ParTejas: A Parallel Simulator for Multicore Processors

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In this paper, we present the design of a novel parallel architecture simulator called ParTejas. ParTejas is a timing simulation engine that gets its execution traces from instrumented binaries using a fast shared memory based mechanism. Subsequently, the waiting threads simulate the execution of multiple pipelines and an elaborate memory system with support for multi-level coherent caches. ParTejas is written in Java, and primarily derives its speedups from the use of novel data structures. In specific, it uses lock-free slot schedulers to design an entity called a parallel port that effectively models the contention at shared resources in the CPU and memory system. Parallel ports remove the need for fine grained synchronization and allow each thread to use its local clock. Unlike conventional simulators that use barriers for synchronization at epoch boundaries, we use a sophisticated type of barrier, known as a phaser. A phaser allows threads to perform additional work without waiting for other threads to arrive at the barrier. Additionally, we use a host of Java specific optimizations, and use profiling to effectively schedule the threads. With all our optimizations we demonstrate a speedup of 11.8X for a multi-issue in-order pipeline and 10.9X for an out-of-order pipeline with 64 threads, for a suite of 7 Splash2 and Parsec benchmarks. The simulation error is limited to 2-4% as compared to strictly sequential simulation.

CCS Concepts: • Computing methodologies → Discrete-event simulation; Distributed simulation; Simulation tools; Systems theory;

Additional Key Words and Phrases: Parallel simulation, Architectural simulator, Tejas, ParTejas, Phasers, Parallel ports, Slot scheduling

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1 INTRODUCTION

An architectural simulator is arguably the most important tool that a computer architect uses to design, and evaluate novel processor architectures. Simulators such as SimpleScalar [4], Sesc [23], M5 [6], GEMS [17], and Multi2Sim [26], are widely used by computer architects in both industry and academia. Along with their use in industrial settings, they are also heavily used by instructors to teach courses on computer architecture. Consequently, it is necessary to design efficient architectural simulators that capture the needs of the computer architecture community.

As compared to detailed RTL level simulators, architectural simulators use a fairly high level description of a processor, and are consequently at least two orders of magnitude faster. The price

1This paper is an extension of a poster presented at ISPASS 2014 with the same title [16]
of this additional speed is accuracy. Architectural simulators are primarily used to estimate broad high level trends, and are typically not used for getting extremely fine grained performance data. Secondly, as compared to other simulation technologies such as FPGA emulators, architectural simulators are much easier to design, and it is possible to create prototypes of futuristic architectures very quickly.

Architectural simulators were traditionally sequential in nature. This is because, they were up till now being used to simulate multicore processors with at most 4-16 cores. The requirements of simulators are changing however. There are predictions that in the near future we shall have hundreds and thousands of cores per chip. In line with this trend, researchers have begun proposing architectural techniques aimed at thousand core systems [18, 24]. With sequential simulators [4, 12, 15, 22], such research is highly cumbersome. Consequently, there is a consensus view that parallel simulators are required. In the world of RTL-level simulators, parallel simulators are already commercially available, e.g., the Synopsys VCS Cheetah.

In the last 4 years, several parallel architectural simulators have been proposed for modern processors such as Graphite [18], ZSim [24], Sniper [7] and Slacksim [8]. These C/C++ based parallel simulators have mostly been written from scratch, and are typically 10x faster than their sequential counterparts. The total error as compared to sequential simulation is typically limited to 10%. Since architectural simulation is a method for estimating the broad high level trends, the research community has by and large found the amount of additional simulation error in parallel simulators acceptable.

In this paper, we present the design of a soon to be released, open source, simulator called ParTejas. ParTejas is the parallelized version of the popular Tejas simulator [25]. Tejas has been released under an open source Apache 2 license. It is a fast, trace driven, cycle accurate simulator, which is platform independent. It supports a plethora of architectural features. It simulates out-of-order pipelines, non uniform caches (NUCA), complex networks on chip (NOCs) and CUDA based GPGPUs. It also simulates relaxed memory models – specifically, processor consistency. It supports applications written for relaxed memory models by supporting the fence instruction, which requires write acknowledgements to the pipeline. Tejas has been validated against real hardware (Dell PowerEdge R620 server). It has been observed that Tejas is more accurate than most of the existing simulators (for which the validation results have been published). Tejas incurs an average absolute error of 11.45% for sequential workloads (SPEC CPU2006 benchmark suite) and 18.77% for parallel workloads (SPLASH2 benchmark suite). The reader is encouraged to read [25] for a detailed treatment of Tejas’ design, implementation and evaluation.

The important differentiating factor of ParTejas is that unlike recent work that achieve their performance by relying on high level models (of the cores and memory system), or advanced sampling based approaches, we primarily rely on novel concurrent data structures. Specifically, we use the recently proposed lock free slot scheduling technology [1], and flexible barriers (newly introduced in Java 6) known as phasers to derive our speedups. We demonstrate that by using these novel data structures, we can simulate tightly coupled multiprocessors. We view approaches such as sampled simulation and statistical simulations as complementary approaches that can always be added to our simulation setup.

Specifically, our contributions in this paper are as follows:

(1) We observe that the pipelines operate in parallel. The rendezvous points between the threads are in the memory system. The crux of our technique for implementing a parallel memory system, is a novel lock-free data structure that we term as a parallel port. The parallel port helps us use local clocks in each Java thread, yet model the contention at shared resources accurately.
(2) We propose a simulation algorithm that uses *phasers* (see Section 4.1) to dynamically reduce the time that threads spend in synchronization. This ensures that the time wasted in waiting for a barrier is significantly reduced.

(3) We design a fast shared memory based transfer mechanism between the native PIN application threads, and our Java based simulator threads.

(4) For improving the performance of Java programs, we propose a host of Java specific optimizations, namely selection of appropriate data structures, fine grained locking mechanisms, and pooling techniques.

(5) We show 10-15X speedups on in-order and out-of-order pipelines with 64 threads.

1.1 Improvements over Base Tejas

In the base *Tejas* simulator, the number of simulation threads, $n_{sim}$, is equal to 1. The efforts taken (i) to enable $n_{sim}$ to be greater than 1, (ii) to ensure that the induced error is modest and (iii) to ensure that the parallelization is maximized constitute *ParTejas*. These include spawning and scheduling of multiple simulator threads, updating the transfer engine to support $n_{sim}$ consumers, equally dividing the simulation responsibilities among the simulator’s threads, and accurately modeling the causality and contention in the memory system. These are done in a manner such that a minimal amount of synchronization overhead is introduced.

Additionally, *ParTejas* adds support for the simulation of various artefacts of parallel benchmarks. Specifically, *ParTejas* simulates the following synchronization functions of the *pthread* library: wait, signal, broadcast, mutex lock, mutex unlock and barrier wait. Importantly, causality is maintained – the order in which the different application threads invoke the different synchronization functions are exactly mimicked by their simulated counterparts. This greatly improves the accuracy of simulation. *ParTejas* also supports different atomic instructions such as *compare-and-swap* (CAS). Such support requires modifications to the coherence protocol to ensure that the core performing the CAS holds the cache line exclusively until the operation is completed to ensure atomicity.

2 SYSTEM ARCHITECTURE

2.1 Overview

Figure 1 shows an overview of *ParTejas*. We instrument regular x86 or x86-64 binaries using an instrumentation engine such as Intel’s PIN [14] to provide us with execution traces. An execution trace consists of a stream of packets, where each packet contains the details of an executed instruction such as its program counter, memory addresses accessed, and its branch outcome (if any).

This stream of packets is passed to waiting Java threads. Each Java thread simulates a set of cores, each having a separate pipeline (in-order or out-of-order). A Java thread first translates the CISC instruction packet into a sequence of VISA (a custom RISC instruction set) instructions. It then sends the VISA instructions to the appropriate pipeline. The timing of the instructions is simulated by the individual pipelines and memory accesses are passed to the parallel memory system. The memory system supports private caches, shared caches and coherence mechanisms.

This decoupling of the instrumentation and the simulation engines has important advantages. First, since the simulation engine does not have to emulate the benchmark’s functionality, it does not have to deal with register or memory values at all. The duty of the simulation engine now is to simply provide an accurate estimate of the time taken to execute the benchmark on the target architecture. It merely has to simulate accurately the times at which data is available for consumption, rather than the value of the data itself. Second, instrumentation engines such as PIN and QEMU are highly optimized, and far outperform any custom emulator. Custom emulators also...
suffer from low coverage of the intended functionality, given the complexity of the x86 instruction set. Third, the decoupling allows the two engines to run on different machines, enabling faster simulations. Fourth, the instrumentation engine may be run once and the traces may be stored in files. These files can then be used to feed the simulation engine independently. Such an approach ensures a deterministic input trace across multiple simulation runs, something that is often useful when working with parallel benchmarks. In this paper, we do not assume the file interface in our discussions, for the sake of simplicity. We assume that the emulator runs in parallel with the simulator, the former passing traces to the latter.

The Java threads do not operate in lock step. This creates issues in the memory system, which needs to cater to requests from multiple threads. We shall design novel solutions to model the causality (load-store order), and contention in the memory system.

### 2.2 PIN Instrumentation

PIN is a binary instrumentation engine that runs on Intel based x86 (32/64 bit systems). It dynamically reads the contents of a binary, and adds its instrumentation code to track events such as the execution of a memory or branch instruction. After it detects an event such as a branch event, PIN invokes a custom function. We use such custom functions to summarize the execution of an instruction in an instruction packet, and send it to the Java threads. Every instruction packet contains the application’s thread id, and the program counter of the instruction. Additionally, in the case of a memory instruction, the instruction packet contains the memory address, and in the case of a branch instruction, it contains the branch outcome. We send synchronization packets...
to indicate the beginning and end of synchronization events such as locks, unlocks, signal-wait events, and barriers. Such packets also contain the address of the synchronization variable, timestamp (real time), and nature of the event. ParTejas uses this information to simulate the effect of synchronization calls.

ParTejas can also employ QEMU as the emulator, as an alternative to PIN. This allows ParTejas to simulate the entire software stack, and not just the application. ParTejas also supports the saving of emulator trace information in files, and reading from these to perform simulation. This helps achieve deterministic outcomes in experiments involving parallel benchmarks. In this paper however, we will assume the PIN emulator mode only. The proposed optimization techniques are independent of the emulator used.

2.3 Transfer Engine

Application threads run in parallel, and also dump their traces to small in-memory circular buffers. Gigabytes of trace data are generated per second. It is necessary to send all of this data to ParTejas using a high throughput channel. We evaluated several options—shared memory, memory mapped files, network sockets, and Unix pipes. Unix pipes were found to be the slowest of all. Hence, we ceased to consider them early in our design process.

![Comparison of various inter-process communication mechanisms](image)

**Fig. 2.** Comparison of various inter-process communication mechanisms

As an experiment, we wrote a small compute intensive loop based benchmark. For this experiment, we measured the time that it takes to transfer 1 GB of data from PIN to Java on an Intel Core i7 desktop (2.4 GHz processor, 4 GB RAM, Ubuntu Linux 12.04). Since we have finite buffers, we need a method for the consumer (Java thread) to indicate to the producer (PIN) that it cannot accept more packets for the time being. In the case of sockets, we use a dedicated socket from the consumer to the producer, and for shared memory or memory mapped files, we use shared variables to indicate the status of the consumer. Figure 2 shows the results. We observe that sockets are the slowest (10 MBps). This is because they make costly system calls to transfer data across the processes, and buffering is done by the kernel. However, they are also very versatile. We can run the application threads on one machine, and run simulator threads on another remote machine.
For high throughput, shared memory is the best option (24 MBps). Communication with memory
mapped files is slower, because we need to synchronize data with the hard disk, or the disk cache
in main memory.

2.3.1 Shared Memory. We use the `shmget` and `shmat` calls in Linux to get and attach shared
memory segments to the PIN processes. We use a single shared memory segment, and avoid using a
separate shared memory segment for each application thread because underlying operating systems
typically place a limit on the number of shared memory segments a process can use. Subsequently,
we divide the segment into $n$ contiguous regions, one for each of the $n$ application threads. Each
region contains a header, and a circular queue. We fix the size of each packet to 192 bytes, and
allocate space for 50 packets in each circular queue. The header contains the status of the thread,
the number of outstanding packets in the queue ($\text{count}$), and its start and end locations.

![Structure of the shared memory segment](image)

The important point to note here is that the custom functions in PIN are written in C++, and the
simulator threads are written in Java. There is no support for Linux shared memory segments in
Java. Hence, we use the Java Native Interface (JNI) to access shared memory. It allows us to write
code in C that can be linked to the JVM at runtime. The second issue is that of locking. We need to
be able to get locks to update the pointers to the circular queue, and the $\text{count}$ variable. We use a
Peterson lock [20] optimized for the Intel x86 TSO (total store order) memory model.

The code shown in Algorithm 1 implements a Peterson Lock for TSO systems with a single fence
(hardware and compiler). Let us elaborate. A Peterson Lock without fences guarantees mutual
exclusion between two processes only for a sequentially consistent system. However, if sequential
consistency cannot be ensured then the algorithm will fail to work because the ordering of memory
accesses cannot be guaranteed. Now, in the TSO model writes to different addresses are globally
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ALGORITHM 1: The Peterson lock for x86 Systems

\[
\begin{align*}
flag[tid] & \leftarrow true \\
turn & \leftarrow tid \\
fence() \\
while flag[1 - tid] = true \text{ and } turn = tid \text{ do} \\
end
\end{align*}
\]

/* Critical Section Begins */
\[
\ldots
\]
/* Critical Section Ends */
flag[tid] \leftarrow false

ordered. As a result, the writes to \( flag \) and \( turn \) will be globally ordered, and we do not need to insert a fence between them. However, TSO relaxes the write-to-read ordering and allows a processor to read its own write early. Hence, it is necessary to have a global ordering between a write and a subsequent read. Therefore, we insert a fence between the update to \( turn \) and the while loop. Subsequently, we are only reading the values of \( flag[1 - tid] \) and \( turn \). Since TSO does not change the read-to-read ordering, there will be no violations in correctness. Similarly, while releasing a lock, there is no problem in not having a fence. We can argue that in the worst case, some other process may see that the process that held a lock has proceeded forward without changing \( flag[tid] \) to \( false \). However, this will not happen if we assume that variables accessed inside the critical section of the Peterson Lock are not accessed outside it (property of properly-labeled programs [11]). For another process to know this fact, it needs to enter the critical section first, and this is not possible unless \( flag[tid] \) is set to \( false \). Hence, our code for the Peterson Lock is correct.

Along with updating the header and its contents, we also read the execution packets that have already been transferred to ParTejas using dedicated JNI routines. JNI uses Intel x86 string instructions to transfer large amounts of data between memory locations in one go. Hence, we homogenized the structure of a packet, at the cost of space. Each packet contains three 64 byte fields. We use a single JNI call, \(\text{memcpy}(\text{void*}, \text{const void*}, \text{size_t})\), to copy an entire memory area into the address space of Java threads. Secondly, note that Java is big-endian while Intel x86 is little-endian. JNI handles this by providing its own datatypes such as \( jint \), and \( jlong \). While transferring values to the JVM, JNI routines seamlessly convert little endian values to big endian values.

2.3.2 Comparison of the Peterson Lock with other Atomic Instruction based Approaches. To figure out what is the best synchronization mechanism, we conducted a simple experiment. We created a 100 element shared queue of integers and shared it between two processes (1 written in C++, and 1 in Java). This mimics our PIN-Tejas setup. On an Intel Xeon 2.2 GHz CPU with GCC 4.7.2, we compiled our C++ code, and we compiled our Java code with Sun Java 7. We then transferred 1 million integers between the processes. The time required for transfer is shown in Table 1 using different mechanisms: Peterson Lock optimized for the TSO memory model, and different implementations using atomic primitives.

We observe the Peterson lock to be 100X faster for this workload. This is per-se not a new observation for a system with two threads. The reasons can easily be understood by looking at the data produced by David et al. [9] (SOSP 2013). An atomic operation takes somewhere between 120–310 cycles on a dual socket Xeon server. This is because it involves locking a line in the modified state, and at least a single fence instruction. A fence imposes a total order on memory accesses, and significantly increases overall latency, because loads can be issued only after the fence is committed. Most of the advantages of out-of-order processors will be lost. In comparison, only loads and stores are used in the Peterson’s lock, and thus are cheap. Most of them hit in the local cache, and are thus
very fast (1-3 cycles). In the waiting phase of a Peterson’s lock (while loop), there are no fences, and thus it is much faster. There is only one fence before the while loop, and this does not determine the overall timing.

### 2.4 Translation Engine and Pipelines

We initialize the simulation by populating a static table called the *instruction table* by reading the contents of the executable that we wish to simulate. We use the GNU `objdump` command for this. The instruction table contains the contents of each instruction indexed by the program counter in a hashtable. To avoid issues with dynamically linked libraries, we statically compile our benchmarks. If this is not possible, we can use one of several options: use trace driven simulation (file interface), or use programs such as statifier and Ermine to create statically linked binaries out of dynamically linked binaries.

During the simulation, we need to only transfer the details of memory and branch instructions from PIN. We have to transfer memory addresses because we need to simulate the memory system. For branch instructions, we transfer the branch outcome from PIN. The outcome of a branch instruction indicates the beginning of a basic block. Other than branch, memory and synchronization instructions, it is not necessary to transfer any other type of instruction such as ALU instructions from the basic block, because we can infer its execution from the instruction table. We do not need operand values because we only simulate the timing of instructions. After this offline component, we start the application emulation with PIN.

Now, for \( n_{app} \) application threads, and \( n_{sim} \) Java threads, each Java thread simulates \( n_{app}/n_{sim} \) parallel pipelines (see Figure 1). In ParTejas, each Java thread independently reads the circular queues of all the application threads that it simulates. We assume that each Java thread is physically mapped to a separate core. Subsequently, we translate each CISC instruction into a set of RISC instructions. Our instruction translation scheme is similar to PTLSim [29]. PTLSim has been validated with commercial x86 processors such as AMD K8. Next, instructions are passed to a data structure that simulates the pipeline. All the pipelines in ParTejas inherit from a generic Pipeline interface. Individual pipelines can be either in-order, or out-of-order.

For in-order pipelines, we support both single and multiple issue pipelines. For out-of-order pipelines, we model a 12 stage pipeline with load latency speculation, and multi-level bypass. Since modeling pipelines is established technology, we shall avoid explaining the operation of our pipelines in detail. In our simulator we distinguish between two types of events: regular, and non-deterministic. Regular events need some processing in the current cycle, and also need some processing in the subsequent cycle. Examples of such events are events generated by instructions moving from the fetch to the decode stage. Unless there is a pipeline stall, these instructions will progress from one pipeline stage to the next, every subsequent cycle. In comparison, non-deterministic events such as memory accesses do not have a fixed latency. It is possible that a memory event leaving the load-store queue might return after 2 cycles (L1 hit) or after hundreds
of cycles if it goes to the main memory. For non-deterministic events we use a per thread event queue, whereas for regular events, the standard iterative simulation model is employed.

3 PARALLEL SIMULATION

We should note that the process of transferring packets, translation, and execution, is inherently parallel. The challenges to parallelism arise because of interactions between threads in the form of synchronization operations, and memory operations. Traditional parallel discrete event simulators follow one of the two alternatives – a pessimistic strategy where we explicitly enforce ordering among all inter-thread interactions, or an optimistic strategy that detects causality violations and later recovers from them. When reasoning about ParTejas along these lines, we find that it adopts a mostly pessimistic strategy.

We assume that every Java thread has its local clock. The threads periodically synchronize using a barrier at every epoch boundary (clocks become the same). This periodic synchronization of clocks ensures that most interactions between threads are simulated in the same order as emulated, or at the very least reduces the timing skew. An epoch is defined as a given number of cycles measured using the thread’s local clock. We discuss the tradeoffs for choosing an optimal epoch size in Section 6.5.

We now discuss further efforts at enforcing synchronization so as to minimize the simulation error, while still maintaining the performance benefits of parallelization.

3.1 Synchronization Operations

The POSIX pthread library primarily uses three kinds of synchronization operations: signal – wait, lock – unlock, and broadcast – wait. The lock-unlock primitive is used to implement critical sections by establishing exclusive ownership of a lock variable. The signal-wait and broadcast-wait primitives are used to make a set of threads wait till another thread releases them. Barriers internally use the broadcast-wait primitive. Let us consider signal-wait. During simulation, it is possible that a waiting thread might move ahead of the signaling thread. This is because it might be the case that PIN was able to quickly send the packets for the waiting thread, and the packets for the signaling thread got delayed, possibly due to a context switch. Thus ParTejas will see the waiting thread resume before seeing the signal event.

We have two modes. In the first mode, nosync, we allow this situation to happen in the interest of simulation speed. We shall see that the jitter introduced by PIN is not very large. However, we have a more accurate (but pessimistic, in terms of its effect on performance) mode, fullsync, which tracks dependences across threads, and enforces them. We track the following events le (lock enter), lx (lock exit), ue (unlock enter), ux (unlock exit), se (signal enter), sx (signal exit), be (broadcast enter), bx (broadcast exit), we (wait enter), and wx (wait exit) using the emulator. For each event, the emulator appends its timestamp (real time), and address in a synchronization packet, and sends it through the transfer engine. The simulator process has a table called syncops that maintains a list of all the synchronization operations indexed by the address of the synchronization variable. syncops is protected by a single lock.

Let us first define some terminology. we(tA) refers to the wait enter event of thread A. We further say that we(tA) < we(tB), if the timestamp of the former event, as recorded by the emulator, is less than the timestamp of the latter. Based on these timestamps, we want to establish a partial order → across events in the simulation. A → B means that A needs to happen before B. Table 2
shows the timing condition, as recorded by the emulator, on a synchronization variable for different synchronization operations, and the consequent invariant that we need to maintain in the simulation.

<table>
<thead>
<tr>
<th>Condition</th>
<th>Invariant enforced in ParTejas</th>
</tr>
</thead>
<tbody>
<tr>
<td>Signal-Wait</td>
<td></td>
</tr>
<tr>
<td>$w_e(t_B) &lt; s_e(t_A) &lt; w_x(t_B)$</td>
<td>$s_e(t_A) \rightarrow w_x(t_B)$</td>
</tr>
<tr>
<td></td>
<td>$w_e(t_B) \rightarrow s_e(t_A)$</td>
</tr>
<tr>
<td>Broadcast-Wait</td>
<td></td>
</tr>
<tr>
<td>$w_e(t_B) &lt; b_e(t_A) &lt; w_x(t_B)$</td>
<td>$\forall t_B, w_e(t_B) \rightarrow b_e(t_A)$</td>
</tr>
<tr>
<td></td>
<td>$\forall t_B, b_e(t_A) \rightarrow w_x(t_B)$</td>
</tr>
<tr>
<td>Lock-Unlock</td>
<td></td>
</tr>
<tr>
<td>$(l_e(t_A) &lt; l_e(t_B) &lt; u_e(t_A))$ \land $(u_e(t_A) &lt; l_e(t_B))$</td>
<td>$u_e(t_A) \rightarrow l_x(t_B)$</td>
</tr>
<tr>
<td></td>
<td>$l_e(t_A) \rightarrow l_e(t_B) \rightarrow u_e(t_A)$</td>
</tr>
</tbody>
</table>

Table 2. Conditions for maintaining the synchronization order across threads

For the sake of brevity, let us explain the most complex lock-unlock condition (the others follow the same pattern). The first subcondition says that if a lock-enter for thread B arrives when the same lock is held by thread A, B needs to wait till the lock is released. We thus have: $u_e(t_A) \rightarrow l_x(t_B)$. Secondly, we need to ensure that the lock-enter call for B is processed by ParTejas during the duration that A holds the lock. Hence, we have, $l_e(t_A) \rightarrow l_e(t_B) \rightarrow u_e(t_A)$. Note that the right column is the invariant we need to enforce by stalling the pipeline of either A or B, and the left column is the set of conditions that the timestamps of the events must satisfy. To implement any dependency of the type $A \rightarrow B$, whenever we receive B, we stall all the threads till they have received events with greater timestamps. This ensures that A has reached one of the Java threads. B needs to retire from its pipeline after A has retired from its pipeline. After B retires, we can free the entry for A in the syncops table.

### 3.2 The Memory System

#### 3.2.1 Parallel Ports

Implementing the memory system (both shared and coherent caches) is difficult because there are happens-before relationships in memory ($\text{store} \rightarrow \text{load}$), and it is necessary to model the contention in memory banks and directory structures. The interactions between the threads in the memory system can be classified into three categories: (1) contending for access to the shared resource (cache, directory, network); (2) memory behavior when they access the same shared data; (3) memory behavior when they access private blocks that happen to map to the same set in the shared cache. In the third case, changing the order of accesses may change the eviction pattern in the shared cache and the directory, leading to results different from the native run. Fortunately, prior work [24], and our experiments (Section 6.5) demonstrate that the number of such conflicting accesses between threads in a small window of time is very small. The second type of interaction, that is, accessing the same shared data is much more frequent. Since synchronization primitives are modeled accurately in ParTejas (see Section 3.1), and most programs are free of data races (accesses to shared data are wrapped in critical sections), there is no possibility for any deviation due to shared data access.

We thus primarily focus in this section on the first type of interaction – modeling contention. The most important assumption that we make is that every thread issues requests using its local...
The memory system does not have a separate clock. For example, if a thread with its local clock equal to $\tau$ records a miss in the L1 cache, then it accesses the shared L2 cache at time $\tau$. At this point, it essentially assumes that its local clock is equal to the global clock. It performs the L2 access using its local clock. As discussed, we need to model contention at the L2 cache. For example, if $\tau = 10$, we need to consider the state of the L2 cache at the 10th clock cycle, and delay the current request if there is contention at that cycle. Unlike other simulators, there is no separate thread that simulates the memory system.

To model contention, we assume that every structure (cache bank, controller, directory, NOC element) has a parallel port at its inputs, which acts as a gate keeper. The parallel port is implemented as a matrix of slots, where the number of rows is equal to the number of simultaneous requests that can be processed in the same cycle (capacity), and each column represents the number of the cycle (as per the local clock of a thread) in the current epoch. Each slot can either be available or busy (if a request is scheduled). For example, if an L2 request requires 3 clock cycles, it needs to reserve 3 contiguous slots.

A parallel port that can accept only one request per cycle is shown in Figure 4. Let us assume that two requests arrive simultaneously. The request is in the form of a tuple having the requested starting slot(cycle) number, and the number of consecutive slots as its two parameters. Since slots 3 and 5 are busy, we need to start scheduling from slot 6. The parallel port arbitrarily chooses request 2 to be scheduled in slots 6 and 7. Henceforth, it schedules request 1 in slots 8, 9, and 10. The main aim of a parallel port is to provide a linearizable (all requests appear to execute instantaneously) data structure that tries to schedule requests as close as possible to the requested time. It can effectively model contention because it never schedules more requests in a cycle than the capacity of the structure.

We implement the parallel port using lock free algorithms based on the work on non-blocking slot schedulers by Aggarwal and Sarangi [1]. They propose a method to implement large 10,000 to 100,000 entry parallel ports using algorithms that do not use any kind of locks. Each cell is an
atomic integer, and we need to only use the compare-and-set primitive. This approach is at least two orders of magnitude faster than approaches that use locks. The main idea of their approach is to first temporarily reserve the earliest available slots. If there is a conflict with another thread while performing temporary reservations, then one thread helps the other thread in completing its request. After a thread completes its request either on its own, or with the help of other threads, it proceeds to permanently mark them as reserved. This data structure is linearizable, i.e., operations appear to take effect instantaneously at some point of time between its invocation, and completion. Our implementation of the parallel port can typically process a request in less than 2-3 $\mu$s.

Since lock free algorithms can sometimes suffer from starvation, we have the option of using a wait free parallel port that guarantees completion of the request in a finite amount of time. This is again based on the work by Aggarwal and Sarangi [1]. A thread desirous of booking some slots first places its request in an array of requests (one entry per thread). Subsequently, it traverses the older entries of the list to see if there is any request, which has been waiting for a long time (determined by a threshold). If there is such a request, then the current request first helps this request. This is done by taking the parameters of the request (starting slot, number of slots) and then trying to mark the slots in the parallel port. This is again a two step process. First, the thread needs to temporarily reserve the slots, and then tries to permanently reserve them. At any point of time in the temporary reservation process, it is possible that another thread might take a slot away. In this case, the thread needs to undo its work, and start from another position. It is also possible that another helper completes the work. This also needs to be checked periodically. This approach is much slower than the lock-free algorithm, yet it guarantees that every thread makes progress and there is no starvation. The wait free implementation was measured to be 3-5X slower than the lock free implementation, and is only suitable for a system with a very large number of simulation threads, or a system that has a lot of jitter. The rest of this paper is based on the lock free parallel port.

Another subtle point to note is that as we reach the end of an epoch, we need to place requests in the next epoch. We thus use a pair of parallel ports. When, we reach the end of the epoch, we place requests in the other parallel port. Thus, for a short duration of time both the parallel ports are used, and finally when all the threads reach the end of the epoch, the old port is not used anymore. The old port will be reused at the end of the next epoch. To avoid cleaning up entries eagerly, and thus wasting time, we lazily clear them by prefixing the least significant bit of the epoch number with the state bit (vacant or reserved).

3.2.2 Directory Coherence. The moment, there is a read/write miss in a private cache, the cache controller of the requesting core, $C$, sends a message timestamped with its local clock. After passing through the parallel port, the Java thread looks up the list of sharers. The directory implements a MOESI protocol, and subsequently sends a request to the owner core. The owner core uses the clock domain (local clock) of $C$ to access its private cache, and sends the block directly to the cache controller of $C$. The reader should note that all these operations happen in the clock domain of the requesting core, $C$. All the relevant parallel ports are accessed using this clock domain. Secondly, the Java thread simulating $C$ performs all of these accesses. A memory request never passes to another thread.

We admit that this approach introduces some error, as the state of the cache being accessed corresponds to a time different from that of the requesting cache. However, the error introduced is low, as our results confirm (see Section 6.5). The low error is due to two reasons: (i) accesses to shared data, which are more critical when determining performance, are simulated accurately as synchronization operations are captured meticulously (see Section 3.1), and (ii) accesses to shared structures like the directory are modeled using the parallel port, thereby capturing the contention.
among accesses (see Section 3.2.1). Our aim in following this coherence approach is to provide an abstraction that a simulator thread operates independently of others. At a small price in terms of error, significant gains are attained in simulator performance by not forcing a synchronization between simulator threads for every directory access.

4 OPTIMIZATIONS

Aside from the high performance parallel ports, we also propose the following optimizations.

4.1 Relaxed Synchronization using Phasers

We shall observe in Section 6.3 that the periodic synchronization at the end of an epoch is one of the largest limits to scalability. The reason for the lack of scalability is that a considerable imbalance exists between the slowest thread and some of the fastest threads. The faster threads wait for the slowest thread to catch up at the barrier, resulting in a considerable amount of idling in the system. For example, it is possible that one thread simulates cores that have high IPC, and thus it needs to do a lot of work to reach the end of the epoch. In comparison, another thread might simulate cores that suffer a series of L2 cache misses. Most of the time, its pipelines will be idle, and thus the simulation will proceed quickly. When both of the threads synchronize at a barrier, one of the threads has to remain idle, and wait for the other thread to finish its epoch. We have observed in our experiments that most of the time the interaction between threads is relatively low, and thus it does not make sense for threads to wait for each other all the time.

Consequently, we propose to replace barriers with phasers to cater to programs which have long phases of low interaction between threads. A phaser is a special kind of barrier that has two distinct points. After a thread arrives at the first point, it informs the rest of the threads. No thread can cross the second point, unless all the threads have reached the first point. Note that a regular barrier is a special kind of phaser that merges both the points. Java 6 (with backports) or Java 7 has native support for phasers. We use phasers to consider two epochs at a time. After the end of the first epoch, a thread enters the first point of the phaser. Instead of suspending itself, the thread continues till the end of the second epoch or till the time it has to access a shared resource such as the shared cache or the communication interface. At this point it enters the second point of the phaser. Now, if there is any thread that has not completed the first epoch, then the thread waits for it to finish the first epoch. Thus phasers give us the same synchronization benefit as barriers and at the same time we are able to reduce the idle time by a considerable amount (see Section 6.3). The reason we can reduce idling with a phaser is because it is better suited for adjusting to timing jitter in simulation. Since we allow a limited amount of divergence between the threads, the jitter can get balanced across two epochs. On the other hand, a phaser does not allow the divergence between the cycle count of two threads to be arbitrarily large. Any two threads can at the most be one epoch away. Thus there is a tradeoff between accuracy and the amount of idle time here.

The addition of phasers requires us to modify the structure of our parallel ports. Previously, the size of a parallel port was equal to the size of an epoch, or phase, and we used two parallel ports to seamlessly place requests at the epoch boundary. Now, we maintain three parallel ports (one for each phase, and one buffer). To reuse our ports, we again lazily clear them by appropriately encoding our states. We can extend this idea, to consider phases with \( n \) points, and use \( n + 1 \) copies of parallel ports for each structure. There is a space, speed, and accuracy tradeoff here. We can switch between barriers and phasers depending on the amount of inter-thread communication. If the load at all the parallel ports is higher than a threshold, then we may switch to using regular barriers at epoch boundaries to ensure additional accuracy.
4.2 Core Partitioning and Mapping

We have $n_{app}$ application threads. In comparison we have $n_{sim}$ Java simulator threads, where each thread typically simulates more than one application thread. We need to carefully choose the number of cores that we assign to the application, and the number of cores that we assign to the Java threads. We observed in our experiments that statically partitioning the cores among these two class of threads was better in terms of performance than allowing the OS scheduler to manage application and Java threads.

To maximize performance, we need to carefully compute the right sizes of the two partitions. For our workloads, we did not observe a lot of intra-application phases. Hence, we divide the execution into two phases. The first is a short profiling run that runs a benchmark for a pre-determined number of instructions with varying partition sizes. We choose the best result for the partition, and use the `sched_affinity()` call in Linux to create the partitions.

4.3 Java Specific Optimizations

A key principle in our design is to avoid locks as much as possible. Most of our structures are non-blocking. We use Java’s built in concurrent data structures that are known to have lock-free implementations whenever possible. We observed that the appropriate choice of the data structure is important.

For example, some read only structures such as the instruction table should be implemented using HashMaps rather than HashTables. HashTables are synchronized (use locks) by default, whereas HashMaps are not synchronized (do not use locks). Similarly, Vectors are synchronized by default, whereas the ArrayList does not use locks. Some data structures such as the cache bank’s tag array has both read and write accesses. The probability of a simultaneous access to the same entry is very low. Hence, in the interest of speed, we implement it with a HashMap. The chances of an occasional error is very small. Similarly, thread specific state in shared data structures such as the directory are implemented using Array Lists.

To avoid false sharing, we try to minimize the number of instances in which we place data belonging to different threads in the same cache line, even if the threads access disjoint sets of data. Instead, we use `ThreadLocal` variables to localize the data of each thread.

Lastly, there is a tradeoff between manual memory allocation, and garbage collection. Garbage collection is fairly efficient, and boosts programmer productivity for data structures that are not instantiated very frequently. However, classes such as events and instructions are instantiated extremely frequently. For events, we follow a policy of object reuse. The event class is so engineered that it can be used for a variety of communication scenarios between different architectural elements. Therefore, instead of instantiating new events every time, existing objects are reused. Instructions, in `ParTejas`, have elegant points of entry (fetch stage) and exit (commit stage). So for them, we adopt object pooling. A pool of instruction objects are created. Objects are drawn from the pool when required during the fetch stage, and returned to the pool at commit. Per thread pools are maintained to avoid expensive synchronization.

5 RELATED WORK

For the sake of brevity, we shall only discuss parallel architecture simulators in this section. For a background of sequential simulators, the reader can look at the following simulators: Simics [15], SimFlex [27], GEMS [17], COTSon [3], SESC [23], ESESC [2], Multi2Sim [26], M5 [6], and SimpleScalar [4].

Wisconsin Wind Tunnel [21] was one of the earliest parallel simulators. It was designed to run on CM-5 machines, and used barriers for synchronization similar to `ParTejas`. Wisconsin Wind
Tunnel II [19], was later proposed as an extension to Wisconsin Wind Tunnel. This proposal relied on extensive binary instrumentation to estimate the execution time of basic blocks rather than instructions. Secondly, for simulating memory, the authors proposed to replace memory instructions by small code snippets that quickly estimate the time an instruction will take to execute. The main limitation of these two projects was that they could not be used to simulate complex architectures, complex cache coherence, and were not very customizable. On similar lines, BigSim [30] and FastMP [13] were designed to simulate large message passing computers such as the IBM Blue Gene. They primarily simulate the message complexity across processors. In comparison we simulate complex out-of-order pipelines and elaborate cache coherence protocols as well.

SlackSim [8] uses the pthreads API to parallelize the Simplescalar simulator. Each core of the target CMP is simulated by one pthread and the L2 cache is simulated by another dedicated manager thread. Unlike ParTejas, a single thread (the manager thread) can become a bottleneck (also pointed out by the authors of the paper). Graphite [18] uses PIN based instrumentation, and runs a single binary on a cluster of machines. A major focus of Graphite is to provide the abstraction of a single address space for all the threads running on different machines. This is achieved by trapping each memory access, and converting it into a message that is sent over the network to another node that contains the value of the memory address. Along with functionally simulating a single binary on multiple machines, Graphite simulates cores, and the memory system at each node. The authors use Graphite to simulate private caches, and private DRAM controllers using Graphite. In our opinion, it is difficult to use Graphite to simulate a CMP with a tightly coupled cache coherence system, complex NOC, and a multitude of shared structures, because each message needs to be sent over the network. ParTejas on the other hand is designed to simulate more tightly coupled systems.

Sniper [7] speeds up simulation by using high level models. The authors define an interval to be a compute bound phase of a thread delimited by misses in the data or instruction caches. Sniper uses high level models of cores to speed up the simulation time of intervals, instead of performing cycle accurate simulation. Between intervals, it simulates the misses in the memory system, and computes their latency. It simulates an approximate OOO model. Only the functionality of an OOO pipeline is simulated, whereas the structures such as the load-store queue, rename buffer and register-allocation table are not simulated. This makes Sniper unfit to study and design novel structures that are part of a detailed architectural pipeline. The authors also suggest that interval simulation is not a substitute to cycle accurate simulation [10]. It is rather a complementary approach, which can be used when the need for accuracy is not paramount and the focus is on the speed of simulation.

Similarly, ZSim [24] uses high level models for in-order and out-of-order cores. It quickly estimates the execution time for a sequence of instructions. It introduces a novel method called “bound and weave”. The bound phase runs the instructions assuming zero cycle latency for memory instructions, and it simultaneously collects traces. The weave phase executes the traces on dedicated nodes that simulate the memory system. The final latency is obtained by a combination of the results in the bound and weave phases. Sampling, modeling, and other approximate simulation approaches, are orthogonal to our approach. It is possible to seamlessly augment our simulator to support a variety of high level models, and sampling approaches.

6 RESULTS
In this section, we shall discuss the performance gains achieved through our parallelization techniques. We shall also discuss the error, or rather the deviation, induced due to parallelization.
6.1 Experimental Setup

We evaluated the performance of ParTejas on a four socket, 64 bit, Dell PowerEdge R820 server. It had four 8 core 2.20GHz Intel® Xeon® cpus (with hyper-threading enabled), 16 MB L2 cache, and 64 GB of main memory. We thus had a maximum of 64 cores visible to software. This server runs Ubuntu Linux 12.10 using the generic 3.5.0-17 kernel. All our code is written in Java 6 using Sun OpenJDK 1.6.0_27 with the latest patches. We use Intel PIN (rev:49306) [14], with gcc 4.7.2. Table 3 shows the details of the simulated architecture. We show our results with multi-issue in-order (MII) and out-of-order (OOO) pipelines. We use private L1 caches (instruction and data), and a large shared L2 cache. We group 4 cores into a cluster (also referred to as tile in the literature). Each cluster has a slice of the shared L2 cache. A 64 core system, divided into 16 clusters, is simulated for all our experiments.

We simulate 7 benchmarks from the Parsec [5] (blackscholes, fluidanimate, swaptions, canneal, streamcluster) and Splash2 [28] (ocean_cont, ocean_ncont) benchmark suites. We chose benchmarks that have large parallel sections, and that spawn parallel threads early in their execution (for quicker simulation). For Parsec we use simsmall inputs (trends were the same across all the input sets), and for Splash2 we use the default inputs. An epoch size of 1000 cycles is used for all our simulations.

<table>
<thead>
<tr>
<th>Architectural Parameters</th>
</tr>
</thead>
<tbody>
<tr>
<td>Frequency</td>
</tr>
<tr>
<td>Cores (4 per cluster)</td>
</tr>
<tr>
<td>NOC</td>
</tr>
<tr>
<td>Issue width</td>
</tr>
<tr>
<td>Decode width</td>
</tr>
<tr>
<td>ROB size</td>
</tr>
<tr>
<td>Branch predictor</td>
</tr>
<tr>
<td>L1 size (I and D) (per core)</td>
</tr>
<tr>
<td>L2 size (per cluster)</td>
</tr>
<tr>
<td>Associativity</td>
</tr>
<tr>
<td>Latency</td>
</tr>
<tr>
<td>Cache coherence</td>
</tr>
<tr>
<td>Directory</td>
</tr>
<tr>
<td>Memory Controllers</td>
</tr>
<tr>
<td>Memory Latency</td>
</tr>
</tbody>
</table>

Table 3. Parameters of the simulated architecture

6.2 Performance Results

For all our experiments, we simulate a 64 core processor (details shown in Table 3), that is, the application has 64 threads, each mapped to a different simulated core. The number of Java threads, njava, that simulate the application’s execution on the target architecture is varied – 8, 16, 32 and 64. Each Java thread is responsible for the simulation of 64/njava application threads. As elaborated in Section 6.1, our simulation infrastructure has 64 physical cores. These physical cores are statically partitioned between the application threads and the Java simulation threads. That is, njava physical cores are allocated to the Java threads, and the remaining are allocated to the application threads. For each benchmark, we show the normalized speedup with respect to the sequential execution time on 1 core. The bar chart shows the results with phasers and the results with simple barriers are shown using the diamond shaped dots. Note that for some benchmarks, such as streamcluster,
intermittently some more threads (up till 69) are created. We assign each such additional thread (that is short lived) to a new virtual pipeline with the same configuration as a regular pipeline, and new Java threads.

Our performance results are shown in Figure 5. Let us first discuss the general trends with phasers. With the MII pipeline, blackscholes and fluidanimate show speedups of 4.2-5.1X for 8 threads, 10-15X for 16 threads. However, the scalability tapers off beyond 32 threads. In case of the OOO pipeline, the speedup is 5-5.5x for 8 threads, 7.5-9.3x for 16 threads and 9.7-11.5x for 64 threads. swaptions an incremental benefit beyond 32 threads for the in-order pipeline. canneal, ocean_cont, and ocean_ncont show a steady rise in speedup as we move from 16-64 cores. For 16 threads, their speedups are between 2.2-6.6X, for 32 threads, their speedups are between 2.3-9.43X, and for 64 threads, their speedups are in the range of 6.2-12.5X respectively for both the pipelines.

Fig. 5. Performance results (speedups with respect to 1 core)
streamcluster is an outlier. We see a speedup of 7.8-9.8X for 8 threads. The performance goes down for 16, and 32 threads in the case of the MII pipeline. With the OOO pipeline, we observe a slow down for 32 threads. The speedup increases to 10.6-12.3X for 64 cores. streamcluster creates a large number of auxiliary threads, and there is a sizeable amount of interaction between the threads. However, as we increase the number of hardware threads, it is easier to schedule the execution of these auxiliary threads. Hence, the performance improves with 64 threads.

Let us now look at the results with barriers (diamonds in the graph) for the MII pipeline. The first trend is that on average, barriers are 42.6% slower than phasers in terms of the speedup for 64 threads. The second point to note is that other than blackscholes, and streamcluster, simulations with 32 or 16 threads are faster than those that use 64 threads. The synchronization overheads dominates in this case. We can thus infer a net loss in scalability with the use of barriers in this case.

In the case of the OOO pipeline, the difference in performance between barriers and phasers is not so large. On an average phasers are 7.23% better than barriers for 64 threads. Some benchmarks such as swaptions and ocean_cont show 9.23% and 28.86% improvement respectively for 64 threads. The reasons for the diminished difference is as follows. The OOO pipeline is fairly complicated and slow as compared to inorder pipelines. Consequently, there are fewer idle cycles and rarely any outlier threads that have a lot of idle time at an epoch boundary (refer to Section 4.1).

Table 4 shows the simulation speed in MIPS (millions of instructions per second). For 64 cores, we observe simulation speeds of 3.5-8.2 MIPS with the MII pipeline and simulation speeds of 0.2-4.0 MIPS with the OOO pipeline across 7 benchmarks. Note that canneal is an exception (0.06 MIPS (16 cores), and 0.2 MIPS (64 cores)) because of the large sequential sections in the benchmark which slowed down the OOO pipeline. For 16 cores, the simulation speeds for both pipelines are of the order of 2.8 MIPS and 1.95 MIPS respectively. In comparison, Sniper [7] which uses high level models reports a simulation speed of 2.2 MIPS for simulating 16 cores, on a 24 core SMP machine.

### 6.3 Analysis of the Speedups

<table>
<thead>
<tr>
<th>Application</th>
<th>MIPS</th>
<th>Barrier</th>
<th>Par. Port (%)</th>
<th>Mem. Sys. (%)</th>
<th>Pipeline (%)</th>
<th>Phaser</th>
<th>Par. Port (%)</th>
<th>Mem. Sys. (%)</th>
<th>Pipeline (%)</th>
<th>PIN</th>
<th>Java Port</th>
<th>Synch. Error (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>blackscholes</td>
<td>3.5</td>
<td>6.99</td>
<td>1.98</td>
<td>0.91</td>
<td>21.02</td>
<td>40.56</td>
<td>35.03</td>
<td>0.01</td>
<td>24.40</td>
<td>3</td>
<td>39</td>
<td>0</td>
</tr>
<tr>
<td>fluidanimate</td>
<td>4.9</td>
<td>73.84</td>
<td>1.28</td>
<td>0.60</td>
<td>24.28</td>
<td>32.36</td>
<td>19.19</td>
<td>0.60</td>
<td>27.85</td>
<td>5</td>
<td>56</td>
<td>0.02</td>
</tr>
<tr>
<td>swaptions</td>
<td>3.1</td>
<td>83.27</td>
<td>1.23</td>
<td>1.07</td>
<td>14.43</td>
<td>39.53</td>
<td>37.92</td>
<td>1.07</td>
<td>21.48</td>
<td>15</td>
<td>39</td>
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<tr>
<td>canneal</td>
<td>1.7</td>
<td>88.52</td>
<td>4.83</td>
<td>0.10</td>
<td>6.35</td>
<td>32.98</td>
<td>47.17</td>
<td>0.62</td>
<td>19.23</td>
<td>8</td>
<td>56</td>
<td>0</td>
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<tr>
<td>streamcluster</td>
<td>4.9</td>
<td>85.00</td>
<td>1.29</td>
<td>0.01</td>
<td>13.10</td>
<td>61.43</td>
<td>19.84</td>
<td>0.42</td>
<td>18.29</td>
<td>6</td>
<td>58</td>
<td>0</td>
</tr>
<tr>
<td>ocean_cont</td>
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<td>0.94</td>
<td>0.08</td>
<td>7.61</td>
<td>35.47</td>
<td>44.63</td>
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<td>19.89</td>
<td>7</td>
<td>37</td>
<td>0.05</td>
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<tr>
<td>ocean_ncont</td>
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<td>89.17</td>
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<td>0.06</td>
<td>10.00</td>
<td>39.40</td>
<td>46.74</td>
<td>0.90</td>
<td>22.96</td>
<td>10</td>
<td>54</td>
<td>0.06</td>
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<tr>
<td>mean</td>
<td>2.8</td>
<td>84.11</td>
<td>1.76</td>
<td>0.28</td>
<td>13.85</td>
<td>41.68</td>
<td>35.78</td>
<td>0.52</td>
<td>22.02</td>
<td>9</td>
<td>55</td>
<td>0.02</td>
</tr>
</tbody>
</table>

Table 4: Simulator speed, break-up of time taken, and optimal partition sizes for 64 threads

Table 4 shows the break up of the total time in terms of the average (across all threads) time taken to wait for barriers (or phasers), time spent in the parallel ports (and NOC), the time taken to
access memory structures (caches, directory), and the time the pipeline takes to execute. Let us explain with an example. Consider a system with two threads that runs for one epoch (4s long). Let the first thread take 2s, and the second thread take 4s. Assume that there were no accesses to the memory or the parallel ports. In this case, the average time in the pipeline is 3s and the average time to wait for a barrier is 1s.

To measure these parameters we instrument the code with calls to read the current time. For example, when we start a function that simulates the pipeline we record the time at which we enter and leave the function. We have similar instrumentations for the core functions of the memory system, parallel ports, and the barrier/phaser. We use Java’s built-in routines that access the CPU’s built-in high resolution timer. An astute reader may argue that these instrumentations may skew the readings and change the behavior of the simulator. However, we took care to minimize their use, and since we have well defined access points for each module, we did not require a lot of instrumentation. At the end we collate all the data.

As we discussed in Section 6.2, the OOO pipeline shows similar trends with both phasers and barriers. A large amount of time (70-90%) is spent waiting at barriers or phasers respectively. On the other hand, phasers perform better than barriers by 29.87% with the MII pipeline. With phasers, we spend much less time waiting for a phaser to release (32-61%). This is because phasers allow threads to move forward between the first and second points. Note that streamcluster is an exception (61%) because it spawns a large number of additional threads. We start spending higher amounts of time at the parallel ports (19-47%). Since, all the shared structures (cache banks, NOC routers, directories) have parallel ports, the delay becomes large when there is a higher degree of parallelism enabled by phasers. In both cases, memory structures are accessed very quickly (because we use HashMaps).

6.4 Core Partitioning

For each benchmark, Figure 6 shows the simulation speed (in MIPS), as a function of the partition sizes with 64 cores. The x-axis shows the number of cores assigned to PIN, and the number of cores assigned to the Java threads. The format, \(m - n\), means that the PIN threads are assigned \(m\) cores, and the Java threads are assigned \(n\) cores. From Figure 6, we can infer that there is an optimal partition size, beyond which the speedup decreases. For example, for the MII pipeline, the optimal partition size for streamcluster is 6-58, and for blackscholes is 5-59. The second important point to infer from Figure 6 is that choosing an optimal partition can increase the speedup by as much as 50-100%. Table 4 shows the optimal partition sizes for each benchmark for the experiments with 64 cores.

6.5 Simulation Errors and Deviations

Let us divide the simulation error into two categories – error in enforcing proper synchronization (lock-unlock, signal-wait, broadcast-wait), and errors introduced due to the altered nature of the memory system. For the former case, let us compare the mean simulated execution time across the two configurations, and report the error (relative difference). The first configuration uses a strict synchronization order as explained in Section 3.1. The second configuration does not obey any synchronization constraints at all. The error is shown in the last column of Table 4. The error is limited to 0.23%, which is negligible. This is because PIN enforces synchronization strictly, and the additional jitter introduced in the transfer, and translation engines is minimal. Hence, it is not necessary to strictly enforce synchronization for Parsec/Splash benchmarks. The packets come to ParTejas (Java side) in the proper order.

Let us now evaluate the deviations due to the altered nature of the memory system. Let us first evaluate the nature of sharing of memory addresses between threads in an epoch. Figures 7, 8, and
Fig. 6. Speedup as a function of the sizes of partitions (64 threads)

9 show representative address diagrams for an epoch from the fluidanimate benchmark. Owing to the fact that the fluidanimate benchmarks has high level of synchronization [5], it serves as a good representative for this evaluation. In an address diagram, the application threads are arranged in concentric circles. Thread 1 is assigned the innermost circle, and thread 64 is assigned the outermost circle. The entire physical address space is split radially. Now, for each thread, we find out all the non-read-only addresses that it has accessed in an epoch, and mark them in the address diagram. For addresses that are close by and map to the same cache line, we create a bigger circle, instead of a cloud of points. Also if the same address is accessed more than once, we increase the size of the circle. Thus, in each of the address diagrams, we see a multitude of circles of various sizes. Now, we can infer memory contention if we have multiple circles along the same radial line (line passing through the center of a circle). When we use an epoch size of 500 cycles(Figure 7), the contention
is minimal. There are at most two circles on any radial line. In comparison, we have two highly contended memory regions with an epoch size of 1000 (Figure 8), and for an epoch size of 10,000, we have multiple (> 4) highly contended memory regions. There is a tradeoff between the epoch size, and the interaction between threads. We choose an epoch size of 1000 aiming to balance speed, and the level of interaction between threads.

If there is a lot of interaction between threads in the memory system, then the order of simulating the accesses is important. Changes in the order of simulating memory accesses can change the simulated execution time of the benchmark as follows. Let us assume that thread $T_1$ has two writes to the same address $W_1$ and $W_2$ interspersed in time. Similarly, thread $T_2$ has two reads to that address ($R_1$ and $R_2$). The order, $W_1 \rightarrow W_2 \rightarrow R_1 \rightarrow R_2$, will lead to lower simulated execution time because of higher locality. In comparison, $W_1 \rightarrow R_1 \rightarrow W_2 \rightarrow R_2$, will result in additional transfers between caches leading to a higher simulated execution time. Hence, it is necessary to choose the epoch size carefully. Let us take a look at the average deviation as compared to a sequential execution in Figure 10. The deviation in mean simulated execution time varies from 0.5 to 8.6%, and the deviation in the total L2 miss rate varies from 2-7.4% for both the pipelines. There is some variance (<2%) for different runs of the same benchmark (shown in error bars). The reason, we use the word deviation here, instead of error is that once we enforce synchronization, and the program is data race free, the simulator does not execute incorrectly. The order of memory accesses might differ as compared to a sequential execution, and the delays of some memory operations might appear to be unusually large because of the parallel port; however, the execution of memory accesses still follows a valid ordering. Recall that the parallel port is a linearizable (stronger than sequential consistency) structure, and tries to schedule requests as close as possible to the requested time. Nonetheless, the consensus view in the parallel simulator community is to have simulation results as close as possible to that of a sequential run. In this regard, FluidCheck achieves mean deviations of 6% and 1.3% in the simulated execution time, for the MII and the OOO pipeline cases respectively, and a mean deviation of 3% in the L2 miss rate (in both pipeline cases). This is similar to competing simulators [18, 24].

7 CONCLUSION

We outlined the design of a novel Java based parallel simulator called ParTejas that can simulate a suite of Splash2 and Parsec benchmarks, which can provide a mean speedup of 11.8X for a multi issue in-order pipeline, and 10.9x for an out-of-order pipeline with 64 threads, as compared to sequential execution. ParTejas is different from the other conventional proposals that rely on high level models, and sampling. Unlike others, we base our performance on novel concurrent data structures such as the parallel port, and phasers, along with Java specific features, and intelligent
core partitioning. As we observed from the results, 70-80% of the total simulation time is consumed in waiting at the barriers, hence we might be able to find improvements in the near future using better simulation techniques and data structures.

REFERENCES


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