborah Weisser, Nick Nystrom, Chad Vizino, Shawn T. Brown, and John Urbanic. Optimizing Job Placement on the Cray XT3. 48th Cray User Group Meeting 2006 Proceedings, 2006.
- [10] Shahid H. Bokhari. On the mapping problem. *IEEE Trans. Computers*, 30(3):207–214, 1981.
- [11] Soo-Young Lee and J. K. Aggarwal. A mapping strategy for parallel processing. *IEEE Trans. Computers*, 36(4):433–442, 1987.
- [12] P. Sadayappan and F. Ercal. Nearest-neighbor mapping of finite element graphs onto processor meshes. *IEEE Trans. Computers*, 36(12):1408–1424, 1987.
- [13] S. Wayne Bollinger and Scott F. Midkiff. Processor and link assignment in multicomputers using simulated annealing. In *ICPP* (1), pages 1–7, 1988.
- [14] S. Arunkumar and T. Chockalingam. Randomized heuristics for the mapping problem. *International Journal of High Speed Computing (IJHSC)*, 4(4):289– 300, December 1992.
- [15] N. Mansour, R. Ponnusamy, A. Choudhary, and G. C. Fox. Graph contraction for physical optimization methods: a quality-cost tradeoff for mapping data on parallel computers. In *ICS '93: Proceedings of the 7th international conference* on Supercomputing, pages 1–10. ACM, 1993.
- [16] F. Ercal, J. Ramanujam, and P. Sadayappan. Task allocation onto a hypercube by recursive mincut bipartitioning. In *Proceedings of the third conference on Hypercube concurrent computers and applications*, pages 210–221. ACM Press, 1988.
- [17] Tarun Agarwal. Strategies for topology-aware task mapping and for rebalancing with bounded migrations. Master's thesis, Dept. of Computer Science, University of Illinois, 2005.
- [18] Vikas Mehta. Leanmd: A charm++ framework for high performance molecular dynamics simulation on large parallel machines. Master's thesis, University of Illinois at Urbana-Champaign, 2004.
- [19] G. Bhanot, A. Gara, P. Heidelberger, E. Lawless, J. C. Sexton, and R. Walkup. Optimizing task layout on the blue gene/l supercomputer. *IBM Journal of Research and Development*, 49(2/3):489–500, 2005.
- [20] Laxmikant V. Kale, Eric Bohm, Celso L. Mendes, Terry Wilmarth, and Gengbin Zheng. Programming Petascale Applications with Charm++ and AMPI. In D. Bader, editor, *Petascale Computing: Algorithms and Applications*. Chapman & Hall / CRC Press, 2008.

- [21] L. V. Kale and Sanjeev Krishnan. Charm++: Parallel Programming with Message-Driven Objects. In Gregory V. Wilson and Paul Lu, editors, *Parallel Programming using C++*, pages 175–213. MIT Press, 1996.
- [22] Sameer Kumar, Chao Huang, Gheorghe Almasi, and Laxmikant V. Kalé. Achieving strong scaling with NAMD on Blue Gene/L. In Proceedings of IEEE International Parallel and Distributed Processing Symposium 2006, April 2006.
- [23] Sameer Kumar, Chao Huang, Gengbin Zheng, Eric Bohm, Abhinav Bhatele, James C. Phillips, Hao Yu, and Laxmikant V. Kalé. Scalable Molecular Dynamics with NAMD on Blue Gene/L. *IBM Journal of Research and Development: Applications of Massively Parallel Systems (to appear)*, 52(1/2), 2007.
- [24] R. Car and M. Parrinello. Unified approach for molecular dynamics and densityfunctional theory. *Phys. Rev. Lett.*, 55:2471–2474, (1985).
- [25] G. Galli and M. Parrinello. Computer Simulation in Materials Science, 3:283, (1991).
- [26] M.C. Payne, M. Teter, D.C. Allan, T.A. Aria, and J.D. Joannopolous. Iterative minimization techniques for ab initio total energy calculations : molecular dynamics and conjugate gradients. *Rev. Mod. Phys.*, 64:1045, (1992).
- [27] M. E. Tuckerman. Ab initio molecular dynamics: Basic concepts, current trends and novel applications. J. Phys. Condensed Matter, 14:R1297, 2002.
- [28] H.S. Lee, M.E. Tuckerman, and G.J. Martyna. Efficient evaluation of nonlocal pseudopotentials via euler exponential spline interpolation. *Chem. Phys. Chem.*, 6:1827, (2005).
- [29] Abhinav Bhatele, Sameer Kumar, Chao Mei, James C. Phillips, Gengbin Zheng, and Laxmikant V. Kale. NAMD: A Portable and Highly Scalable Program for Biomolecular Simulations. 2007.
- [30] Klaus Schulten, James C. Phillips, Laxmikant V. Kale, and Abhinav Bhatele. Biomolecular modeling in the era of petascale computing. In D. Bader, editor, *Petascale Computing: Algorithms and Applications*, pages 165–181. Chapman & Hall / CRC Press, 2008.
- [31] Laxmikant Kalé, Robert Skeel, Milind Bhandarkar, Robert Brunner, Attila Gursoy, Neal Krawetz, James Phillips, Aritomo Shinozaki, Krishnan Varadarajan, and Klaus Schulten. NAMD2: Greater scalability for parallel molecular dynamics. *Journal of Computational Physics*, 151:283–312, 1999.
- [32] Laxmikant V. Kale, Gengbin Zheng, Chee Wai Lee, and Sameer Kumar. Scaling applications to massively parallel machines using projections performance analysis tool. In *Future Generation Computer Systems Special Issue on: Large-Scale System Performance Modeling and Analysis*, volume 22, pages 347–358, February 2006.

[33] Filippo Gioachin, Amit Sharma, Sayantan Chakravorty, Celso Mendes, Laxmikant V. Kale, and Thomas R. Quinn. Scalable cosmology simulations on parallel machines. In VECPAR 2006, LNCS 4395, pp. 476-489, 2007.

Appendix A



Figure 8.1: Performance of 3D Jacobi on Blue Gene/L (VN mode)



Figure 8.2: Hops for 3D Jacobi on Blue Gene/L (VN mode)



Figure 8.3: Performance of 3D Jacobi on Cray XT3



Figure 8.4: Hops for 3D Jacobi on Cray XT3

Appendix B



Figure 9.5: Performance of LeanCP using the WATER_32M_70Ry benchmark on Blue Gene/L



Figure 9.6: Performance of LeanCP using the WATER_256M_70Ry benchmark on Blue Gene/L

Appendix C

The Topology API in CHARM++ provides the user application with a lot of useful functions which facilitate the implementation of mapping algorithms. It should be noted that these functions are only applicable for 3D mesh and torus topologies:

- void rankToCoordinates(int pe, int &x, int &y, int &z) This function returns the physical co-ordinates on the machine for a given processor rank or ID (assigned by the system).
- int coordinatesToRank(int x, int y, int z) This function returns the system assigned processor rank for given physical co-ordinates (x, y, z) on the machine.
- void sortRanksByHops(int pe, int *pes, int *idx, int n) Given a list of processors, this function sorts the list by their distance in hops from a given processor.
- int pickClosestRank(int mype, int *pes, int n) This function picks the closest processor to a given processor from a given list.
- int areNeighbors(int pe1, int pe2, int pe3, int distance) This function determines if the sum of the distance in hops between three processors is less than a certain distance.

These functions are used in LEANCP and NAMD extensively to simplify the mapping algorithms. For example, to sort a list of processors by their distance from their centroid, we can do the following:

```
void sortByCentroid(int *list, int npes) {
    int pe, x, y, z, x1, y1, z1;
    int idx[npes];
    for(int i=0; i<npes; i++) {
        rankToCoordinates(list[i], x1, y1, z1);
        x += x1;
        y += y1;
        z += z1;
    }
    x = x/npes;
    y = y/npes;
    z = z/npes;
    pe = coordinatesToRank(x, y, z);
    sortRanksByHops(pe, list, idx, npes);
}</pre>
```

Vita

Office Address:	Residential Address:
Thomas M. Siebel Center for Comp. Sc.	505, E White St, Apt. #06
201, N. Goodwin Avenue,	Champaign, IL 61820
Urbana, IL 61801	Ph: +1-217-417-7579
http://charm.cs.uiuc.edu/~bhatele	E-mail: bhatele2@uiuc.edu

Abhinav Bhatele

Educational Qualifications

Doctor of Philosophy in Computer Science	2005 - Present
University of Illinois at Urbana Champaign	CGPA: 3.89/4.0

Bachelor of Technology in Computer Science and Engineering2001-2005Indian Institute of Technology, KanpurCPI: 9.1/10.0

Research Experience

Graduate Research Assistant, Parallel Programming Laboratory, UIUC 2005-Present

Working with Prof. Laxmikant V. Kale in the Parallel Programming Group at UIUC since August 2005.

• I work on molecular dynamics applications called NAMD and LeanCP written

in Charm++. My work is related to topology sensitive mapping and loadbalancing.

• Working on BigSim, a Parallel Discrete Event Simulator developed at PPL.

Summer Intern, IBM T J Watson Research Center, NY, USA May-July 2007

Worked with Dr. Sameer Kumar in the Blue Gene Software Group

• Devised topology sensitive load balancing algorithms to be applied to Charm++ programs in general and NAMD in particular.

Summer Intern, IBM T J Watson Research Center, NY, USA May-July 2006

Worked with Dr. Guojing Cong at the Advanced Computing Technology Center

• Developed a tool for automatic and detailed profiling of programs at finer levels like functions and individual statements.

Summer Intern, INRIA Labs, Nancy, France May-July 2004

Worked with Prof. Dominique Mery and Stephan Merz in the MOSEL group

• Developed a GUI for TLC and Xprove: TLC is a model checker and Xprove is a theorem prover being developed at INRIA Labs.

Summer Intern, IIT Kanpur, India May-July 2003

Worked with Prof. Shashank K. Mehta at IIT Kanpur

• Studied a Picoblaze microprocessor design in VHDL (developed by Xilinx) and implemented and tested it using Verilog.

Recent Publications

- Abhinav Bhatele, Sameer Kumar, Chao Mei, James C. Phillips, Gengbin Zheng, Laxmikant V. Kale, NAMD: A Portable and Highly Scalable Program for Biomolecular Simulations, *PPL Technical Report*, 2007
- Abhinav Bhatele, Guojing Cong, A Selective Profiling Tool: Towards Automatic Performance Tuning, In Proceedings of SMTPS 07, IPDPS, 2007
- Abhinav Bhatele, Shubham Satyarth, Sanjeev K. Aggarwal, Compiler Algorithm Language (CAL): An Interpreter and Compiler, In Proceedings of ACST, 2007
- Klaus Schulten, James C. Phillips, Laxmikant V. Kale, Abhinav Bhatele, Biomolecular modeling in the era of petascale computing, In Petascale Computing: Algorithms and Applications, pp. 165-181, D. Bader, Ed., Chapman & Hall / CRC Press, New York, 2007
- Eric Bohm, Abhinav Bhatele, Laxmikant V. Kale, Mark E. Tuckerman, Sameer Kumar, John A. Gunnels, Glenn Martyna, Fine grained parallelization of the Car-Parrinello ab initio MD method on Blue Gene/L, *In IBM J. Res. Dev.*, *Volume 52, No. 1/2*, 2007
- 6. Sameer Kumar, Chao Huang, Gengbin Zheng, Eric Bohm, Abhinav Bhatele, Jim Phillips, Gheorghe Almasi, Hao Yu, Laxmikant V. Kale, Achieving Strong Scaling with NAMD on Blue Gene/L, In IBM J. Res. Dev., Volume 52, No. 1/2, 2007

Relevant Courses

Graduate – Advanced Computer Architecture (CS533), Formal Methods of Computation (CS475), Parallel Programming Methods (CS498lvk), Programming Languages and Compilers (CS421), Advanced Topics in Compiler Construction (CS526),
Social Computing (CS598kgk), Improving your Research Skills (CS598lrs)

Undergraduate – Advanced Compiler Optimizations (CS738), Computer Architecture (CS422), Compilers (CS335), Computer Networks (CS425), Operating Systems (CS330), Algorithms II (CS345), Theory of Computation (CS340), Data Structures and Algorithms (CS210), Discrete Mathematics (CS201)

Programming Skills

Languages: Charm++, C, C++, JAVA, Verilog, VHDL, Scheme, ML, Ocaml Platforms: Most flavors of Windows and Linux Tools: Lex, Yacc, LaTeX, Make, Perl

Graduate Projects

Optimizing LeanMD on Blue Gene/L – Topology mapping for this basic molecular dynamics code on the Blue Gene machine

Review of Barnes-Hut Implementation in Several Parallel Programming Models – Survey of which parallel programming are a good fit for the Barnes Hut problem

SocViz: Visualization of Facebook Data – A HCI Study and development of a visualization tool for social networking data

Undergraduate Projects

Compiler Algorithm Language (CAL): An Interpreter and Compiler – Designed a new language to help compiler writers express their algorithms in a more natural way.

Picoblaze 8-bit Microprocessor – Studied this multiprocessors design and implemented and tested it in Verilog HDL.

Performance of TCP over Multiple Wireless Interface – Solved the problem of Loss Isolation in Multiple Wireless Interfaces by using TCP-in-TCP mechanism.
LaTeX Compiler – (A reduced subset of it) was developed as a part of the compiler course CS335.

Extended Nachos Project – Implemented a subset of the OS as a part of the operating systems course CS330.

Awards and Achievements

- Instructor for a class of 40 students for the "Data Structures and Algorithms" course
- Nominated among the six best B. Tech. projects (2005) in the Computer Science department
- Awarded the Student Benefit Fund Scholarship for excellent performance in academics in 2002
- Awarded the Academic Excellence Award at IIT Kanpur for the year 2001-2002
- Received a certificate of merit in the Indian National Physics Olympiad 2001
- Received a certificate of merit in the Indian National Chemistry Olympiad 2001
- Adjudged the best student and awarded the Gold Medal in 2000 in high school for excellent academic performance
- Received the SAIL Scholarship in high school in 1999 for Grade 10th performance.

References

Prof. Laxmikant V. Kale (kale@cs.uiuc.edu), Professor, Dept. Of Computer Science, UIUC

Dr. Glenn J. Martyna (martyna@us.ibm.com), Researcher, IBM T J Watson Research, NY

Dr. Sameer Kumar (sameerk@us.ibm.com), Researcher, IBM T J Watson Research, NY

Dr. Guojing Cong (gcong@us.ibm.com), Researcher, IBM T J Watson Research, NY

Prof. Sanjeev K. Aggarwal (ska@iitk.ac.in), Professor, Dept. Of Computer Science, IITK