Theory Comput. Systems **35**, 289–304 (2002) DOI: 10.1007/s00224-002-1055-5

Theory of Computing Systems New York Inc.

# Online Scheduling of Parallel Programs on Heterogeneous Systems with Applications to Cilk\*

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**Abstract.** We study the problem of executing parallel programs, in particular Cilk programs, on a collection of processors of different speeds. We consider a model in which each processor maintains an estimate of its own speed, where communication between processors has a cost, and where all scheduling must be online. This problem has been considered previously in the fields of asynchronous parallel computing and scheduling theory. Our model is a bridge between the assumptions in these fields. We provide a new more accurate analysis of an old scheduling algorithm called the *maximum utilization scheduler*. Based on this analysis, we generalize this scheduling policy and define the *high utilization scheduler*. We next focus on the Cilk platform and introduce a new algorithm for scheduler is inspired by the high utilization scheduler and is modified to fit in a Cilk context. A crucial aspect of our algorithm is that it keeps the original spirit of the Cilk scheduler. In fact, when our new algorithm runs on homogeneous processors, it exactly mimics the dynamics of the original Cilk scheduler.

<sup>\*</sup> This work appeared in preliminary form in the *Proceedings of the 12th Annual ACM Symposium on Parallel Algorithms and Architectures (SPAA)*, pages 13–21, Bar Harbor, Maine, July 9–12, 2000 [7]. M. A. Bender was supported in part by HRL Laboratories, ISX Corporation, NSF Grant EIA-0112849, Sandia National Laboratories, and Grant NSF-CCR-97-00365 at Harvard University. M. O. Rabin's work was supported in part by Grant NSF-CCR-97-00365 at Harvard University.

# 1. Introduction

One of the basic problems in parallel computing is how to execute a parallel program on a collection of *heterogeneous* processors, that is, processors of different and possibly changing speeds. In this paper we focus on the scheduling issues that arise when processors are heterogeneous. We develop scheduling algorithms that are designed to run efficiently in parallel computing environments. We consider general parallel computing environments, but with a particular focus on the Cilk platform [10].

One of the most important constraints of the parallel setting is that the schedulers must make *rapid* decisions about how to assign tasks to processors; otherwise, the time to run the scheduler may actually delay the execution of the parallel program. These scheduling decisions must be made with only partial knowledge of the actual scheduling problem because both the structure of the parallel program and the speeds of the processors are only known *online*, that is, as the computation unfolds.<sup>1</sup> Furthermore, the entire state of the system is not automatically visible to any processor. Each processor *i* is only aware of its *own* local state; in order to determine the state of another processor *j*, processor *i* must explicitly communicate with *j* and this communication has a cost. Consequently, a *centralized* scheduler, which repeatedly gathers all the information about the states of the processors, may be too expensive. This paper describes a scheduling algorithm that is distributed.

Our scheduler is optimized for the following pattern of speed changes, which seems to be the common case in parallel computing environments.

- 1. Most of the time the processor speeds are fairly consistent, and therefore a processor can maintain a *good estimate* of its own speed. This estimate naturally is not completely accurate, but most of the time it will be mostly accurate.
- 2. Processor speeds may occasionally change dramatically, but these changes are limited. The efficiency of our scheduler is allowed to degrade gradually as processors become more erratic.

The model in this paper is a bridge between asynchronous parallel computing and scheduling theory; these two fields attack the general problem of executing parallel programs on processors of different speeds. However, both of these fields make assumptions that differ dramatically from the parallel setting described above. For example, in asynchronous parallel computing the processor speeds are assumed to change arbitrarily and adversarially. This worst-case assumption is often too pessimistic and may lead to inefficient schedules. In scheduling theory the processor speeds are assumed to remain constant, and the scheduler is provided with global knowledge of the state of the system, a large amount of time to run, and offline knowledge of the structure of the computation. Based on these assumptions, the system is unrealistically predictable and the scheduler is unrealistically powerful.

<sup>&</sup>lt;sup>1</sup> In some special cases, such as numerical algorithms, the structure of the parallel program may be known in advance. This paper considers general parallel computations (e.g., parallel chess programs) and does not assume that the programmer provides the running times of the parallel tasks and a mapping from tasks to processors.

We further describe why it is useful to bridge these fields and then proceed to the main results in this paper.

# 1.1. Asynchronous Parallel Computation

Executing parallel programs on heterogeneous processors is studied intensely in the area of *asynchronous parallel computation* [20], [19], [34], [32], [28], [5], [3], [2a], [2b]. In this field the goal is to run a parallel program that is written assuming synchronization barriers, on a collection of asynchronous processors that do not have a synchronization primitive.

Processors are assumed to be *arbitrarily erratic*. That is, a processor may initially run so slowly that it is essentially stopped, change speed abruptly so that it runs extremely (even infinitely) fast, and then stop once more. Correctness proofs typically assume that processor speeds are determined by an adversary, whose goal is to prevent the parallel program from executing correctly or efficiently. Because processors may change speeds to an arbitrary degree, processors are not assumed to have knowledge of their own speed.

The machinery of asynchronous parallel computation is useful for mission critical applications, in which a program must run correctly and steadily, regardless of the erratic behaviors of the individual processors. On the other hand, it may not be worth paying the overhead of these schemes if the application is not mission critical; similarly, it may not be worth paying the overhead if the processors are not arbitrarily erratic, that is, if they change speeds, but most of the time by too much.

#### 1.2. Scheduling on Related Processors

Executing a parallel program on heterogeneous processors is a common problem in scheduling theory. In this field there is an underlying assumption that processors may have different speeds but that the speeds do not change. The goal is to schedule a parallel program represented as a directed acyclic graph (dag) to minimize the *makespan*, that is, the maximum completion time of the jobs. Using terminology from scheduling theory, the problem is that of *scheduling precedence-constrained tasks on related processors to minimize the makespan*.

Because this problem is NP-hard [35] even when all processors have the same speed, the scheduling community has concentrated on developing approximation algorithms for the makespan. Early papers introduce  $O(\sqrt{p})$ -approximation algorithms [23], [24], and more recent papers propose  $O(\log p)$ -approximation algorithms [16], [17], [14], [15]. Unfortunately, some common assumptions from scheduling theory often do not apply to parallel computing, and consequently many scheduling algorithms from this field are not usable in our setting. For example, many of these scheduling algorithms run *offline*, that is, after seeing the entire structure of the parallel program. In addition, the schedulers usually have full knowledge about the state of the system and have the unlimited ability to apply the scheduling decisions.

Finally the quality of many of the scheduling algorithms are measured using the approximation ratio. Even in the *homogeneous setting*, i.e., when all processors run at the same speed, it is known that the approximation ratio may be misleading [12] by a factor as large as 2. The approximation ratio is dramatically less reliable when processors are heterogeneous for several reasons that we describe shortly.

#### 1.3. The Heterogeneous Setting

To develop intuition about the heterogeneous setting, consider the natural class of *greedy schedules*, in which no processor is allowed to stay idle if there is a task that can be assigned to it. When processors are homogeneous, all greedy schedules have essentially comparable makespans (within a factor of 2 of each other). However, when processors are heterogeneous there may be an unbounded ratio between the makespan of the best greedy schedule and the makespan of the worst greedy schedule. To obtain a schedule having a good makespan, fast processors should be assigned to longer paths in the dag and slower processors should be assigned to shorter paths. This assignment process is computationally difficult because nodes in the dag may belong to many interleaving paths of different lengths.

Thus, for any p homogeneous processors, consider p heterogeneous processors that have the same average speed. The optimal makespan in the heterogeneous setting may be much smaller than in the homogeneous setting. However, practical and computational limitations usually prevent this elusive schedule from being found. On the other hand, it is easy to encounter a poor schedule, especially when the processors' speeds can change. This is why users prefer homogeneous processors to heterogeneous ones, even though in ideal conditions the heterogeneous processors may allow shorter schedulers. Thus, in this paper the objective of an efficient scheduler is to use its heterogeneous processors *as efficiently as if they were homogeneous*.

### 1.4. Results

We present the following results:

1. We provide a new analysis of an old scheduling algorithm called the *maximum utilization scheduler* [23]. In particular, we prove a bound on the makespan and on the number of preemptions. Based on this analysis, we generalize this scheduling policy and define the *high utilization scheduler*. We explain why these scheduling policies have close to optimal makespans on dags that represent most parallel programs.

The algorithms presented so far are not directly implementable because the schedulers require too much centralized control. However, they provide insight into how to schedule parallel programs on heterogeneous systems.

2. We next focus on the Cilk platform and present the main result of the paper. We introduce a new algorithm for scheduling Cilk multithreaded parallel programs on heterogeneous processors. This scheduler is inspired by the high utilization scheduler, modified to fit in a Cilk context. A crucial aspect of our algorithm is that it retains the original spirit of the Cilk scheduler. In fact, when our new algorithm runs on homogeneous processors, it exactly mimics the dynamics of the original Cilk scheduler.

#### 1.5. Definitions and Notation

There are p processors labeled 1, ..., p where processor i has speed  $\pi_i$  steps/time. For the sake of convenience, we assume that  $\pi_1 \ge \pi_2 \ge \cdots \ge \pi_p$ . In much of the paper

we assume that the processor speeds do not change; later we mention how our solutions behave when speeds change. Let  $\pi_{tot}$  steps/time be the *total computing power* of all of the processors, that is,  $\pi_{tot} = \sum_{i=1}^{p} \pi_i$ . Let  $\pi_{ave}$  steps/time be the *average speed* of the processors, that is,  $\pi_{ave} = \pi_{tot}/p$ .

A dag G = (V, E) describes the structure of a parallel program. The nodes of the dag represent *tasks* that the processors must complete, and the edges represent *dependencies* between the tasks. Thus, if there is an edge  $(u, v) \in E$ , then v cannot be executed until after u completes. In this case we say that u is a *parent* of v. Tasks are grouped into larger segments of code called *threads*; a *thread* is a path in the dag, where all nodes in the thread, except possibly the first and the last, have outdegree and indegree of 1.

A series parallel dag G = (V, E) is a directed acyclic graph with two distinguished vertices, a source s and a sink t. The family of series parallel graphs are described using the following grammar. A series parallel dag G = (V, E) is one of the following: (1) A single edge extending from s to t, that is,  $V = \{s, t\}$  and  $E = \{(s, t)\}$ . (2) Two series parallel graphs  $G_1$  and  $G_2$  composed in parallel. The sources  $s_1$  and  $s_2$  of  $G_1$  and  $G_2$ , respectively, are merged into a single source s and the sinks  $t_1$  and  $t_2$  of  $G_1$  and  $G_2$  are merged into a single sink t. (3) Two series parallel graphs  $G_1$  and  $G_2$  composed in series. The sink  $t_1$  of  $G_1$  and the source  $s_2$  of  $G_2$  are merged into a single node.

Cilk parallel programs are modeled by *fully strict dags*. A fully strict dag is series parallel, all of the nodes in the dag have outdegree at most 2, and there is one node with indegree 0 and one node with outdegree 0. The *root thread* is a path extending from the first node in the dag to the last node. A node in the root thread with outdegree 2 *spawns* another thread, which continues until it joins the root thread once more. This thread may spawn *child threads*, which may in turn spawn other child threads.

Let  $W_1$  represent the *total work*, that is the total number of nodes in the dag G. Let  $W_\infty$  represent the *critical path length* of the graph, that is, the number of nodes in the longest chain in G. Consider a modified dag G' in which all nodes with indegree and outdegree of 1 are removed, that is, all paths of such nodes are replaced by a single edge. Let  $S_1$  represent the total number of edges in G', and let  $S_\infty$  be the critical path in G'. Let  $T_p$  represent the *time* to execute G on p processors. A task or thread is *ready* if all of its predecessors in G have been executed.

We say that a thread is *preempted* if it is interrupted and later resumed, possibly on a different processor. We say that there is a *migration* whenever the state of the system is moved from one processor to a different processor. Thus, there may be a migration if a previously idle processor begins executing a thread because the processor may have obtained the thread from another processor. There is *not* a migration if a processor finished executing a thread and then executes a successor thread in the dag. Thus, there may be a migration without a preemption, or a preemption without a migration. All migrations entail an additional cost, which we take into account.

We say that an event *E* occurs with high probability if for any c > 0 there exists a proper choice of constants such that  $Pr{E} \ge 1 - n^{-c}$ .

#### 1.6. Related Work

Graham [21], [22] proved that a *list schedule* is a (2-1/p)-approximation to the optimal makespan, and this result holds for any greedy schedule. (In a *list schedule* the jobs have

fixed priorities and the processors execute the ready tasks in the system with the highest priorities.) This results derives from the following theorem:

**Theorem 1** [21], [22], [13]. A greedy schedule (or list schedule) has makespan

$$T_p \leq \frac{W_1}{p} + \left(\frac{p-1}{p}\right) W_{\infty}.$$

Jaffe [23] shows that the following preemptive scheduling policy, called a *maximum utilization schedule*, is a  $O(\sqrt{p})$ -approximation algorithm. At all times maintain the following invariant: if there are i, i < p, ready threads, assign these threads to the i fastest processors. Note that threads may be *preempted*; that is, in the middle of the execution of a thread, a faster processor may take up the responsibility for executing the thread. Jaffe [24] then showed that the following nonpreemptive is also a  $O(\sqrt{p})$ -approximation algorithm for the makespan. Consider the following two schedules and select the one having the better makespan: (1) assign all jobs to the fastest processor, and (2) assign all jobs greedily to processors having speed faster than half the average. More recently, Chudak and Shmoys [16], [17] obtained a  $O(\log p)$ -approximation by using a linear programming relaxation to decide at which speed each task should run. Chekuri and Bender [14], [15] developed a combinatorial approximation algorithm having the same asymptotic approximation ratio.

*Cilk Scheduler.* Cilk is a parallel system with a scheduler that has provable performance guarantees. The Cilk scheduling algorithm is entirely distributed and uses the idea of *work stealing*. Namely, if a processor is idle, it randomly chooses another processor, checks if the processor has extra work, and if so, steals some. The work is stolen in a way that avoids a large increase in memory usage or in running time. The Cilk scheduler works as follows. Each processor maintains a double-ended queue, which is called a *ready deque*. Threads can be inserted and removed from either end of the ready deque. If a processor has no local work to do, it begins work stealing. The processor uses its own ready deque as a stack but other processors' deques as queues. Each processor *i* operates as shown in Figure 1. (For a more complete introduction to the Cilk scheduler see for example [12] and [10].)

#### 2. High Utilization Schedules

We now provide a new analysis of the maximum utilization scheduling policy. This scheduler maintains the following invariant. During each time interval in which there are exactly *i* ready threads, for each i < p, the fastest *i* processors execute these tasks. If there are  $i \ge p$  ready threads, then all of the processors work. Beyond this basic restriction, any processor may execute any task. Note that in order to maintain this invariant, the scheduling policy must allow preemptions.

The maximum utilization scheduling policy is a  $O(\sqrt{p})$ -approximation algorithm, but there are other scheduling algorithms that have comparable approximation ratios and that do not even require preemption. Thus, the advantages of the maximum utilization

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- 1. The processor chooses a victim processor *j* uniformly at random.
- 2. If the victim *i*'s ready deque is empty, processor *i* attempts to steal again.
- 3. Otherwise, it steals the thread T from the *top* of the deque and begins executing it. The processor begins working on thread T until one of three situations:
  - (a) Thread *T* spawns a thread T'. In this case the processor puts *T* on the *bottom* of the ready deque and starts work on thread T'.
  - (b) The thread *T* returns or terminates. If the deque is not empty, the processor begins working on the *bottom* thread. If the deque is empty, it tries to steal and execute thread *T*'s parent. Otherwise, if the parent is busy, the processor attempts to work steal.
  - (c) The thread reaches a synchronization point. In this case the processor attempts to work steal. (Note that the deque is empty.)

Fig. 1. The Cilk scheduler.

scheduler are more subtle, and consequently this scheduling strategy has hardly been revisited. However, many of the other scheduling strategies suffer from the following drawbacks: either they are too complicated to be implemented efficiently, or they produce schedules that are qualitatively unsatisfactory.

The maximum utilization scheduler has a straightforward generalization, which we call a *high utilization scheduler*. In this scheduler we relax the invariant so that at all times: if there are *i*, *i* < *p*, ready threads, the fastest idle processor is at most  $\beta$  times faster than the slowest busy processor. Thus, when  $\beta = 1$ , we obtain a maximum utilization schedule. This makespan of a high utilization schedule appears inferior to the makespan of a maximum utilization schedule, but may have the advantage of fewer preemptions.

We demonstrate two advantages of high utilization schedules: (1) in the common case in parallel computing, high utilization schedules are almost optimal, and (2) they convey a straightforward message to practitioners, run your parallel program on the fastest processors that you can find, and this may be all the optimization that is required. On actual systems such as the Cilk platform, the unembellished high utilization schedule may be too complicated to implement. However, the straightforward concept of using the fastest available processors can be generalized. Thus, high utilization strategies are important because of the guidance that they give in actual situations.

**Theorem 2.** Any maximum utilization schedule has makespan

$$T_{p} \leq \frac{W_{1}}{p \,\pi_{\text{ave}}} + \left(\frac{\pi_{2}}{\pi_{1}} + \frac{\pi_{3}}{\pi_{2}} + \dots + \frac{\pi_{p}}{\pi_{p-1}}\right) \frac{W_{\infty}}{p \,\pi_{\text{ave}}} \leq \frac{W_{1}}{p \,\pi_{\text{ave}}} + \left(\frac{p-1}{p}\right) \frac{W_{\infty}}{\pi_{\text{ave}}}$$

*Proof.* We introduce an accounting tool. We postulate p - 1 disjoint *shadow threads*  $ST_2, ST_3, \ldots, ST_p$ . Each shadow thread is an imaginary chain of tasks. When a processor

*i* is unable to do any work on an *actual thread*, we say that the processor begins working on its *shadow thread*  $ST_i$ .

Consider any time interval in which processor *i* is idle and thus working on its shadow thread  $ST_i$ . Since not all processors have actual work, we are assured that progress is being made on the critical path at the rate of the slowest working processor. That is, since only faster processors  $1 \cdots i - 1$  may be working on the computation, the critical path is advancing at a rate of at least  $\pi_{i-1}$  steps/time.

Because the critical path has length  $W_{\infty}$ , processor *i* can work on  $ST_i$  for  $\pi_i/\pi_{i-1} W_{\infty}$  time units. Processor 1 is never idle. Therefore the total amount of work the processors dedicate to actual and shadow threads is at most  $W_1 + (\pi_2/\pi_1 + \pi_3/\pi_2 + \cdots + \pi_p/\pi_{p-1}) W_{\infty}$ . Because the processors operate at  $\pi_{tot}$  steps/time we obtain the desired bound.

Note that from Theorem 2, we obtain Theorem 1 as a corollary. The makespan can be marginally improved by placing processors on threads more strategically. Namely, put the *i*th fastest processor on the *i*th longest critical path. This policy guarantees that the critical path progresses at least at the average speed of the working processors.

**Claim 3.** Suppose that the maximum utilization strategy additionally maintains the invariant that the ith fastest processor executes the thread that is ith farthest from the end of the dag. This amounts to putting the fastest processor on the critical path. Then the computation has makespan

$$T_p \leq \frac{W_1}{p \,\pi_{\text{ave}}} + \left[\frac{\pi_2}{\pi_1} + \frac{2 \,\pi_3}{\pi_1 + \pi_2} + \frac{3 \,\pi_4}{\pi_1 + \pi_2 + \pi_3} + \cdots + \frac{(p-1) \,\pi_p}{\pi_1 + \pi_2 + \cdots + \pi_{p-1}}\right] \frac{W_\infty}{p \,\pi_{\text{ave}}}.$$

*Proof.* As in Theorem 2, we introduce p-1 disjoint *shadow threads*  $ST_2$ ,  $ST_3$ , ...,  $ST_p$ , where each shadow thread is an imaginary chain of tasks. When a processor *i* is unable to do any work on an *actual thread*, we say that the processor works on its *shadow thread*  $ST_i$ .

Consider any time interval in which processor *i* is idle and thus working on its shadow thread  $ST_i$ . Since not all processors have actual work, progress is being made on the critical path at least as fast as the *average* speed of the working processors. That is, since only faster processors  $1 \cdots i - 1$  may work on the computation, the critical path advances at a rate of at least  $(\pi_1 + \pi_2 + \cdots + \pi_{i-1})/(i-1)$  steps/time.

Because the critical path has length  $W_{\infty}$ , processor *i* can work on  $ST_i$  for  $\pi_i$   $(i - 1)/(\pi_1 + \pi_2 + \cdots + \pi_{i-1}) W_{\infty}$  time units. Processor 1 is never idle. Therefore the total amount of work the processors dedicate to actual and shadow threads is at most

$$W_1 + \left[\frac{\pi_2}{\pi_1} + \frac{2\pi_3}{\pi_1 + \pi_2} + \frac{3\pi_4}{\pi_1 + \pi_2 + \pi_3} + \cdots + \frac{(p-1)\pi_p}{\pi_1 + \pi_2 + \cdots + \pi_{p-1}}\right] W_{\infty}.$$

Because the processors operate at  $\pi_{tot}$  steps/time we obtain the desired bound.

Unfortunately, this gain in makespan seems small in comparison with the potentially infinite number of additional preemptions that this policy entails.

The proof of Theorem 2 extends to prove the following theorem that provides a bound on the makespan of a high utilization schedule.

**Theorem 4.** Any high utilization schedule has makespan

$$T_p \leq \frac{W_1}{p \,\pi_{\text{ave}}} + \left(\frac{p-1}{p}\right) \frac{\beta \, W_\infty}{\pi_{\text{ave}}}.$$

We now provide a bound on the number of migrations in a high utilization schedule.

**Theorem 5.** Consider a high or maximum utilization schedule of an arbitrary dag. If there are a total of  $S_1$  threads, then there are at most  $2S_1$  migrations.

*Proof.* We divide the computation into phases,  $S_1$ ,  $S_1 - 1, ..., 2$ , 1, where in phase  $\Pi$  the computation has  $\Pi$  (incomplete) threads. Within a phase, a computation has no migrations at all. A phase begins when the number of *active threads* (e.g., threads currently being executed by processors) changes.

Assume without loss of generality that at most one thread completes at any time. (If two threads complete simultaneously, we break the tie arbitrarily.) There are two cases for the dynamics of the schedule when a thread completes. (1) When a thread  $T_{\alpha}$  completes, no new threads active become active. Then the slowest currently active processor *k* migrates to the idle pool, and the processor *j* on  $T_{\alpha}$  migrates to *k*'s thread. (If we are lucky, the slowest currently active processor *k* is already on thread  $T_{\alpha}$ .) (2) When a thread  $T_{\alpha}$  completes, *x* new threads become active. Then x - 1 processors migrate from the idle pool to a new active thread and one processor migrates from the completed thread  $T_{\alpha}$  to a new active thread.

If each migration requires an extra cost of M, then we have a bound on the increase in makespan from Theorem 6 when migrations have a cost, namely  $2MS_1/p$ . The quantity M may include the cost to send the system state from one processor to another or may even include the cost to restart a thread from some previous checkpoint.

Theorem 2, Claim 3, and Theorem 4, which bound the makespan of maximum and high utilization schedules, hold even when the speeds of processors change. Theorem 5, however, no longer applies. Instead, the number of migrations increases as the processors become more erratic. An open question is to choose the value of  $\beta$  that optimizes the makespan while avoiding too many migrations.

#### 2.1. Performance in the Common Case

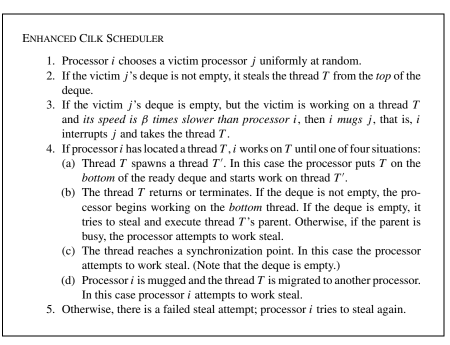
Even though the high utilization schedule is a  $O(\sqrt{p})$ -approximation algorithm for general dags, on dags that represent most parallel programs, the algorithm has a substantially better performance. In most parallel programs  $W_1/p \gg W_{\infty}$  [12]. An interpretation of this inequality is that the parallel program has enough inherent parallelism to justify the use of *p* processors. Observe that in Theorems 2 and 4,  $W_1/\pi_{tot}$  is a lower bound on

the makespan, and when  $\beta > 1$  is sufficiently close to 1, this quality dwarfs  $\beta W_{\infty}/\pi_{ave}$ . Therefore, even though the high utilization schedule is a  $O(\sqrt{p})$ -approximation for general dags, in the case of dags representing typical parallel programs, it is almost optimal. This closeness to optimal is not true of the nonpreemptive  $O(\sqrt{p})$ -approximation algorithm.

# 3. An Enhanced Cilk Scheduler

Direct implementation of the the scheduling policies in the previous section are impractical because they rely on global control. However, the general design principle of high utilization is critical, and we apply this concept in Cilk scheduling. In this section we describe an enhanced Cilk scheduler that runs correctly and robustly even when processors have different speeds. Moreover, when the processors run at *similar* speeds, our new schedule behaves identically to the standard Cilk scheduler. Thus, an important feature of our scheduler is that it is extremely similar to the original scheduler at a small cost in algorithmic complexity.

In this algorithm there are two kinds of migrations: *steals* and *muggings*. In a steal, a processor begins working on a thread at the top of another processor's ready deque. In a mugging, there is no work on another processor's ready deque, and so the processor "mugs" a processor that is slower by at least a  $\beta$  factor and takes the thread that the slower processor was working on. The pseudocode for the Enhanced Cilk Scheduler appears in Figure 2.





If all processors operate at speeds within a  $\beta$  factor of each other, then there are no muggings and the scheduler behaves like the standard Cilk scheduler. The parameter  $\beta$  can be tuned to optimize system performance.

Indeed, it is not even necessary to define a particular value of  $\beta$ . That is, our algorithm still works for any  $\beta > 1$ , i.e., processor *i* mugs processor *j* only if  $\pi_i > \pi_j$ . The advantage of introducing  $\beta$  is that it reduces the number of migrations. Optimizing the value of  $\beta$  is a topic of future work.

#### 3.1. Design Assumptions and Changing Speeds

We make the following additional assumptions: (1) Each processor steals at a rate proportional to its speed. (2) Steals and steal attempts are completed in an amount of time that is proportional to the speed of the processor doing the stealing/mugging. It is important that the steal responses on the platform do not depend on the speed of the victim processor because otherwise the slowest processor can delay the entire system.<sup>2</sup> There are several ways to ensure this design principle. For example, there might be a bound on the ratio between the fastest and slowest processor. We could also require some mechanism for communicating steal attempts, such as a shared memory, that allows one processor to look directly into the deques of other processors.

The Enhanced Cilk Scheduler is designed to be efficient when speeds change. This is because the scheduler relies on brief interactions between pairs of processors, rather than global control. The processors do not have to store information about the speeds of other processors, which might quickly become out of date. However, as the processors become more erratic, there may be additional steals and muggings.

The following section bounds the running time and number of steals and muggings in the case when the processors speeds do not change by too much. The performance of the algorithm degrades gracefully as the speeds become more erratic. An important open question is to optimize the value of  $\beta$  to remove unnecessary muggings.

#### 3.2. Analysis

We now analyze the running time of the Enhanced Cilk Scheduler. We prove the following performance guarantee.

**Theorem 6.** With high probability the execution time  $T_p$  of the enhanced Cilk Scheduler is bounded as follows:

$$T_p \leq \frac{W_1}{p \,\pi_{\mathrm{ave}}} + O\left(\frac{W_\infty}{\pi_{\mathrm{ave}}}\right).$$

We use an accounting argument to prove Theorem 6. Observe that at all times a processor is either (1) executing an instruction, or (2) attempting to steal (and perhaps actually

<sup>&</sup>lt;sup>2</sup> If the steal attempts run at the speed of the victim processor, then the work-stealing approach cannot have guaranteed good performance. This is because the root thread of the computation may reside on a processor that is entirely stopped, and the computation cannot proceed.

stealing or mugging). For simplicity of analysis, we assume that each of these operations requires one unit of work. (In fact, executing an instruction is likely to be much faster and so in our analysis we can group multiple instructions together.)

We postulate two *buckets* that we use for accounting, a *work bucket* and a *steal bucket*. Each time a processor completes a unit of work on the dag it puts one dollar into the work bucket; each time a processor completes a steal attempt (successful or not) it puts one dollar into the steal bucket. (This approach was used in the original paper of [12] and in much of the subsequent work on Cilk.) There are  $\pi_{tot}$  dollars that enter the buckets per unit of time. Therefore, if at the end of the computation, there are a total of *D* dollars in both buckets, then the computation ran in time  $D/\pi_{tot}$ .

Computing the number of dollars in the work bucket is straightforward, because each time the processor completes one unit of work, it puts a dollar in the work bucket.

**Observation 1.** At the end of the computation there are a total of exactly  $W_1$  dollars in the work bucket.

We now use a potential-function argument to prove a bound on the number of dollars in the steal bucket. This argument is an extension of the result in [1] and [9] and begins with some definitions.

**Definitions.** For any (nonroot) node v, suppose that node u is the last of v's parents to be executed. Then we say that the execution of node u enables node v. Node u is called the *designated parent* of v and edge (u, v) is called the *enabling edge*. The graph composed of all the enabling edges is called the *enabling tree*. The node that is being executed at time t by processor i is called the *assigned node of processor* i *at time* t. We assign weights to all of the nodes, so that we can use these weights in a potential function argument. Let d(u) denote the *depth* of node u in the dag, i.e., the distance to the root node. Each node u has weight  $w(u) = W_{\infty} - d(u)$ , so that nodes closer to the root have larger weight.

We now present the potential function from [1] and [9], which we will use. Let  $R_t$  be the set of ready nodes at time t. Each node is either in some deque or assigned to and executed on some processor. For each ready node  $v \in R_t$ , we define its potential  $\varphi_t(v)$  as

 $\varphi_t(v) = \begin{cases} 3^{2 \cdot w(v) - 1} & \text{if } v \text{ is assigned;} \\ 3^{2 \cdot w(v)} & \text{otherwise.} \end{cases}$ 

We let  $\Phi_t(i)$  denote the sum of the potentials of the nodes on processor *i* at time *t*. We let  $\Phi_t = \sum_{i=0}^{p} \Phi_t(i)$  be the value of the potential function at time *t*. Thus, the initial potential is  $3^{2 \cdot W_{\infty}}$  because the root node has depth 0 and is initially unassigned. The final potential is 0 because all nodes have been completed.

Now supplied with these definitions, we show that the Structural Lemma of the deques from [1] and [9] still holds in the heterogeneous setting. This lemma guarantees that for any deque at all times during the execution of the work-stealing algorithm, the

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designated parents of the nodes in the deque lie on the root-to-leaf path in the enabling tree.

**Lemma 7** [1], [9]. Let k be the number of (ready) nodes in a given deque at any time t, and let  $v_1, v_2, \ldots, v_k$  denote these nodes ordered from bottom to top. Let  $v_0$  be the assigned node. In addition, for  $i = 1 \cdots k$ , let  $u_i$  be the designated parent of  $v_i$ . Then for  $i = 1 \cdots k$ , node  $u_i$  is an ancestor of  $u_{i-1}$  in the enabling tree. Moreover, although it may be that  $u_0 = u_1$ , for  $i = 2 \cdots k$ ,  $u_{i-1} \neq u_i$ . Thus, the weights of the nodes increase from bottom to top, that is,  $w(v_0) \leq w(v_1) < w(v_2) < \cdots < w(w_k)$ .

*Proof.* The proof is by induction on times in which the structure of the deque changes, as in [1] and [9]. There are five possible ways that the deque may change: (S) the top node of the deque is stolen; (E0) the assigned node enables 0 children; (E1) the assigned node enables 1 child; (E2) the assigned node enables 2 children; (M) The processor is mugged and the assigned node is moved to a faster processor. The first four cases are described and analyzed in the proof in [1] and [9].

The case of muggings, which is unique to the heterogeneous setting, is trivially integrated into the correctness proof. After a mugging, the mugged processor has no assigned tasks and an empty deque, and the mugging processor has an assigned task but an empty deque. Thus, the claim follows trivially in the case of muggings because there is that most one node.

The Structural Lemma enables us to prove the following observation:

**Observation 2** [1, Lemma 6]. For any processor at time t during the execution of the scheduling algorithm, the potential of the topmost nodes in the deques contributes at least three-quarters of the potential associated with the processors that have nonempty deques.

We now divide the computation into *phases*, which are defined inductively by when steal attempts occur. The first phase begins at time t = 0, the start of the computation, and it ends after  $(\beta + 2)p$  steal attempts have occurred. (Recall the definition of  $\beta$ : in order for a processor *i* to mug a processor *j*, it must be that  $\pi_i > \beta \pi_j$ .) The *i*th phase begins at the end of the (i - 1)th phase and completes after  $(\beta + 2)p$  additional steal attempts have been made.

**Theorem 8.** There is at least a constant probability that within each phase, the potential drops by at least a constant factor. Therefore, there are at most  $O(\log n)$  phases, both expected and with high probability.

*Proof.* At any time *t* we partition the potential  $\Phi_t = D_t + S_t + F_t$  into three disjoint components. The component  $D_t$  is associated with processors whose deque contains nodes. The rest of the potential is associated with processors that have empty deques, but which may have assigned nodes. We divide this remaining potential into components associated with processors we define as *slow* and *fast*, respectively. A processor *i* is

called *slow in phase*  $\ell$ , if, during phase  $\ell$ , the processor does not have time to finish executing the node that it was working on when the phase began. A processor *i* is called *fast* otherwise.

We first consider the potential  $D_t$  associated with the set of processors whose deques are *not* empty. Recall that at least three-quarters of the potential from nodes in the deques is exposed to steals at the top of the deques. Consequently, because there are  $(2 + \beta)p$ steal attempts in any phase and the probability that a given steel attempt does not steal from a given deque is (1 - 1/n), the probability that there is no steal attempt in a deque is at most  $e^{-(2+\beta)}$ . When the node at the top of the deque is stolen, the potential of this node decreases by a factor of two-thirds because the node is now assigned to a processor.

Let value Q be the sum of the potentials of the nodes at the top of the deques. Then the expected value of the remaining potential of these nodes after the phase ends is at most  $e^{-(2+\beta)}Q + (1 - e^{-(2+\beta)}) 2Q/3$ . Therefore, by the Markov inequality, there is at least a constant probability that the potential associated with these nodes decreases by at least a constant factor. Consequently, by Observation 2, with at least a constant probability the potential associated with all the nodes in those deques decreases by at least a constant factor.

We now examine the component  $F_i$  of the potential, that is, the potential associated with fast processors having empty deques at the start of phase  $\ell$ . For any such processor *i*, the completion of *i*'s assigned node causes the potential to decrease by at least a constant factor because *i*'s original assigned node will be completed.

Finally, we examine the component  $S_t$  of the potential, that is, the potential associated with slow processors having empty deques at the start of phase  $\ell$ . In order to reduce the potential of a slow processor *i* that contributes to  $S_t$ , another processor *j* must (1) choose to mug processor *i*, and (2) complete one node of the thread that it obtained from processor *i*. In order to mug *i*, processor *j* must be more than  $\beta$  times faster than processor *i*. How many steal attempts are there in phase  $\ell$  that satisfy these conditions? Any processor that makes  $\beta + 2$  steal attempts in the phase must be more than  $\beta$  times faster than processor *i*, which does not even finish executing one node. Consequently, in  $(\beta + 2)p$  steal attempts, there will be at least *p* steal attempts that satisfy all of these conditions. Therefore, the probability that any given slow processor is not mugged is at most 1/e. Let value Q' be the sum of the potential of the nodes being executed by the slow processors. Then the expected value of the remaining potential of these nodes after the phase ends is at most Q'/e. Therefore, by the Markov inequality, there is at least a constant probability that the potential associated with these nodes decreases by at least a constant factor.

By considering all three cases, we conclude that there is at least a constant probability that the total potential decreases by at least a constant factor. Therefore, an application of Chernoff Bounds [33] demonstrates that after at most  $O(W_{\infty})$  phases the potential has decreased until it is zero, both expected and with high probability.

From Theorem 8, we conclude that there are at most  $O(\beta W_{\infty} p)$  steal attempts and consequently  $O(\beta W_{\infty} p)$  dollars in the steal bucket. Therefore, the running time of the algorithm is  $W_1/(p\pi_{ave}) + O(\beta W_{\infty}\pi_{ave})$ , which finishes the proof of Theorem 6.

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#### Acknowledgment

The first author warmly thanks Charles Leiserson for suggesting this problem, for enjoyable meetings in the earlier stages of this work, and for much excellent advice.

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Online publication May 14, 2002.