### Efficient Resource Scheduling in Multiprocessors

by

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### Abstract

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Professor Katherine Yelick, Computer Science, Chair

As multiprocessing becomes increasingly successful in scientific and commercial computing, parallel systems will be subjected to increasingly complex and challenging workloads. To ensure good job response and high resource utilization, algorithms are needed to allocate resources to jobs and to schedule the jobs. This problem is of central importance, and pervades systems research at diverse places such as compilers, runtime, applications, and operating systems. Despite the attention this area has received, scheduling problems in practical parallel computing still lack satisfactory solutions. The focus of system builders is to provide functionality and features; the resulting systems get so complex that many models and theoretical results lack applicability.

The focus of this thesis is in between the theory and practice of scheduling: it includes modeling, performance analysis and practical algorithmics. We present a variety of new techniques for scheduling problems relevant to parallel scientific computing. The thesis progresses from new compile-time algorithms for message scheduling through new runtime algorithms for processor scheduling to a unified framework for allocating multiprocessor resources to competing jobs while optimizing both individual application performance and system throughput.

The compiler algorithm schedules network communication for parallel programs accessing distributed arrays. By analyzing and optimizing communication patterns globally, rather than at the single statement level, we often reduce communication costs by factors of two to three in an implementation based on IBM's High-Performance Fortran compiler.

The best parallelizing compilers at present support regular, static, array-based

parallelism. But parallel programmers are out-growing this model. Many scientific and commercial applications have a two-level structure: the outer level is a potentially irregular and dynamic task graph, and the inner level comprises relatively regular parallelism within each task. We give new runtime algorithms for allocating processors to such tasks. The result can be a twofold increase in effective megaflops, as seen from an implementation based on ScaLAPACK, a library of scientific software for scalable parallel machines.

Compilers and runtime systems target single programs. Other system software must do resource scheduling across multiple programs. For example, a database scheduler or a multiprocessor batch queuing system must allocate many kinds of resources between multiple programs. Some resources like processors may be traded for time, others, like memory, may not. Also, the goal is not to finish a fixed set of programs as fast as possible but to minimize the average response time of the programs, perhaps weighted by a priority. We present new algorithms for such problems.

Most of the above results assume a central scheduler with global knowledge. When the setting is distributed, decentralized techniques are needed. We study how decentralization and consequent local knowledge by per-processor schedulers affects load balance in fine-grained task-parallel applications. In particular, we give new protocols for distributed load balancing and bounds on the trade-off between locality and load balance. The analysis has been supported by experience with implementing task queues in Multipol, a library for coding dynamic, irregular parallel applications.

> Professor Katherine Yelick, Computer Science Dissertation Committee Chair

রবীন্দ্রনাথ ঠাকুর

না গো, এই যে ধূলা আমার না এ। ডোমার ধূলার ধরার °পরে উড়িয়ে যাব সন্ধ্র্যাবায়ে॥ দিয়ে মাটি আগুন জ্বলি রচলে দেহ পূজ্যার থালি— শেষ আরতি সারা করে ভেঙ্গে যোব তোমার পায়ে॥ ফুল যা ছিল পূজ্যার তরে যেতে পথে ডালি হতে অনেক যে তার গেছে পড়ে। কত প্রদীপ এই থালাতে সাজিয়েছিলে আপন হাতে— কত যে তার নিবল হাওয়ায়, পৌঁছলনা চর+ছায়ে॥

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### Chapter 1

## Introduction

Parallelism exists in most forms of computation, and exploiting it is a promising means to boost the performance of many important scientific and commercial applications beyond peak uniprocessor performance. However, exploiting parallelism is not trivial; it often requires complete algorithm redesign and implementation for different parallel machines. It has taken almost a decade to identify a few cost-effective parallel architecture configurations, and as a result, system software is only starting to mature and reach a level of portability and standardization that can motivate application writers to invest in parallel programming.

This point in time is unique in the development of parallel computing, since some basic design issues have been resolved, so that one need not chase moving targets any more, but there is a large scope for algorithm design in systems software. The surviving architectural choices are clear for the foreseeable future: cost-effective parallel machines will be built from shared memory "symmetric" multiprocessors (SMP's). These will scale up to a few dozen processors at most. For applications that demand more resources, these SMP's will be interconnected by a network. Processes on different processors will communicate by passing messages to each other, either explicitly by subroutine calls, or implicitly, by accessing remote data and triggering a software abstraction of global memory to issue messages. In units of CPU cycles, communication overhead is expected to be large. Each message has some fixed startup overhead time and then some proportional time per byte of data transfered. It will be the interconnect that will define the character and the price-tag of the machine; the processors will be commodity.

Apart from interface and portability issues, the challenge for system software, such

as compilers, runtime systems, and application libraries, is to export a clean performance model to the client while adjusting to diverse communication cost parameters and diverse application behavior in terms of both locality and load balance.

The work in this dissertation is based on the following premises.

- *Performance engineering* is of paramount importance in parallel computing, because, "Parallelism is not cheaper or easier, it is *faster*" [66].
- Scheduling the resources of the multiprocessor (mainly, processors, network, memory, and IO activity) judiciously between the units of work is essential for achieving high utilization of the resources. Very few compilers or runtime systems take such a global view of the scheduling problem yet, and we shall show the benefits of doing so throughout the thesis.
- These resource scheduling algorithms must use simple models of the machine and the workload, so that the resulting system is performance-portable across diverse platforms without need for ad-hoc tuning.

The contribution of this thesis is a resource scheduling framework for scientific and commercial applications on scalable multiprocessors, comprising techniques at the compiler, runtime, and application level.

Despite the attention that the area of resource allocation and job scheduling has received, many problems in parallel system scheduling lack satisfactory solutions. First, simple application of existing results to new domains is rarely possible, because the setting often has crucial details not addressed by classical scheduling literature. Second, the practical situation is usually complex, and it is hard to abstract a model that is both simple and faithful to reality. Third, most of the problems are computationally intractable, and knowledge of complexity and approximations is needed to design algorithms and analyze them. Finally, practical algorithmics, reflecting both the constraints and simplifications offered by practical scenarios, are more important than exploiting powerful but impractical primitives from theory.

The rest of this chapter summarizes the problems and new solutions described in the thesis.

### 1.1 Message scheduling via compiler analysis

Parallelism expressed through arrays and loops is very common in scientific numerical applications. Many languages have been designed for expressing array parallelism, commonly called "data parallelism." The success of data parallelism has reached the stage where a language standard called High Performance Fortran (HPF) has been defined. HPF extends Fortran with two features: one can refer to regular array sections rather than individual elements (this extension is also part of Fortran 90 [49]), and give directives to the compiler to distribute and align arrays on parallel machines. The compiler partitions computation to match the data distribution and generates communication for remote arrays. However, compiling HPF code to get performance close to hand-coded message-passing programs is still a major challenge. On scalable multiprocessors such as the IBM SP2 or Intel Paragon, a single extra communication step can cost tens of thousands of CPU cycles.

There is a large body of literature on compiler analysis and optimization of communication generated by loops involving distributed arrays. Initial progress was mostly restricted to local optimizations at the single loop-nest level; these could not detect and eliminate communication that is redundant between different loop nests or subroutines. As a simple example, a single value of a distributed array could be used in several loops after its definition throughout a subroutine; if the array value is unchanged, communication for the uses should be done once after the definition of the value, but without global analysis it would be done before each loop enclosing an use.

Several factors complicate the situation. The main existing technique to detect redundancy has been to trace uses of values back to the definition and cancel all communication that is rendered redundant by other communication placed at that point. This is called *earliest placement*, and is also good for overlapping communication with computation. However, this technique places communication for a remote array access at a place depending only on that access and not others, which may prevent some highly profitable optimizations like combining messages from multiple arrays into one. While message combining amortizes communication overhead, it requires memory copies for packing and unpacking data. This has to be done carefully so that the communication buffers are smaller than the cache, or the benefits of amortizing the fixed overhead of a message may be canceled by cache miss penalty.

In Chapter 2 we will describe a new compiler algorithm for communication analysis

and optimization that achieves redundancy elimination while maximizing network and cache performance across loops and statements in a global manner [27]. This significantly improves on the recently proposed array dataflow analyses which use the "earliest placement" policy: our algorithm takes into account all legal positions for all remote accesses before deciding on the final placement of any access. One implication of this is that our algorithm may *avoid* early communication placement if it can lead to larger message overhead. We implemented the algorithm in IBM's production compiler, called pHPF. Although pHPF already generated highly optimized code, the new technique cut down communication time by another factor of two, which meant a 20-40% overall savings for many well-known scientific benchmarks.

### 1.2 Scheduling jobs with mixed parallelism

The underlying execution model of HPF and similar compilers is called *bulk synchronous*: there are alternating phases of computation and communication. Commodity networks perform well if the latter are few and far between; the compiler algorithm outlined in the previous section enhances this effect. However, loop-level parallelism still has limited scalability because many processors have to communicate frequently and cooperatively on a data-parallel problem.

Some applications have a natural two-level structure: at the outer level there is a directed acyclic task graph, where each vertex is a task that can run only after all its predecessors have completed. Tasks not ordered with respect to each other can be executed in parallel. At the inner level, each task is parallelizable; it can run on some set of processors provided they work on it simultaneously. (The inner level parallelism is the data parallelism mentioned earlier.) There is a growing body of such applications, including eigensolvers, adaptive mesh codes, circuit simulation, and parallel database query trees. We shall call applications with these two forms of parallelism *mixed* parallel. Recent versions of the HPF language proposal includes support for such applications, and some parallelizing compilers have been extended to express such applications.

Given the importance of mixed parallel applications, it is not surprising that there has been significant research on scheduling them. Most of the work has been in the area of compile-time scheduling. Array parallelism is expressed as in HPF, and task parallelism is typically expressed using annotations. The compiler uses estimates or profiles of computation and communication in the data parallel modules and uses expensive off-line optimization algorithms to allocate processors to modules.

Unfortunately, the amount of static task parallelism in such problems is so small (typically 4–8) that except for the smallest problems, the gains over pure data parallelism are small. Problems where the task parallelism grows with problem size, such as in divide and conquer, are therefore much more promising from the perspective of exploiting mixed parallelism. On the other hand, these applications have to be scheduled dynamically, and therefore cannot use the expensive optimization algorithms, such as linear-programming, that are used in compile-time scheduling.

In Chapter 3 we will describe a simple and effective heuristic for scheduling divide and conquer applications with mixed parallelism [26]. The algorithm classifies the tasks into two types. The large problems near the root are allocated all the processors in turn, while small problems close to the leaves are packed in a task-parallel fashion, each task being assigned exactly one processor. There is some internal frontier at which the execution model *switches* from data to task parallelism. Some previous implementations have used switching based on an ad-hoc switch point tuned to a given problem and machine. In contrast, our switching algorithms are parameterized on machine and problem parameters and are provably close to optimal. Control on the switching frontier also gives a simple means for adjusting to diverse network cost parameters, while also trading communication cost with the load imbalance inherent in the task graph.

We have implemented the switching algorithms as an extension of the ScaLAPACK runtime system and used it to run a non-symmetric eigensolver. A significant performance improvement is seen over a purely data-parallel implementation.

We have also analyzed the performance gap between the switched mode execution (which is simple to program) and the general setting of allocating arbitrary processor subsets to jobs (for which many algorithms exist in theory) for a large class of applications. Our analysis shows that the gap is often small. From a language or runtime design perspective, these results indicate that this limited form of mixed parallelism will produce most of the benefits with a much simpler implementation.

### 1.3 Scheduling resource constrained jobs

Chapters 2 and 3 focus on the performance of a single application on a dedicated machine. In realistic installations many users submit jobs that contend for system resources. Since these jobs arrive over time, it makes little sense to measure the performance of a scheduler by giving it a batch of jobs off-line and measuring the finish time of only the last job (this is called the *makespan* of the set of jobs). A better measure to minimize is the average response time, which is the average time from arrival to completion of a job. For reasons elaborated later, this problem is very hard. A useful step in that direction is to solve the special case of minimizing the prioritized (also called *weighted*) average completion time, WACT, over a fixed batch of jobs.

The situation is complicated by the jobs having diverse running times and resource requirements. In real life, jobs need not only processors (which is the predominant case dealt with in scheduling theory) but also other resources, such as memory and network and disk bandwidth. While some known algorithms can deal with general resources, they assume the jobs are independent, while in many situations, there are precedence constraints between them.

Until recently, few machines allowed multiple jobs to execute concurrently. As most machines are going towards a multiprogrammed full-UNIX node model, resource scheduling is becoming more crucial at the system level. Vendors like IBM, SGI and Convex place significant emphasis on load management software, as do academic communities [50]. It is also becoming apparent that applications should dynamically allocate and deallocate resources to avoid under-utilization and long job response times [48]. While significant software infrastructure is being built, the algorithms at the heart of these systems are often variants of list-based greedy schedulers, which perform poorly under heavy load.

In Chapter 4 we give several simple and near-optimal algorithms for variations of this scenario. They depend on a framework for extending makespan algorithms into WACT algorithms. By applying the framework, we first give new algorithms for processor scheduling under the WACT metric. There appear to be qualitative differences between different types of resources. Some resources, like processors, are *malleable*, i.e., they can be traded for time relatively gracefully, while others, like memory are *non-malleable*, i.e., admitting little or no such flexibility. For jobs with both precedence and non-malleable resource constraints, the problem appears to be very different from most precedence-constrained scheduling results that are based on basic lower bounds like average work per processor or the critical path length in the precedence graph. We obtain an  $O(\log T)$  approximation for both WACT and makespan [28], where T is the longest to shortest job time ratio. Our algorithm uses a technique that deliberately introduces delays to improve on a greedy schedule. We also show that the log T blowup is unavoidable for certain instances. Since our models were carefully abstracted from real-life scheduling problems in parallel databases and operating systems, we expect that these results will prove useful in practice.

### 1.4 Dynamic scheduling using parallel random allocation

The results summarized thus far work well in problem domains where the resource requirements of jobs can be estimated with reasonable accuracy at the time the job is submitted to the scheduler. Sparse, irregular, and dynamic problems do not always have this benign feature. In Chapter 5 we explore randomization as an effective means for dealing with such dynamic and unpredictable problems.

In the first part we consider applications where the tasks each run on only one processor, but may have a precedence relation (the task DAG) unfolding dynamically. The running time of a task may be unknown before completion, and the scheduler has to be *non-clairvoyant*, i.e., it cannot know and use the running time of a job to schedule it. It is not necessary for a centralized scheduler (e.g. Graham's list algorithm [64]) to be clairvoyant, when evaluated using the finish time metric, also called "makespan." But this solution has a communication bottleneck. We analyze the running time using the following decentralized strategy: each processor has a local task pool, new tasks are sent to a random pool, and each processor removes and executes tasks from its local pool. Although used in practice, the effect of random allocation on dynamic, irregular task graphs was unresolved before this work; the closest known analysis was for unit-time tasks [78]. The resulting bounds are supported by practical experience with irregular applications [30, 25], which show that randomization is an effective tool for load balancing a certain class of applications.

It is well-known from the literature on occupancy problems that one round of random allocation leads to roughly a logarithmic smoothing of load. In more concrete terms, if n tasks are randomly assigned to n processors, each processor has  $O(\log n / \log \log n)$  tasks with probability at least 1 - O(1/n). In the second part of Chapter 5 we study an extended

model of random allocation, where assignment of tasks to processors are made in multiple rounds so as to further smooth down the load imbalance to sub-logarithmic amounts. In addition to the task scheduling application, this setting can be motivated by the problem of allocating file blocks in distributed "serverless" file systems like xFS [6]. In this setting there are n client workstations whose local disks comprise the file system, which can be abstracted as n servers. As clients write files, new disk blocks must be allocated in a decentralized way without overloading any particular server. There is a trade-off between the cost of communication to obtain more global information and the cost of load imbalance owing to imperfect knowledge. At one extreme, one can send each new block to a random server; at the other, one can get optimal load balance by maintaining exact global load information. We give a precise characterization of this trade-off [1]. Initial simulation results agree with theoretical predictions.

Extensive work has been done on all the problems studied. In Chapter 6 we will present a survey and classification of known results based on job and machine models. Finally, in Chapter 7 we will summarize the thesis and suggest directions for future work.

### Chapter 2

# Message scheduling via compiler analysis

### 2.1 Introduction

Distributed memory architectures provide a cost-effective method for building scalable parallel computers. However, the absence of a global address space and the resulting need for explicit message passing makes these machines difficult to program. This has motivated the design of languages like High Performance Fortran (HPF) [55], which allow the programmer to write sequential or shared-memory parallel programs that are annotated with directives specifying data decomposition. The compilers for these languages are responsible for partitioning the computation and generating the communication necessary to fetch values of non-local data referenced by a processor [72, 123, 20, 5, 21, 68].

Accessing remote data is usually orders of magnitude slower than accessing local data, for the following reasons. It is getting increasingly cost-effective to build multiprocessors from commodity hardware components and system software. Most current generation CPU's are well beyond the 100 MFLOPS mark, and are consistently out-pacing network performance. Most vendors support UNIX-like environments on each processor, in particular with multiprogramming and virtual memory. This means that low-overhead message passing implementations like Active Messages [115], which work best with gang-scheduled time slicing, user-level access to the network interface, and register-based data transfer, are not the best choice. As a result, communication startup overheads tend

Machine	NW software	$\alpha$	$\beta$
Paragon	NX	$7.8  imes 10^3$	9
Cray T3D	BLT	$2.7 imes10^4$	9
CM5+VU	CMMD	$1.4  imes 10^4$	103
IBM SP1	MPL	$2.8  imes 10^4$	50
IBM SP2	MPL	$1.2  imes 10^4$	60

Table 2.1: Estimates of message startup overhead  $\alpha$  and transfer time per double  $\beta$  scaled to the peak floating point operation time for different machines. Estimates are in part from [107, 117, 88, 7]; the network software are described in these references.

to be very large on most distributed memory machines, although reasonable bandwidth can be supported for sufficiently large messages [109, 106]. See Table 2.1 for some idea of the CPU and communication speeds of current multiprocessors.

Therefore, parallelizing compilers have to be extremely careful about generating communication code. A single suboptimal choice may waste time equivalent to several thousand CPU operations. Unfortunately, communication optimization is a very difficult problem, with many potentially conflicting considerations. Consider a parallel program where a data value is produced by some statement, after which it is sent out of the owner processor, received by other processors, and finally used in some other statement. There is usually some flexibility about the placement of the send and receive subroutine calls between the definition and the use. An early placement favors potential overlap between CPU and network activity, but may consume more temporary communication buffers and cache problems if the main CPU is involved in packing and unpacking data, e.g., for strided accesses. A late placement may break up communication of a large array section into smaller sections or individual elements, suffering increased overhead. Furthermore, placing one communication at some point in the code also affects the choice of the best place for other communication.

There is a clear need for global *scheduling* of communication. "Scheduling," in this chapter, means judicious placement of message-passing subroutine calls in the compiled code by the compiler. In this chapter we present a novel compiler algorithm that recognizes the global nature of the communication placement problem. Our algorithm derives from static single assignment analysis, array dependence analysis, and data availability analysis [69], which is extended to detect compatibility of communication patterns in addition to redundancy. We differ significantly from previous research, in which the position of communication code for each remote access is decided independent of other remote accesses; instead, we determine the positions in an interdependent and global manner. The algorithm achieves both redundancy elimination and message combining globally, and is able to reduce the number of messages to an extent that is not achievable with any previous approach.

Our algorithm has been implemented in the IBM pHPF prototype compiler [68]. We report results from a preliminary study of some well-known HPF programs. The performance gains are impressive. Reduction in static message count can be up to a factor of almost nine. Time spent in communication is reduced in many cases by a factor of two or more. We believe that these are also the first results from any implementation of redundant message elimination across different loop nests, and add significant experimental experience to research on communication optimization.

### 2.2 Motivating codes

We will use a few code fragments to show the importance of recognizing the global nature of the message placement problem. We are interested in the communication patterns and the flexibility in message placement revealed by data dependence analysis, not the particulars of the applications.

In the code fragments that we present, we will elide actual operations and show each right hand side (RHS) as a list of variables accessed. Frequently we deal with Fortran 90 (F90) style shift operations that involve nearest-neighbor communication; we show this pictorially using arrows. For simplicity, the combinable messages in our examples have identical patterns on the processor template; in practice, combining is feasible when one pattern is a subset of another.

Specifically, we demonstrate the following.

- Redundancy elimination is useful, but often not enough to reduce the number of messages. Reducing message count is crucial for our target architectures, especially for synchronous and collective communication.
- The traditional mechanisms of redundancy elimination can sometimes *prevent* the compiler from generating the best communication code.
- The well-known redundancy elimination technique of earliest communication placement is sensitive to minor syntactic differences in the high-level source, and may

```
Timestep loop:

glast(:,:) = g(1,:.:)
for i = 2 to nx - 1

\cdots = g(i,:,:)\uparrow \leftarrow \downarrow \rightarrow

\cdots = sum(g(i,ny,:)), sum(g(i,ny-1,:)), sum(g(i,0,:)), sum(g(i,1,:))

\cdots = glast(:,:)\uparrow \leftarrow \downarrow \rightarrow

\cdots = sum(glast(:,:)\uparrow \leftarrow \downarrow \rightarrow

\cdots = sum(glast(ny,:)), sum(glast(ny-1,:)), sum(glast(0,:)), sum(glast(1,:))

glast(:,:) = g(i,:,:)

g(i,:,:) = \cdots
```

Figure 2.1: A simplified form of the NPAC gravity code illustrating the need for combining communication.

Source code	Earliest placement	Combined placement
Loop:	Loop:	Loop:
	$\underline{\texttt{COMM}} \rightarrow \texttt{p}, \texttt{v}  \uparrow \texttt{p}, \texttt{u}  \leftarrow \texttt{u}  \downarrow \texttt{v}$	$\underline{\texttt{COMM}} \rightarrow \texttt{p}, \texttt{v}  \uparrow \texttt{p}, \texttt{u}  \leftarrow \texttt{u}  \downarrow \texttt{v}$
${ m cu} = { m p}  ightarrow$	${\tt cu} = {\tt p} { ightarrow}$	$\texttt{cu}  = \texttt{p} {\rightarrow}$
	<u>COMM ←cu ↑cu</u>	
$cv = p\uparrow$	$c \mathbf{v} = \mathbf{p} \uparrow$	$cv = p\uparrow$
	<u>COMM →cv ↓cv</u>	
$z = u\uparrow, v \rightarrow, p \rightarrow, p\uparrow, p\nearrow$	$\mathbf{z} = \mathbf{u} \uparrow,  \mathbf{v}  ightarrow,  \mathbf{p}  ightarrow,  \mathbf{p} \uparrow,  \mathbf{p} \nearrow$	$z = u\uparrow, v\rightarrow, p\rightarrow, p\uparrow, p\nearrow$
	$\underline{\texttt{COMM}} \downarrow_{\textbf{Z}} \leftarrow \underline{\textbf{z}}$	
h = $\mathbf{u} \leftarrow, \mathbf{v} \downarrow$	$\mathbf{h} = \mathbf{u} \leftarrow, \mathbf{v} \downarrow$	$\mathbf{h} = \mathbf{u} \leftarrow, \mathbf{v} \downarrow$
	$\underline{\texttt{COMM}} \rightarrow \underline{\mathbf{h}}  \uparrow \underline{\mathbf{h}}$	$\underline{\texttt{COMM}} \rightarrow \texttt{cv}, \texttt{h}  {\downarrow} \texttt{z}, \texttt{cv}  {\leftarrow} \texttt{z}, \texttt{cu}  {\uparrow} \texttt{cu}, \texttt{h}$
$unew = z \downarrow, h \rightarrow, cv \rightarrow, cv \downarrow, cv \searrow$	$unew = z\downarrow, h \rightarrow, cv \rightarrow, cv\downarrow, cv \searrow$	$\texttt{unew} = \texttt{z} {\downarrow},  \texttt{h} {\rightarrow},  \texttt{c} \texttt{v} {\rightarrow},  \texttt{c} \texttt{v} {\downarrow},  \texttt{c} \texttt{v} {\searrow}$
$\texttt{vnew} = \texttt{z}\leftarrow, \texttt{h}\uparrow, \texttt{cu}\leftarrow, \texttt{cu}\uparrow, \texttt{cu}\checkmark$	$\mathtt{vnew} = \mathtt{z} \leftarrow,  \mathtt{h} \uparrow,  \mathtt{cu} \leftarrow,  \mathtt{cu} \uparrow,  \mathtt{cu} \land$	$\mathtt{vnew} = \mathtt{z} \leftarrow, \mathtt{h} \uparrow, \mathtt{cu} \leftarrow, \mathtt{cu} \uparrow, \mathtt{cu} \checkmark$
$\mathtt{pnew} = \mathtt{cu} \leftarrow,  \mathtt{cv} \downarrow$	$pnew = cu \leftarrow, cv \downarrow$	$pnew = cu \leftarrow, cv \downarrow$

Figure 2.2: The NCAR shallow benchmark illustrating that redundancy elimination via earliest placement may lose valuable opportunity for message combining.

F90 Source code	Scalarized code	Hand coded F77
distribute $a, b, c ::$ (BLOCK) a = 3 b = 4 c(2:n) = a(1:n-1) + b(1:n-1)	$do \ i = 1 : n$ $a(i) = 3$ $COMM \ Earliest(a)$ $do \ i = 1 : n$ $b(i) = 4$ $COMM \ Earliest(b)$ $do \ i = 2 : n$	$do \ i = 1 : n$ $a(i) = 3$ $b(i) = 4$ $COMM \ Earliest(a), Earliest(b)$ $do \ i = 2 : n$ $c(i) = a(i - 1) + b(i - 1)$
	c(i) = a(i-1) + b(i-1)	C(i) = G(i - 1) + O(i - 1)

Figure 2.3: Syntax sensitivity of earliest placement. The right hand expressions are constants for simplicity only; for this particular code constant propagation will lead to communication-free code.

produce suboptimal code.

### 2.2.1 Beyond redundancy elimination

Redundancy elimination seeks to avoid unnecessary repetitions of communication for the same data. Programs often exhibit similar communication patterns involving different data as well. At least two types of additional communication optimization are possible. To see this, regard each (bulk-synchronization) communication as a  $P \times P$  matrix, where P is the number of physical processors, and element (i, j) denotes the volume of data that processor i sends to processor j.

- If, in two distinct communications, processor 1 sends to processor 2, these messages can be combined, if permitted by data dependency analysis, to amortize the fixed overhead for transferring a message.
- If, in one communication, processor 1 sends to processor 2, and in the other, processor 5 sends to processor 9, and data dependency permits these calls to be made by the respective processor pair at the same point in the code, these transfers can be parallelized.

It is quite amazing that in spite of the intense research in the area of communication optimization, these optimizations are often missed (or not even attempted) by most compilers.

Figure 2.1 shows a simplified form of a code called gravity. In this code, the 2-d arrays are of dimension (ny,nz) distributed (BLOCK, BLOCK)<sup>1</sup> and the 3-d arrays are of dimension (nx,ny,nz) distributed (\*,BLOCK,BLOCK)<sup>2</sup> In this code it is easy to detect that the nearest neighbor communication for g and glast can be combined, as can the sum operations. Thus, we can combine the eight nearest neighbor messages into four and the eight global sums into two parallel sets of four global sums.

#### 2.2.2 Earliest placement may hurt

In the last example, earliest placement did not preclude other optimizations; we merely needed a message combining post-pass. But in general, the situation could be more

<sup>&</sup>lt;sup>1</sup>Assuming for simplicity that P divides n, an array a[n] is block distributed over P processors by assigning  $\{a[in/P], \ldots, a[(i+1)n/P - 1]\}$  to processor  $i, 0 \le i < P$ . This extends naturally to multiple dimensions by treating the processors as a grid of the same number of dimensions.

<sup>&</sup>lt;sup>2</sup>The notation \* means that for all *i*, element a[i, j, k] are on the same processor.

complicated. In particular, redundancy elimination via earliest placement can prevent the combining possibilities from being exploited. To demonstrate this, we study the benchmark called shallow.

A simplified form of the code is shown in Figure 2.2. Notice that each statement in the array syntax expands into a two-deep loop-nest. If no redundant message elimination is done across different loop nests, there would be 18 exchanges generated per processor. (The IBM compiler already optimizes diagonal communication like  $p \nearrow$  by decomposing it into  $p\uparrow$  and  $p\rightarrow$ . This is reflected by the message counts.) Earliest placement will move up a communication as far as possible within the loop, communicating data right after definition. After a combining post-pass this will result in 12 array sections being communicated per processor. In contrast, if minimizing the number of messages is the objective, then we can obtain a schedule with only 8 exchanges per processor, in which placement of communication is not at the earliest point detected by dataflow analysis.

### 2.2.3 Syntax sensitivity

Since earliest placement pushes communication close to the production of the data value, placement is sensitive to the structure of intervals containing the production. As a case in point, consider the semantically equivalent codes in the three columns of Figure 2.3. Suppose arrays a and b have identical layout, say blocked. Using earliest placement, the messages for the two arrays can be combined in the third column whereas they cannot in the second code. Even if the programmer were to write the code in the first column, intermediate passes of compilation may destroy the loop structure by fission. In fact, the IBM HPF scalarizer [68] will translate the F90-style source to the scalarized form in the second column. If loop fusion can be performed before this analysis, as in this case, the problem can be avoided. But this is not always possible [120, § 9.2]. Thus, limited communication analysis at a single loop-nest level or a rigid placement policy may not work well. Our framework, by not relying on any restricted placement (like earliest or latest) but evaluating many choices globally, proves to be a much more robust strategy that is not easily perturbed by minor syntactic differences.

### 2.3 Network performance

By profiling our target networks, we justify the need for global message scheduling and identify simplifying assumptions that can be made about the optimization problem. We pick two platforms: the IBM SP2 with a custom network, and a network of Sparc workstations (NOW) connected by a commodity network (Myrinet). The SP2 uses IBM's message passing library MPL; the NOW uses MPICH, a portable implementation of the MPI standard from Argonne National Labs. Details of the networks can be found in [109, 106, 79]. We want to measure the benefits of large messages, while estimating the local block copy (bcopy) cost to collect many small messages into a large one. Figure 2.4 shows the profiling code and results.

We will be mostly interested in effective bandwidth since the HPF runtime does bulk transfer as the common case. For fine-grained communication, there is a distinction between CPU overhead per message, which cannot be overlapped with computation, and network latency, which can. In the bulk-synchronous execution model that HPF compiles down to, the benefits of overlap are very small. Kennedy *et al* estimate this to be typically 5-9% for 2-d stencil problems of size  $256 \times 256$  on a 16-processor iPSC/860, in the context of the Fortran-D execution model and runtime system [81].

The top curve shows the bandwidth of local block copy (bcopy) as a function of buffer size. The bottom curve plots network bandwidth as a function of message length, based on the time that the receiver waits for completion.

The top curve shows that as long as the message buffers fit in cache, we can ignore the overhead of **bcopy**. Fortunately, for both machines, most of the message startup amortization benefits occur at message sizes much smaller than the cache limit; given typical cache sizes, we believe this is a fairly general feature.

On the other hand, for messages much larger than cache, it may be important to suppress combining communication from non-contiguous array sections. E.g., for the SP2, the bandwidth of copying buffers larger than the cache is barely twice the message-passing bandwidth beyond cache size.

If there is a network co-processor or DMA, it is possible for the sender to quickly inject the message and the receiver to retrieve it more slowly. The middle curve shows bandwidth computed using the time the sender takes to inject the message. While the injection bandwidth is much lower than **bcopy**, it is larger than receive bandwidth for

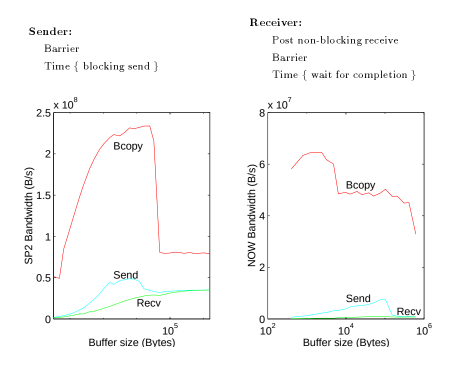


Figure 2.4: Buffer copying and network bandwidth studies on the IBM SP2 using MPL and the Berkeley NOW using MPICH. The x-axis is to a log scale.

certain message sizes. Injection bandwidth also declines around the cache limit.

Given our execution model and typical machine characteristics, we consider message aggregation and parallelization as the first-order concerns, and overlap as a second-order concern [81]. The latter also depends on the co-processor and network software. E.g., the implementors of MPL minimize co-processor assistance because the i860 coprocessor is much slower than the RS 6000 CPU, and the channel between the CPU and the co-processor is slow [106]. However, our algorithm permits additional techniques like Give-n-Take to be used to overlap latency with computation at the sender [114].

### 2.4 Compiler algorithms

In this section we describe our algorithm for placing communication code. This analysis is done after the compiler has performed transformations like loop distribution and loop interchange to increase opportunities for moving communication outside loops [68]. Wolfe provides an excellent overview of the compiler terminology we use [120]. The steps of our algorithm are described below and shown in Figure 2.5.

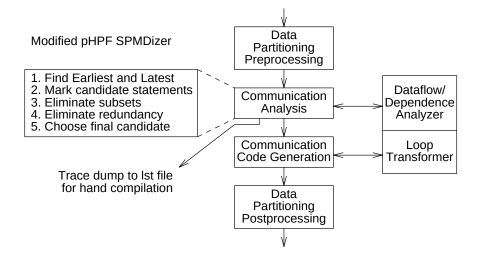


Figure 2.5: Prototype modifications to the IBM pHPF SPMDizer.

- 1. For each array expression on the right hand side of a statement that may need communication, identify the earliest (§2.4.3) and latest (§2.4.2) safe position to place that communication. One of our key innovations is to exploit the static single assignment (SSA) information [39, 35] already computed in an earlier phase by pHPF, refined by array dependence-testing [121]. In contrast, previous proposals for such analysis typically use a bidirectional dataflow approach with array section descriptors and/or bit-vectors [69].
- 2. For each non-local reference, identify a set of candidate positions, any one of which can be potentially chosen as the final point to emit a call to a message-passing runtime routine (§2.4.4).
- 3. Perform the "array-section" analog of common subexpression elimination: detect and eliminate subsumed communication (§2.4.6).
- 4. For the remaining communication, choose one from the set of candidate placements. In the prototype implementation we do this in two substeps that will be explained later (§2.4.5 and §2.4.7).

The above algorithms have been added to a prototype version of the pHPF compiler as shown in Figure 2.5. Throughout this section, we will use the code in Figure 2.6 as a

		CommSet(S) after									
S	Source code	Candidate	Subset	Redundancy	Earliest						
		marking	elimination	elimination	placement						
	distribute a, b, c, d :: (BLOCK,*)										
1	b(:, 1:n:2) = 1	$b_1$			$b_1$						
2	b(:, 2: n: 2) = 2	$b_{1}, b_{2}$			$b_2$						
3	if ( cond )										
4	a = 3				a 2						
5	else										
6	a = 4				a 2						
7	endif	$a_1, a_2, b_1, b_2$	$a_1, a_2, b_1, b_2$	$\{a_2, b_2\}$							
8	do $i = 2: n$										
9	do $j = 1 : n : 2$										
10	$c(i, j) = a_1(i - 1, j) + b_1(i - 1, j)$										
11	do $j = 1: n$										
12	$d(i,j) = a_2(i-1,j) - b_2(i-1,j)$										

Figure 2.6: Running example for analysis and optimization steps. Different uses of a and b are subscripted to distinguish their communication entries. Code for each communication entry is executed *after* executing the statement. The notation  $\{a_2, b_2\}$  means the messages for these accesses can be combined. The results of traditional earliest placement is shown in the last column for comparison.

running example to illustrate the operation of the steps of the algorithm.

#### 2.4.1 Representation and notation

We represent the program using the *augmented* control flow graph (CFG), which makes loop structure more explicit than the standard CFG by placing *preheader* and *postexit* nodes [3, 100] before and after loops. These extra nodes also provide convenient locations for summarizing dataflow information for the loop.

The CFG is a directed graph where each *node* is a basic block, a sequence of statements without jumps. Execution starts at the ENTRY node. A statement S may have a use u or definition d (abbreviated to "def") of an array variable. There are two kinds of defs. A "regular" def is one corresponding to the left hand side variable in a source statement. Conversion to single static assignment (SSA) form also introduces other defs called " $\phi$ -defs." These look like " $v_i = \phi(\ldots, v_j, \ldots)$ ," where each variable<sup>3</sup> v is renamed to  $v_1, v_2$ , etc., and there is only one assignment to a particular renamed version. A  $\phi$ -def  $d = \phi(\ldots, r, \ldots)$  is said to have a set of *parameters*  $\{r\}$ . All regular array defs are considered *preserving*, meaning that (unless proved otherwise) the original value is not assumed to be *killed*. See Cytron *et al* for a detailed treatment of SSA algorithms [39, 35]. We refer interchangeably

<sup>&</sup>lt;sup>3</sup>Arrays are regarded as scalars and the index information is ignored during SSA analysis.

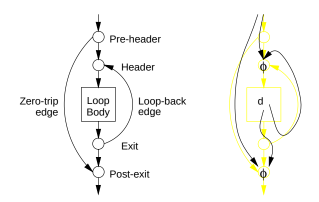


Figure 2.7: The diagram on the left hand side shows an example of a portion of the augmented control flow graph. It is reproduced (faint lines) on the right, and on top of it the SSA structure is super-imposed.

to a use, def, statement, or the node containing them. The node containing S is called CfgNode(S). When we say communication is placed at d we mean immediately after d.

A path  $\pi : v_0 \xrightarrow{+} v_j$  from  $v_0$  to  $v_j$  is a non-empty node sequence  $(v_i)$  with edges  $(v_{i-1}, v_i), 1 \leq i \leq j$ ; we also call  $\pi$  a backward path or backpath from  $v_j$  to  $v_0$ . Possibly empty paths are denoted  $v_0 \xrightarrow{*} v_j$ .  $\pi$  bypasses v if v does not occur on  $\pi$ . Two paths are non-overlapping if they are node-disjoint. Non-empty paths  $\pi_1 : v_0 \xrightarrow{+} v_j, \pi_2 : w_0 \xrightarrow{+} w_k$  are said to converge at z if  $v_0 \neq w_0, v_j = z = w_k$ , and  $(v_p = w_q) \Rightarrow (p = j \lor q = k)$ .

Loops are named L. Every loop has a well-defined *nesting level* called NL(L): this is the number of loops containing it. NL(v) for node v is defined likewise. L or v is *deep* or *shallow* according as NL is large or small. The common nesting level CNL(u,v) of two nodes u and v is the NL of the deepest loop containing them both. Every loop L has a single *preheader* node, PreHdr(L), and there is an edge from PreHdr(L) to Hdr(L), the *header* node. PreHdr(L) dominates all nodes in L. There is a *postexit* node for each distinct loop exit target. Each postexit node of L has an incoming edge, called *zero-trip* edge, from PreHdr(L) (along with the original loop-exiting edges). See Figure 2.7.

L has a  $\phi$ -def at Hdr(L), called  $\phi_{Hdr}$ , for each variable defined in the loop or in a loop transitively nested in L.  $\phi_{Hdr}$  has two parameters,  $r_{pre}$  and  $r_{post}$ , such that there exists a backpath from  $r_{pre}$  to ENTRY that bypasses all nodes in the loop, and there exists a path from any node in the loop to  $r_{post}$  which never takes an exit edge out of the loop.

The postexit node of each loop L has a  $\phi$ -def, called  $\phi_{\mathsf{Exit}}$ , for each variable defined

in the loop or in a loop transitively nested in it. Because of  $\phi_{\mathsf{Exit}}$ , a definition d can reach a use u only through a definition d' at a level  $\mathsf{CNL}(d, u)$ . d' can possibly be d only if  $\mathsf{CNL}(d, u) = \mathsf{NL}(d)$ ; otherwise, d' is a  $\phi$ -def at a level  $\mathsf{CNL}(d, u)$ .

### 2.4.2 Identifying the latest position

We describe how the compiler finds Latest(u), the latest point to place communication for u, which is also placed in as shallow a loop level as possible, so that messages are maximally vectorized. This follows from standard communication analysis: communication is placed just before the outermost loop in which there is no true dependence on u, and is placed just before the statement containing u if no such loop exists [123, 72, 68].

Given a use u, let d range over the reaching regular defs of u. (Reaching defs are defined in Cytron *et al* [39, 35].) Consider some d. Observe that it is never necessary to place communication for u deeper than at CNL(d, u). Given d and u, we can compute all possible direction vectors (each is a CNL(d, u)-dimensional vector) [120]. These vectors are used in the routine lsArrayDep, shown in Figure 2.8(d) on page 21. Let  $DepLevel(d, u) = max\{\ell : lsArrayDep(d, u, \ell)\}$ , the deepest level at which there is a loop-carried dependency between u and d.

Because of the dependency at level  $\mathsf{DepLevel}(d, u)$ , communication for u cannot be moved outside loop level  $\mathsf{DepLevel}(d, u)$ . The overall communication level for use u, denoted  $\mathsf{CommLevel}(u)$ , is set to  $\max_d \{\mathsf{DepLevel}(d, u)\}$ . Finally, to place communication, we check  $\mathsf{CommLevel}(u)$ : if  $\mathsf{CommLevel}(u) = \mathsf{NL}(u)$ , communication is placed immediately before the statement containing  $u^4$ ; if  $\mathsf{CommLevel}(u) < \mathsf{NL}(u)$ , communication is placed in the loop preheader of the loop at level ( $\mathsf{CommLevel}(u) + 1$ ) that contains u. Note that  $\mathsf{CommLevel}(u) > \mathsf{NL}(u)$  is not possible, and that by construction  $\mathsf{Latest}(u)$  dominates u.

### 2.4.3 Identifying the earliest position

We now describe how to compute  $\mathsf{Earliest}(u)$  for use u. Typically, dataflow analysis with array sections marks a set of nodes as "earliest", such that a copy of the communication code has to be placed at *all* these points. This is acceptable if each array section is communicated using a separate call to the communication library, but for our purposes, it greatly complicates code generation. In different control flow paths, communication for u

<sup>&</sup>lt;sup>4</sup>In this case no vectorization has been possible.

```
a. Earliest(u)
      For each def d of use u in depth-first preorder traversal:
         If \mathsf{Test}(d, u) then return d.
b. \mathsf{Test}(d, u)
      If d is a \phi-def, say d = \phi(\ldots, r_i, \ldots)
         For each \phi-parameter r_i
             \operatorname{visit}[\cdot] = 0, \operatorname{visit}[d] = 1
             Let c_i = \mathsf{Rcount}(\mathsf{Reaching}(r_i), u, \mathsf{CNL}(d, u), \text{visit})
         If two or more c_i's are positive
             Return TRUE
      else (d is a regular def)
         If lsArrayDep(d, u, CNL(d, u))
             Return TRUE.
c. \mathsf{Rcount}(d, u, l, visit)
      If d is a \phi-def, say d = \phi(\ldots, r_i, \ldots)
         If visit[d] return 0
         visit[d] = 1
         Return \sum_{i} \mathsf{Rcount}(\mathsf{Reaching}(r_i), u, l, \text{visit})
      else (d \text{ is a regular def})
         If lsArrayDep(d, u, l)
             Return 1
         else if d is a preserving def
             Return \mathsf{Rcount}(\mathsf{Reaching}(d), u, l, \text{visit})
         else return 0.
d. \mathsf{lsArrayDep}(d, u, \ell)
      If d is the pseudo-def at ENTRY then return TRUE
      If \ell > \mathsf{CNL}(d, u) then return FALSE
      If \exists direction vector \vec{v} = (v_1, \dots, v_{\mathsf{CNL}(d,u)}) such that
         • v_i = 0, for i \in \{1, ..., \ell - 1\}, and
         • v_\ell \ge 0
      then return TRUE
      else return FALSE
```

Figure 2.8: (a) Pseudocode for iterating over reaching defs of u. (b) Pseudocode for testing a def d to identify if d is the earliest communication placement point. (c) Pseudocode for recursively counting the number of incoming edges at  $\phi$ -defs or preserving regular defs that bear possible dependences. (In our SSA implementation, there is a pseudo-def at ENTRY for each variable accessed in the routine, which simplifies dataflow analyses.) (d) Routine to check array dependencies at the leaf defs.

may be combined with different references, making it impossible to generate a single version of the original computation containing u. The resulting code expansion can be enormous. Moreover, each message will need to have a descriptor for their contents, because, depending on the control path taken at runtime, the arrays packaged into a message may be different.

Therefore, we restrict our search to the *single* earliest position that dominates the use. Our experience with benchmarks, albeit limited, suggests that further sophistication is often unnecessary. The pseudocode for computing  $\mathsf{Earliest}(u)$  for a use u is shown in Figure 2.8(a-d). Earliest goes through defs reaching u in a certain order, Testing each in turn and returning with the first success. Test calls Rcount to find out if communication can be pushed beyond the candidate def d, or that the def represents a merge of at least two values and is hence a *critical* point to place communication. Rcount recurses through reaching defs of d, calling lsArrayDep at the leaves of the recursion, which are regular defs.

**Claim 2.4.1** Earliest(u) returns the earliest single dominating communication point  $d_1$  for use u.

In Figure 2.6,  $\mathsf{Earliest}(a_1) = \mathsf{Earliest}(a_2) = 7$ . Traditional array dataflow analysis, which does not insist on dominating defs [69], would lead to  $\mathsf{Earliest}'(a_1) = \mathsf{Earliest}'(a_2) = \{4, 6\}$ . In both cases,  $a_2$  subsumes  $a_1$ . We prove Claim 2.4.1 using the following three lemmas.

Lemma 2.4.2  $d_1$  dominates u.

**Proof.** (By contradiction.) We assume  $d_1$  is not the pseudo-def at ENTRY, since the latter dominates all nodes in the CFG. Let  $\ell_1 = \mathsf{NL}(d_1)$ , and  $L_1$  be the loop containing  $d_1$ . Note that  $\ell_1 \leq \mathsf{NL}(u)$  because Earliest will never flag a  $d_1$  with  $\mathsf{NL}(d_1) > \mathsf{NL}(u)$ . Assume  $d_1$  does not dominate u. Then there exist two or more paths: one from ENTRY to u that bypasses  $d_1$ , and another from  $d_1$  to u. If  $\mathsf{NL}(u) = \mathsf{NL}(d_1)$ , these two paths imply that there exists a  $\phi$ -def at level  $\ell_1$  with (at least) two parameters,  $r_1$  and  $r_2$ , such that there exist two non-overlapping backpaths: one from  $r_1$  to  $d_1$ , and the other from  $r_2$  to the pseudo-def at ENTRY that bypasses  $d_1$ . (Because of the zero-trip edges, we can ignore other loops nested in  $L_1$ .). That there is such a  $\phi$ -def at level  $\ell_1$  still holds if  $\mathsf{NL}(u) > \mathsf{NL}(d_1)$ , because the preheader node of each loop containing u dominates u, and the two (or more) paths converge at the preheader node which is at level  $\ell_1$ , at the latest. Test is called on at least one of these  $\phi$ -defs, say p, before  $d_1$  during the traversal of Earliest(u), starting from u. During execution of Test(p, u), Rcount gets called on defs  $\text{Reaching}(r_1)$  and  $\text{Reaching}(r_2)$ , with nesting level  $\text{CNL}(p, u) = \ell_1$ . The call at  $\text{Reaching}(r_1)$  returns a positive number, because some recursive call inspects  $d_1$ . Similarly the call at  $\text{Reaching}(r_2)$  also returns a positive number, because some recursive call inspects ENTRY. Since at least two invocations of Rcount return a positive numbers, the  $\phi$ -def, not  $d_1$ , will be returned as Earliest(u) if  $d_1$ does not dominate u, a contradiction.

**Lemma 2.4.3** Let  $n_3$  be any proper dominator-tree ancestor of  $d_1$ . Then there exists a regular def  $d_2$  such that  $lsArrayDep(d_2, u, CNL(d_1, u))$  returns TRUE, and there also exists a path  $d_2 \xrightarrow{*} d_1 \xrightarrow{+} u$  that bypasses  $n_3$ .

**Proof.** If  $d_1$  is the pseudo-def at ENTRY, there is nothing to prove. Also, if  $d_1$  is a regular def,  $lsArrayDep(d_1, u, CNL(d_1, u))$  must hold for  $d_1$  to be returned as Earliest(u), in which case  $d_1$  serves as the definition  $d_2$  in the statement of the lemma. Therefore, we can assume  $d_1$  is a  $\phi$ -def.

By design,  $\text{Test}(d_1)$  returned TRUE because at least two Rcount calls on the  $\phi$ -parameters of  $d_1$  returned positive counts. But because of the visit[] array, no def is accounted more than once. Therefore the two positive counts can be attributed to two node-disjoint backpaths to two distinct regular defs (one of which could be ENTRY). At most one of these paths contain  $n_3$ . Let  $d_2$  be some regular def on the other path such that  $lsArrayDep(d_2, u, CNL(d_1, u)) = TRUE$ . Then there is a  $d_2 \xrightarrow{+} u$  path bypassing  $n_3$ .

**Lemma 2.4.4** There is no regular def  $d_4$  along a path  $d_1 \xrightarrow{+} d_4 \xrightarrow{+} u$  such that  $\mathsf{lsArrayDep}(d_4, u, \mathsf{CNL}(d_4, u))$  returns TRUE, and there is a path from  $d_4$  to u that bypasses  $d_1$ . That is, it suffices to place communication at  $d_1$ .

**Proof.** (By contradiction.) Assume there exists such a  $d_4$ . According to SSA construction, two cases can occur: either (1)  $d_4$ , as well as  $d_1$ , dominates u, or (2)  $d_4$  has a path, bypassing  $d_1$ , from it to u through one or more  $\phi$ -defs that dominate u.

**Case 1.** If  $d_4$  dominates u,  $d_4$  cannot also dominate  $d_1$ . Otherwise, there exists a path from ENTRY to  $d_4$  to u that bypasses  $d_1$  (second condition in the lemma), in which case  $d_1$ cannot dominate u, contradicting Lemma 2.4.2. Therefore,  $d_1$  dominates  $d_4$  (note that if both  $d_4$  and  $d_1$  dominate u, one of them must dominate the other), which in turn dominates u. Thus  $\text{Test}(d_4, u)$  is called before  $\text{Test}(d_1, u)$  by Earliest(u).  $\text{Test}(d_4, u) = \text{TRUE}$  because  $\text{IsArrayDep}(d_4, u, \text{CNL}(d_4, u)) = \text{TRUE}$ , so  $d_4$  will get returned as Earliest(u); a contradiction. **Case 2.** In the second case,  $d_1$  dominates the  $\phi$ -defs. If not, then  $d_1$  would not dominate u either, (contrary to Lemma 2.4.2) because there is a path  $d_4 \xrightarrow{+} \phi \xrightarrow{+} u$  avoiding  $d_1$ . Hence, these  $\phi$ -defs are dominated by  $d_1$  and are visited before  $d_1$  by  $\mathsf{Earliest}(u)$ . It follows, from a similar argument in the proof of Lemma 2.4.2, that these two paths converge at some node at level  $\mathsf{CNL}(d_4, u)$ , creating a  $\phi$ -def at level  $\mathsf{CNL}(d_4, u)$ . This  $\phi$ -node has (at least) two parameters,  $r_1$  and  $r_2$ , such that there exist two non-overlapping paths: one from  $d_4$  to  $r_1$ , and the other from  $d_1$  to  $r_2$ . When applied to  $r_1$ , Rcount returns positive, possibly because of  $d_4$ , which satisfies  $\mathsf{IsArrayDep}(d_4, u, \mathsf{CNL}(d_4, u))$ . When applied to  $r_2$ , Rcount returns positive, possibly because of the pseudo-def at  $\mathsf{ENTRY}$ . Since (at least) two parameters return positive, the  $\phi$ -def, not  $d_1$ , is returned by  $\mathsf{Earliest}(u)$ , another contradiction.

**Proof of Claim 2.4.1.** Observe that only a node that dominates u can serve as a single communication point for u. Lemma 2.4.2 says that  $d_1 = \mathsf{Earliest}(u)$  dominates u. Consider all dominator-tree ancestors of u. From this set, Lemma 2.4.3 rules out all nodes that strictly dominate  $d_1$  as unsafe. Finally, Lemma 2.4.4 implies that  $d_1$  is a safe communication point for u.

### 2.4.4 Generating candidate positions

Since any safe position to insert a single copy of communication for use u must dominate u, the set of candidate positions has a very simple characterization in terms of the following claims. We omit the proofs; see Figure 2.10 for the justification.

**Claim 2.4.5** Starting at the basic block containing Latest(u), denoted c(Latest(u)), if we follow parent links in the dominator tree of the CFG, we will reach the basic block containing Earliest(u).

**Claim 2.4.6** The statements marked in the basic blocks encountered during the dominator tree traversal from c(Latest(u)) up to c(Earliest(u)) are exactly those that are single candidate positions for communication placement for use u.

Our algorithm for finding candidate placements of communication is thus extremely simple, and shown in Figure 2.9(e). In our example (Figure 2.6), statements 3, 4, 5, and 6 are not candidates for placing communication for accesses  $b_1$  and  $b_2$  because they do not dominate those uses.

e. Mark candidates: c = CfgNode(Latest(u)) While $c \neq CfgNode(Earliest(u))$ do Mark all statements up to Latest(u) in basic block $c$ c = DomTreeParent(c) Mark all statements between Earliest(u) and Latest(u) in CfgNode(Earliest(u)).
f. Eliminate redundancy:
Repeat until no progress: Find statement $S$ and $c_1, c_2 \in CommSet(S)$ such that $c_2$ subsumes $c_1$ For all $S'$ dominated by $S$ disable $c_1$ in $CommSet(S')$
g. GreedyChoose: Let StmtSet( $c$ ) = { $S : c \in CommSet(S)$ } Consider entries $c$ in increasing order of  StmtSet( $c$ ) : For each $S \in StmtSet(c)$ , count the number of entries in CommSet( $S$ ) with which $c$ can combine (see text) Pick $S$ with the highest count to place $c$ Delete $c$ from CommSet( $S'$ ) for all $S' \neq S$ Place each group of combined entries at the latest position common to the candidate placements of the entries it contains, including entries disabled during redundancy elimination.

Figure 2.9: Pseudocode for communication placement. (e) Pseudocode for marking all candidate statements for communication placement. (f) Pseudocode for global redundancy elimination.(g) Simple greedy heuristic to choose a final position from the set of candidates.

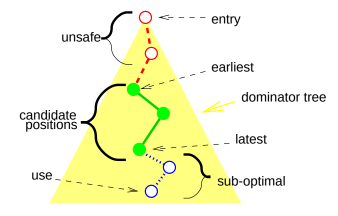


Figure 2.10: The candidate places for a fixed use are a set of consecutive nodes on the dominator chain from the use up to the entry point of the procedure.

#### 2.4.5 Subset elimination

Our current algorithm gives priority to reducing the volume and number of messages over exploiting overlap benefits or reducing contention for buffers and cache. Given this simplification, we can preclude a large number of candidate positions without compromising the quality of the solution. Specifically, let CommSet(S) denote all communication entries associated with the statement S. A given entry can occur in the CommSet of many statements. If for statements  $S_1$  and  $S_2$  we have  $CommSet(S_1) \subset CommSet(S_2)$ , we can reset  $CommSet(S_1) = \emptyset$  without losing opportunities for combining or redundancy elimination. For example, in Figure 2.6, the  $CommSet(S_2)$ , either set may be emptied at this stage, because the actual choice governing the placement of communication would be made in the final step (§2.4.7).

#### 2.4.6 Redundancy elimination

Typically, earlier approaches eliminated redundancy by examining the list of communications placed before each statement, and check each pair of entries to see if one **subsumes** the other. This test is based on the Available Section Descriptor (ASD) representation of communication [69]. Briefly, an ASD consists of a pair  $\langle D, M \rangle$ , where D represents the data (scalar variable or an array section) being communicated, and M is a mapping function that maps data to the processors which receive that data.

A communication  $\langle D_1, M_1 \rangle$  is made redundant by another communication  $\langle D_2, M_2 \rangle$  if  $D_1 \subseteq D_2$ , and  $M_1(D_1) \subseteq M_2(D_1)$ .

In our case, since there can be many entries for a reference, we have to propagate the redundancy information globally. The pseudocode for eliminating redundant communication in the context of our current framework is shown in Figure 2.9(f). The modification is that in each step examining a statement S, the subsumed communication entry is cleared not only from CommSet(S) but from all statements S' such that S dominates S'. (The dominance ordering prevents a cycle of deletions.) We iterate over statements and communication pairs until no more elimination occurs.

**Claim 2.4.7** The subset and redundancy elimination steps are safe, i.e., the remaining communication entries are sufficient.

**Proof.** Subset elimination in an arbitrary order is clearly safe because  $\subseteq$  is transitive. The only concern is that when we throw away all copies of an entry because it was found subsumed at one place, its copy may earlier have subsumed some other communication at another place.

Let entries  $e_1, e_2 \in \mathsf{CommSet}(S_1)$  and  $e_2, e_3 \in \mathsf{CommSet}(S_2)$ . Since  $e_2$  is in both, assume w.l.o.g. that  $S_1$  dominates  $S_2$ . We say that  $e_j$  is redundant if  $\mathsf{ASD}(e_j) \subseteq \mathsf{ASD}(e_i)$ , for some statement S, and  $e_i, e_j \in \mathsf{CommSet}(S)$ .

The possible mishap sequence is that we find  $ASD(e_3) \subseteq ASD(e_2)$  at  $S_2$ , delete  $e_3$ from all S, find  $ASD(e_2) \subseteq ASD(e_1)$  at  $S_1$ , and delete  $e_2$  from all S, including  $S_1$  and  $S_2$ . If the remaining entry  $e_1$  does not remain available at  $S_2$ , it must be due to a def d of some data in  $ASD(e_3)$  between  $S_1$  and  $S_2$ . But in that case, since  $d \in ASD(e_2)$ ,  $e_2$  could not have occurred at both  $S_1$  and  $S_2$  in the first place.

One implication of the above ordering of eliminations is noteworthy. Consider our running example (Figure 2.6), specifically the communication due to the uses  $b_1$ and  $b_2$  (ASD $(b_1) \subset$  ASD $(b_2)$ , since  $b_1$  represents b[2 : n, 1 : n : 2] and  $b_1$  represents b[2 : n, 1 : n]). Since Earliest $(b_1) = 1 \neq$  Earliest $(b_2) = 2$ , an initial test of redundancy based on earliest placement, followed by candidate marking and subset elimination will not catch the redundancy. Thus, by choosing a placement for  $b_1$  later than Earliest $(b_1)$ , we are able to eliminate the communication for  $b_1$  completely. In contrast, the solution proposed in [69] would move each communication to the earliest point, and reduce the communication for  $b_2$  to  $ASD(b_2) - ASD(b_1)$ , while the communication for  $b_1$  would remain unchanged. The solution obtained with our current method is superior because it reduces the communication startup overhead, and it makes code generation much simpler.

# 2.4.7 Choosing from the candidates

At this stage we can still have a communication entry c in multiple CommSet's, and we have to arbitrate in favor of one. The goal is to minimize the total communication cost. In the common message cost model using fixed overhead per message and bandwidth, minimizing the cost is  $\mathcal{NP}$ -hard (also see §2.6). In practice, simple greedy heuristics work quite well; see Figure 2.9(g). It is similar to Click's global code motion heuristic [36]: consider the most constrained communication entry next, and put it where it is compatible in communication pattern (as shown by the test below) with the largest number of other candidate communication. A more refined heuristic would use estimates of message sizes and consider the communication cost if the current entry were combined with a given set of entries.

The entries in the **CommSet** of each statement can now be partitioned into groups, each group consisting of one or more entries which will be combined into a single aggregate communication operation. Any flexibility still available in placing this aggregate can be used to push this communication later, if reducing contention for buffers and cache is more important than overlap benefits (as is folk truism for the SP2), or push it earlier if the situation were reversed. Our algorithm places communication for each group at the *latest* position common to the possible placements of each entry in that group (including positions disabled during the previous step for redundancy elimination). Deferring the placement decision until this final step enables our algorithm to take advantage of any possible placement that leads to redundancy elimination or combining benefits, without the drawback of unnecessary movement of communication that uses up more resources or degrades performance.

**Criteria for communication compatibility.** While in principle, code for arbitrary communications can be combined into code for a single communication operation, we are interested in combining messages only when the startup overheads associated with all but one of them can be eliminated, leading to improved performance. Hence, we view two communications as compatible for combining if the associated sender-receiver relationships

are identical or one is a subset of the other.

Thus, communications for  $\langle D_1, M_1 \rangle$  and  $\langle D_2, M_2 \rangle$  are combined only if  $M_1 = M_2$ or  $M_1 \subset M_2$ . The combined communication is given by  $\langle D_1 \cup D_2, M_2 \rangle$ . In order to ensure better performance and for simplicity of code generation, we impose the following additional constraints on combining.

- The combined data size of  $D_1 \cup D_2$  must be below a threshold (based on our study reported in §2.3, currently set to 20 KB for SP2), beyond which combining messages leads to diminishing returns or even worse performance. When data sizes are unknown, the compiler uses rules of thumb like assuming that nearest-neighbor communication (NNC) and reductions (where volume of data communicated is significantly lower than that involved in computation) are operating within the range suitable for combining.
- The size of D<sub>1</sub> ∪ D<sub>2</sub>, as approximated by a single section descriptor (array sections are not closed under the union operation), should not exceed the combined size of D<sub>1</sub> and D<sub>2</sub> by more than a small constant. This descriptor for D<sub>1</sub> ∪ D<sub>2</sub> refers to identical sections of different arrays if D<sub>1</sub> and D<sub>2</sub> correspond to different arrays, and to a single array otherwise.

The check for  $M_1 \subseteq M_2$  is done in the virtual processor space of template positions, as described in [69]. However, we have incorporated extensions to check for equality of mappings in physical processor space for nearest-neighbor communication and for mappings to a constant processor position [69].

#### 2.4.8 Code generation

As shown in Figure 2.5, the step after communication analysis and optimization is to insert communication code in the form of subroutine calls to the pHPF runtime library routines, which in turn invoke MPL/MPI. The runtime library provides a high-level interface through which the compiler specifies the data being communicated in the form of array sections, and the runtime system takes care of packing and unpacking of data. For NNC, data is received in overlap regions [123] surrounding the local portion of the arrays. For other kinds of communication involving arrays, data is received into a buffer that is allocated dynamically, and the array reference that led to this communication is replaced by a reference to the buffer.

Benchmark	shallow	gravity	gravity	$\operatorname{trimesh}$	trimesh	hydflo	hydflo
Routine	main	main	main	normdot	gauss	flux	hydro
Comm Type	NNC	NNC	SUM	NNC	NNC	NNC	NNC
Original (orig)	18	8	8	24	13	52	12
+ Redundancy elimination (nored)	12	8	8	24	13	30	12
+ Combined messages (comb)	8	4	2	4	4	6	6

Table 2.2: Reduction in compile-time message counts by using the new algorithm.

Redundant message elimination for NNC requires no further change to code generation. For other forms of communication, code generation has been modified to ensure that the array reference corresponding to eliminated communication is also replaced by a reference to the buffer holding non-local data, and that this buffer is deallocated only after its last use is over.

Combining messages for different arrays requires changes in code generation and the HPF runtime library routines. The data being sent or received is still represented by a single section descriptor, but now has a list of arrays associated with it. Correspondingly, the runtime routines now have to take on additional responsibilities of packing and unpacking data for the multiple array sections. Our benchmarks currently emit calls to a rudimentary runtime library with these features, but this has not been integrated into the compiler yet.

# 2.5 Performance

The algorithm described above has been implemented in the pHPF compiler. In order to study the potential performance benefits before the code generator and the run-time library could be modified to take advantage of the superior communication placement, we emitted scalarized code annotated with human readable communication entries after the analysis and optimization pass of the compiler. Table 2.2 shows some compile-time statistics of the reduction in the number of static call sites to the communication library. Static message counts are reduced by a factor of roughly 2–9.

The trace emitted was then used to generate C programs with calls to MPL/MPICH message passing libraries. This enabled us to study performance improvements not only on the IBM SP2, but also on a network of workstations (NOW) consisting of Sparc workstations connected by a Myrinet switch. For each benchmark, the compiler generated two or three versions of code. The baseline pulls communication into outermost possible loops but does not detect redundancy or perform message scheduling.

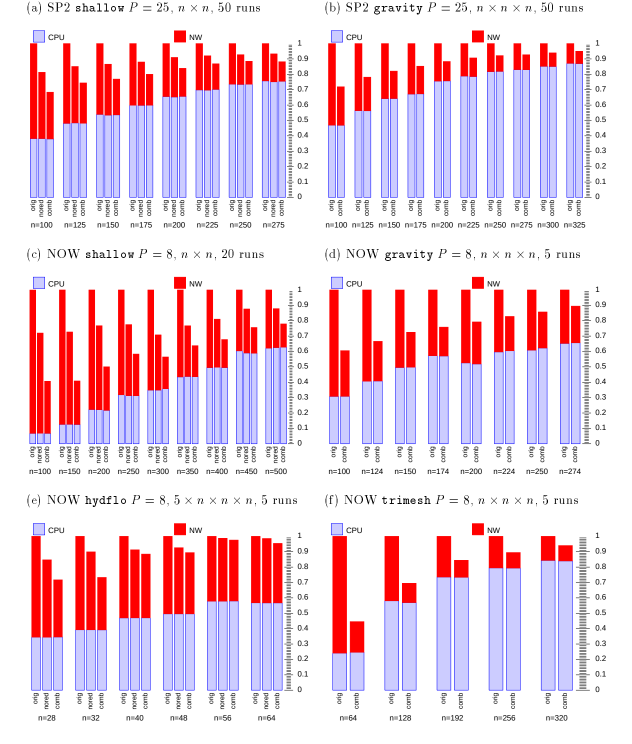


Figure 2.11: Performance benefits of the new algorithm: normalized running times.

The next version uses earliest placement for redundancy elimination but does not perform message scheduling or combining. The final version uses the new algorithm. (Note: gravity and trimesh have no redundancy.) On the SP2, all codes were compiled using the IBM xlc compiler. On the NOW, we used SUNWspro compiler cc. Optimization level -O3 was used.

We report the results in Figure 2.11. Each diagram includes the number of processors and the number of runs over which the median performance is reported. In each bar-chart the x-axis is the problem size. For each size two or three bars are plotted, one for each version of generated code. The y-axis is normalized so that the original code has unit running time, and the dark segment representing network cost shortens as optimizations are applied. These measurements were made with overlap disabled to clearly account for CPU and network activity. All floating point operations are on double words (eight bytes). shallow and trimesh involve 2-d  $n \times n$  arrays distributed (BLOCK, BLOCK); shallow has 13 and trimesh has over 25 such arrays. Thus the problem sizes are realistic in that they occupy several MBytes. Communication time is reduced by a factor of 2–3, which typically translates to 10-40% overall running time reduction. gravity uses a 3-d  $n \times n \times n$  array distributed (\*, BLOCK, BLOCK). Thus memory needed even at moderate *n* is quite staggering, and the graphs again show 10-40% overall gain in this reasonable size range. hydflo uses eight  $5 \times (n+2)^3$  arrays. Therefore even for small n, the memory requirement is enormous, which affects the size range shown. Finally, the SP2 network has lower overhead and higher bandwidth than the NOW<sup>5</sup>, which is evident from the higher overall performance gains on NOW compared to SP2, although the reduction in communication cost alone is roughly proportionate.

### 2.6 Extensions

The basic idea of exploiting flexibility in communication placement is rather general. Although for the sake of practicality our current prototype makes some simplifications; it would be interesting to extend the work in two ways. Currently, network architecture is undergoing considerable flux. If the CPU-network overlap can be exploited more effectively in future generation machines, the compiler could obtain better performance by considering the trade-offs between enhancing the overlap and reducing the number of

<sup>&</sup>lt;sup>5</sup>Note that we use MPI on both machines, not Active Messages [115].

messages and buffer contention. In particular, the simple subset elimination step (§2.4.5) would have to be dropped, as it could easily degrade the quality of the solution. In fact, the general problem becomes intractable when all of these conflicting optimizations are considered. Another future direction comprises enhancements for optimizing special communication patterns like reductions.

#### 2.6.1 General models

In a well-known model of communication cost, the problem of optimally selecting final candidates is  $\mathcal{NP}$ -hard, justifying our heuristic approach. A runtime call to the communication library in general leads to a many-to-many communication pattern. Let the inverse bandwidth of the network be scaled to one, and the message startup cost be Cin these units. The cost of this pattern to a given processor is C times the total number of distinct processors that it sends to or receives from, plus the total volume of data that it sends or receives. Ignoring CPU-network overlap in our bulk-synchronous model, the cost of a pattern is the maximum cost over all processors, and the cost of a set of patterns is the sum of their costs. The maximum number of bytes that can be buffered at any processor for aggregating, sending or receiving messages is B. Unfortunately, we can show the following.

Claim 2.6.1 Picking one candidate position for each reference, such that the total cost of all patterns is minimized, is  $\mathcal{NP}$ -hard.

**Proof.** We will give two reductions with slightly different models. We first outline the common features of the two models. Let there be P processors. A point-to-point message of length b bytes between any two processors costs each processor time C + b, where C is the startup cost scaled to units of per-byte transfer cost. Each array communication entry e gives rise to a problem similar to the h-relation problem. Our problem can be characterized by a  $P \times P$  matrix  $A^{(e)}$ , where  $A_{ij}^{(e)}$  gives the number of bytes to be sent from processor i to j. The time for executing e is given by:

$$t(e) = t(A^{(e)}) = \max_{j} \left\{ C \sum_{i} \left( \operatorname{sign}(A_{ij}^{(e)}) + \operatorname{sign}(A_{ji}^{(e)}) \right) + \sum_{i} \left( A_{ij}^{(e)} + A_{ji}^{(e)} \right) \right\}$$

**Bin packing.** The model for the first reduction assumes that any two patterns can be combined. Assuming any  $e_1$  and  $e_2$  combine, and ignoring buffer copy overhead, we can also assume  $t(e_1 \cup e_2) = t(A^{(e_1)} + A^{(e_2)})$ . But we do have to ensure that if a set of entries  $\{e\}$  are

combined, no processor buffers more than B bytes for sending or receiving. It is trivial to reduce bin-packing to this problem. Let there be communication only from processor 1 to 2; and let the number of bytes in the messages be the item sizes from the bin-packing problem, and let the buffer limit be the bin size. Minimizing the number of bins then corresponds to the minimum number of messages.

Set cover. In practice, code generation issues make it impossible to combine arbitrary entries  $e_1$  and  $e_2$ . The second model ignores the buffer constraint B and focuses on the communication pattern; specifically, assume that similar to §2.4.7, we combine entries  $e_1 = \langle D_1, M_1 \rangle$  and  $e_2 = \langle D_2, M_2 \rangle$  iff  $M_1 \subseteq M_2$  or  $M_2 \subseteq M_1$ .

Initially make the temporary assumption that sending message is free, and receiving message has some enormous startup cost. Under this model, the cost of an entry is the maximum number of messages any processor receives. If sending also had large startup cost, the cost of an entry would be the maximum number of messages any processor sends plus receives.

Given a set cover instance with elems  $\{e_1, \ldots, e_n\}$  and sets  $\{S_1, \ldots, S_m\}$ . Elems are assigned numbers  $1, \ldots, n$ . We will have n + m entries. Each entry is  $P \times P$  where P = n + 1. For each element j there is an entry  $c_j$  of the following form:  $c_j[0, j] = 1$ ; all other  $c_j[] = 0$ . For every set  $S_i$  there is also an entry:  $c_i[0, j] = 1$  iff  $j \in S_i$ . All other  $c_i[] = 0$ . The rules for combining are as above. We say two entries  $c_1$  and  $c_2$  are combinable iff elementwise  $c_1 \leq c_2$  or  $c_2 \leq c_1$ . If there is a set cover of size K, then there is a message schedule with K messages, and conversely. It is not only  $\mathcal{NP}$ -hard to solve set cover exactly, but there is a constant c such that it is not possible to get a polynomial time algorithm that returns a solution within a multiplicative factor of  $c \log n$  of optimum, unless  $\mathcal{P} = \mathcal{NP}$ .

The reason for the temporary assumption was that otherwise processor 0 would have to send many messages, while all other processors receive one message each in parallel for every entry. To remove the assumption, just stagger the first row of each matrix: instead of setting c[0, j] to 1, set c[j - 1, j] to 1. Now for each entry each processor has to send at most one message and receive at most one message. By blowing up P by a factor of 2, we can even make sure that no processor both sends and receives.

Like many other  $\mathcal{NP}$ -hard problems, the optimization problem can be formulated as an integer linear program (ILP). Furthermore, several additional constraints can be incorporated into the ILP, including overlap between CPU and network and message buffer and cache constraints. Profile information would be crucial to specify this ILP and solve it to adequate precision.

### 2.6.2 Special communication patterns

Throughout, we have focused on two-party communication that fetches remote data before a statement involving distributed arrays can be executed. Another important class of communication is collective communication such as scans and reductions, generated, e.g. from statements of the following form involving associative operators.

x[0] = y[0];	s = 0;
for ( i = 1; i < n; i++ )	for ( i = 0; i < n; i++ )
x[i] = x[i-1] + y[i];	s += x[i];

Global operations are dealt with in a special way in the compiler since communication requirement is, in a sense, inverted. Whereas ordinary statements require communication to fill in remote values before computation can proceed, for reduction the computation occurs first (for the partial reduction operation on individual processors), followed by communication for the global reduction operation that must be completed before the use. Our preliminary prototype does not schedule collective communication yet. For communications which are marked as reductions, we need to employ a reversed SSA analysis, i.e., iterating through reached uses of a given definition to determine the latest point at which communication may be safely placed. Conceptually this is identical to the framework in this chapter, but the implementation is left for future work. The current implementation does allow reduction communications placed at the same point to be combined, as in **gravity**.

# 2.7 Conclusion

We have presented an algorithm for global optimization of communication code placement in compilers for data-parallel languages like HPF. Modern parallel architectures greatly reward dealing with remote accesses throughout a program in an interdependent manner rather than naively generating messages for each of them. We achieve precisely this enabling optimization. In particular, we explore later placements of communication that preserve the benefits of redundancy elimination (normally obtained by moving communication earlier), reduce the wastage of resources like buffers for non-local data, and improve performance due to other factors like fewer messages. Preliminary performance results obtained on some HPF benchmarks show significant reduction in communication costs and overall improvements in performance of those programs on the IBM SP2 and a cluster of Sparcs connected by Myrinet. In the future, we will conduct performance studies to investigate the desirability of including partial redundancy elimination as well into our framework. Another area for future work is interprocedural analysis; we believe that the application of our algorithm across procedure boundaries can often lead to further improvements in performance.

# Chapter 3

# Scheduling hybrid parallelism

# 3.1 Introduction

The parallel executable code that compilers for data parallel languages generate usually runs in a bulk synchronous style. There are alternating phases of computation and cooperative communication across most or all processors. Fine-grain asynchronous communication is expensive on most current machines. Reducing this is an active area of parallel architecture research. However, it is also important for compilers and runtime systems to optimize an application to coarsen the grain size and make the communication more coordinated. The compiler algorithm outlined in the previous chapter enhances the bulk-synchrony of the program. Another way to reduce communication overhead is to reduce the number of processors, which makes sense only if the application has other work that can be done on the freed processors. In the context of data-parallel languages like High Performance Fortran (HPF), this is equivalent to applications that show task parallelism as well as the standard *data* parallelism of loops. There is a growing body of such applications, including eigensolvers, adaptive mesh codes, circuit simulation, and query trees in parallel relational databases. Another incentive for investigating such programming models is that future multiprocessors are likely to be clusters of symmetric shared memory (SMP) machines. The data and task parallelism will map efficiently to within and across SMP clusters, since loop-level communication in data parallelism can be supported by shared memory, while relatively infrequent task-level communication can be implemented via message passing across clusters.

Recent versions of the HPF proposal includes support for such applications [55].

Some parallelizing compilers have been extended to express such a *mixed* parallel execution model, but the algorithmic details of the runtime scheduler are still under debate. These applications are characterized by a precedence graph of tasks, each of which can run on many processors in a data parallel fashion. For applications amenable to compile-time scheduling, the task graph has to be known statically, and information about the communication and computation cost of all the task nodes is needed.

Unfortunately, the amount of static task parallelism in most problems is so small (typically 4–8) that except for the smallest problems, the gains over pure data parallelism are negligible. Static task graphs, such as those generated from control flow graphs by parallelizing compilers, have a fixed small degree of task parallelism. For example, the benchmarks in [95] have 4–7 fold effective task parallelism and the signal processing applications in [110] have a 2–5 fold task parallelism. The task parallelism in climate modeling applications is typically no more than 4–6.

Problems where the task parallelism grows with problem size, such as divide and conquer problems, are therefore much more promising from the perspective of exploiting mixed parallelism, and are the focus of this chapter. On the other hand, these applications may have to be scheduled dynamically, because the structure of the task graph and costs of tasks may not be known in advance. With runtime scheduling, the ability to use expensive optimization techniques, such as linear-programming, is limited. This chapter explores how to extend the purely data-parallel execution model of HPF and implement simple and effective dynamic scheduling algorithms. Specifically, we make the following contributions:

- We identify the nature of applications that need task parallelism in addition to data parallelism for enhanced performance. In particular, we focus on dynamic, scalable task parallelism such as in divide and conquer.
- For dynamic scenarios, we formulate and give practical solutions for scheduling such task graphs using a simple *switched* execution strategy: some tasks are allocated all processors, some only one; the system alternates between executing tasks of the first type one after the other, and packing tasks of the second type into the processors evenly.
- We prove that the algorithms are close to the best possible switched execution. We also show by experiments that the gains of switched parallelism over pure data-parallelism

can be dramatic, e.g., up to a factor of three times higher MFLOPS on the IBM SP2. (This  $3 \times$  factor is not to be compared with the  $2-7 \times$  task parallelism mentioned earlier; those amounts of statically available task parallelism do not translate into similar improvements over data parallelism.)

• Using simulation we estimate the gap between the efficiency of switched parallel execution with the best possible efficiency of *any* scheduler. The gap is small.

Data parallelism is the most familiar parallel execution model in scientific computing, as in ScaLAPACK, CMSSL, Fortran D, CM-Fortran, and High Performance Fortran [45, 76, 72, 49, 55]. The computational load can usually be balanced statically, and the important performance issue is generating parallel loops with minimal communication overhead (as in Chapter 2). In general, data parallel applications show cooperative computation and communication within loops, and the parallelism in an operation is exploited over the complete machine partition.

Task parallelism is specified at the language level using statements like "fork", which generate a new thread of control to execute the given function invocation. Tasks may depend on the completion of other tasks, so task parallelism is modeled by an acyclic graph (DAG) whose vertices represent tasks. Each task runs on exactly one processor (i.e., is sequential). The advantage is that communication is potentially required only in setting up the task and retrieving outputs. Since the task runs on one processor, no inter-processor communication occurs during task execution. (In our problem domain we do not need to deal with inter-*process* communication on the same processor.) The challenge is that often no information about the structure of the task graph or the cost of tasks is available at compile-time, making load balancing a more difficult runtime problem.

# 3.2 Notation

We will model mixed parallel applications as macro-dataflow graphs [95]. Each vertex of the (directed acyclic) graph represents a data-parallel task, which could be written using HPF or ScaLAPACK. The edges represent data and/or control dependency. In this chapter we will assume some advance knowledge of a task's running time given the number of processors it is allocated, at the time the task arrives, at the latest. The most general form in which this can be specified is function  $t_j(\mu)$ , which gives the running time of task

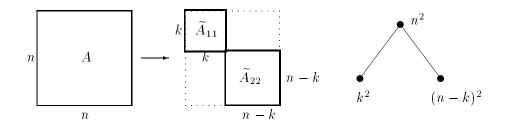


Figure 3.1: An example application with dynamic irregular mixed parallelism.

*j* if it is allocated  $\mu$  processors that are gang scheduled to run task *j* cooperatively, where  $1 < \mu < P$ , where *P* is the number of processors. In our switching algorithms we will schedule tasks on one or all processors; we will assume that  $t_j(1)$  and  $t_j(P)$  are effectively computable to reasonable accuracy.

**Example 3.2.1** Consider the problem of finding all eigenvalues of a dense non-symmetric  $n \times n$  matrix A. To find the n eigenvalues of A, which are complex in general, a new algorithm due to Bai and Demmel divides the complex plane into two half-planes and counts the number k of eigenvalues in one half-plane [12]. In terms of matrix operations, this is a data-parallel operation consisting mostly of several dense LU factorizations. k is not known before this operation. At the end of this operation, A is split into two diagonal blocks  $\tilde{A}_{11}$  which is  $k \times k$  and  $\tilde{A}_{22}$  which is  $(n - k) \times (n - k)$ ; the union of the eigenvalues of  $\tilde{A}_{11}$  and of  $\tilde{A}_{22}$  are the eigenvalues of A. Now we can recursively split  $\tilde{A}_{11}$  and  $\tilde{A}_{22}$ . This generates a dynamic binary divide-and-conquer tree. In this example, as in others that our scheduling framework is suitable for, each task j has a natural notion of "problem size," denoted  $N_j$ ;  $t_j(\mu)$  will depend of  $N_j$  in some well-defined way. For example, for the data-parallel operation above, as in matrix multiplication,  $N_j = O(n^2)$ , where the matrix A being split is  $n \times n$ , and  $t_j(1) = \Theta(N_j^{3/2})$ .

# 3.3 Switched parallel execution

The general scheduling solution for mixed parallel programs requires asynchronously scheduling tasks on subsets of processors. In this section we consider a simpler parallel execution model, where at any given time the scheduler is executing either a single task in data parallel mode over all the processors, or it is executing many tasks each running on only one processor. We call this the *switched* execution model. For divide-and-conquer, there is in fact only *one* switch point: large tasks near the root of the divide-and-conquer tree run in data parallel mode; the remaining tasks are packed into a task parallel mode.

Switching is simple to program and, for divide-and-conquer, has only one global data remapping phase. General processor subset scheduling may have better processor utilization but has much higher communication and synchronization overhead. In this section, we will give simple heuristics to schedule tasks in such a model, and prove some bounds on their performance. Then in §3.4, we study how close the performance of this restricted model is to the best possible. We set up the switching problem as follows.

**Input.** There is a set J of n tasks and P processors. These tasks have data parallelism. Task j specifies running times under two settings: on one processor, it takes time  $t_j(1)$  run in task-parallel fashion; on all P processors it takes time  $t_j(P)$  to run in data-parallel fashion. (Usually  $t_j(P) \ge t_1(P)/P$ , but we will not need this property.) The tasks may be independent as in §3.3.1, or there may be certain precedence restrictions between them, as in §3.3.2.

**Model.** We wish to minimize the execution time of a schedule with the following structure. The tasks are classified (partitioned) into two groups,  $J_P$  and  $J_1$ . Tasks in  $J_P$  are run one after the other in some order consistent with the precedence restrictions, allocating to each task all the P processors. Let the total time for this phase be  $t_P$ . Each task in  $J_1$  is run on one processor. They can run concurrently with each other, again as long as precedence restrictions are not violated. Let the total time for this phase be  $t_1$ . The total execution time is  $t_P + t_1$ .

**Classification.** The classification problem is to find  $J_P$  and  $J_1$  such that there is a two-phase schedule of optimal makespan where  $J_P$  is executed in data parallel mode and  $J_1$  is executed in task parallel mode.

**Scheduling.** Given  $J_P$  and  $J_1$ , the problem is to compute a good schedule. Note that scheduling  $J_P$  is trivial, and Graham's list scheduling suffices for  $J_1$ . Even after classification, constructing an optimal schedule is clearly  $\mathcal{NP}$ -hard via reduction from partition. It is easy to see that even the classification is  $\mathcal{NP}$ -hard.

Claim 3.3.1 The classification problem is  $\mathcal{NP}$ -hard, even for independent tasks.

**Proof.** By reducing partition to the classification problem. Given a partition instance  $A = \{a_1, \ldots, a_n\}$ , where these are positive integers, let there be P = 2 processors and n tasks, task j having  $t_j(1) = a_j$  and  $t_j(P) = \frac{a_j}{2} + \epsilon$  for suitably small  $\epsilon > 0$ . If there is a partition of A, then the best classification is task-parallel for all tasks. If a partition of A does not exist, then the best classification is data-parallel for all tasks.

However, it is often possible to classify tasks to achieve near-optimal makespan, as we see next. Thereafter we can use a standard list-type near-optimal scheduling algorithm to get an overall schedule within a small constant factor of the optimal switched schedule, in the worst case.

#### 3.3.1 Switching a batch

The input instance is a set J of independent tasks. Task i has size  $N_i$  and sequential running time  $t_i(1) = f(N_i)$  which increases with  $N_i$ . Assume all  $N_i$  are large enough that  $t_i(1) > t_i(P)$  (otherwise these small tasks would clearly be better off in the task parallel phase). The problem is to decide for each whether to execute it in data parallel mode or task parallel mode. It is clear that there is an optimal switched schedule that switches exactly once between the data-parallel and task-parallel phase.

A first cut solution is to define  $J_P = \{j : t_j(P) < t_j(1)\}$ ; this has been used in practice [17]. This is overly conservative since it does not consider the complete ready list to decide the fate of a single task. We use the following algorithm: sort the tasks in decreasing order of problem size, and find an index j such that the total time of executing tasks  $1, \ldots, j$  one after the other in data parallel mode followed by tasks  $j + 1, \ldots, |J|$  in a task parallel packing is minimized.

Let  $\operatorname{Pack}(P, S)$  be the makespan (length of schedule) generated by packing tasks from set S in task parallel mode into P processors. There are approximation algorithms that return  $\operatorname{Pack} \leq (1 + \epsilon) \operatorname{Pack}_{\operatorname{OPT}}$  for any fixed  $\epsilon > 0$ , within time that is polynomial in |S| [104]. Here  $\operatorname{Pack}_{\operatorname{OPT}}$  is the optimal makespan. It is easy to see that  $\operatorname{Pack}_{\operatorname{OPT}}(P,S) \leq \frac{1}{P} \sum_{s \in S} t_s(1) + \max_{s \in S} t_s(1)$ , by an averaging argument. Consider the following heuristic, which we call Prefix-Suffix.

1. Sort tasks in decreasing order of  $t_j(1)$ ; i.e., let  $t_1(1) \ge t_2(1) \ge \cdots \ge t_L(1)$ .

- 2. For  $1 \leq i \leq L+1$ , define  $p[i] = \sum_{1 \leq j < i} t_j(P)$  (p[1] = 0), and define  $s[i] = \text{Pack}(P, \{i, \dots, L\})$  (s[L+1] = 0).
- 3. Pick  $1 \le i^* \le L+1$  such that  $p[i^*]+s[i^*]$  is minimal over all i, such that  $1 \le i \le L+1$ .
- 4. Run tasks  $1, \ldots, i^* 1$  in data parallel mode, and tasks  $i^*, \ldots, L$  in task parallel mode.

For the analysis, consider a modified problem where tasks  $i, \ldots, L$  have to be scheduled using switched parallelism, given the constraint that the largest task i has to be run during the task parallel phase. Let  $s^*[i]$  be the optimal switched makespan subject to this new constraint.

Lemma 3.3.2  $\operatorname{Pack}_{\operatorname{OPT}}(P, \{i, \dots, L\}) \leq 2 s^*[i].$ 

**Proof.** Because  $\operatorname{Pack}_{OPT}(P, \{i, \dots, L\}) \leq \frac{1}{P}(t_i(1) + \dots + t_L(1)) + t_i(1), \text{ and } s^*[i] \geq \max\{\frac{1}{P}(t_i(1) + \dots + t_L(1)), t_i(1)\}.$ 

**Theorem 3.3.3** Independent tasks can be classified in polynomial time such that the resulting makespan is within a factor of  $(2 + \epsilon)$  of the makespan of the optimal schedule, for arbitrary constant  $\epsilon > 0$ .

**Proof.** Given an optimal schedule, let  $\ell$  be the smallest index in the above sorted order such that task  $\ell$  is executed in task parallel mode. Prefix-Suffix produces a schedule of length  $p[i^*]+s[i^*] \leq p[\ell]+s[\ell] = p[\ell]+\operatorname{Pack}(P, \{\ell, \ldots, L\}) \leq p[\ell]+(1+\epsilon) \operatorname{Pack}_{\operatorname{OPT}}(P, \{\ell, \ldots, L\}) \leq p[\ell]+2 (1+\epsilon) s^*[\ell] \leq 2 (1+\epsilon) (p[\ell]+s^*[\ell]) = 2 (1+\epsilon) \operatorname{OPT}$ , since  $\operatorname{OPT} \geq p[\ell]+s^*[\ell]$ .

In a dynamic divide-and-conquer tree, we assume that the problem size of tasks decreases down any path in the tree. For such trees, we can proceed as follows.

- 1. Maintain a ready-list of independent tasks/tasks that have been generated but not executed yet.
- 2. Before scheduling a module in data parallel mode, apply the above prefix-suffix heuristic to the ready-list of tasks.
- 3. If the best data parallel prefix is empty, decide to switch to task parallelism at this point.

This is only a heuristic, since in the worst case, an adversary can prune all but the largest task on the frontier immediately after the algorithm decides to switch. While this means that we have no provable bounds, such malicious behavior is rare in practice. We will show that this heuristic performs quite well in §3.5.

#### 3.3.2 Precedence graphs

In contrast to dynamic trees, if the precedence graph is a tree and is known before startup, we can provide a simple polynomial time constant factor algorithm. It uses dynamic programming and is very inexpensive compared to mathematical programming type approaches.

The problem instance is a set of n tasks, each task j specifying  $t_j(1)$  and  $t_j(P)$  as before. We will consider schedules with *one* switch: the large nodes at and near the tree root are processed in fully parallel mode, and then a switch is made to task-parallel execution of the remaining nodes.

**Definition 3.3.4** For task j in a directed acyclic precedence relation  $\prec$ , PathToLeaf $(j) = t_j(1) + \max_{j \prec i}$  PathToLeaf(i), and PathToRoot $(j) = t_j(1) + \max_{i \prec j}$  PathToRoot(i).

We will do binary search for the makespan between a lower and upper bound similar to the classical  $\epsilon$ -approximate decision procedures [104]. Suppose in the current search step the proposed deadline is D.

- 1. Compute PathToLeaf(j) by a bottom-up traversal. Compute  $S_P = \{j :$ PathToLeaf $(j) t_j(1) > D\}$ . If  $\sum_{j \in S_P} t_j(P) > D$  then deadline D is infeasible.
- We will solve a certain knapsack-like problem. The items to pack are tasks, these are assigned *profits* and *weights*. The knapsack has a weight limit; the goal is to maximize the profit of items packed. Let the profit of task j be t<sub>j</sub>(1), and the weight be t<sub>j</sub>(P). Obtain S'<sub>P</sub> = Knapsack(J, D) having total weight at most (1+ε)D such that the profit is at least that of the optimal knapsack packing. The algorithm Knapsack has been described in Chapter 4, §?? (page 82).
- 3. If  $\sum_{j \notin S'_{P}} t_{j}(1) > D \cdot P$  then deadline D is not feasible.

4. Execute  $S_P \cup S'_P$  on all P processors in any order consistent with the precedence constraints given by the tree edges. List schedule the remaining tasks in task parallel fashion.

**Lemma 3.3.5** If deadline D is feasible, the above schedule terminates in time  $(4 + \epsilon)D$ .

**Proof.** If D is feasible, then  $\sum_{j \in S_P \cup S'_P} t_j(P) \leq (2 + \epsilon)D$ . Also by a standard critical path argument the time to execute all  $j \notin S_P \cup S'_P$  using list scheduling is at most

$$\sum_{j \notin S_P \cup S'_P} t_j(1) + \max_{j \notin S_P \cup S'_P} \mathsf{PathToLeaf}(j) \leq 2D.$$

**Theorem 3.3.6** A single switch point for a task tree can be found that gives makespan within a factor of  $(4 + \epsilon)$  of the optimal switch, for an arbitrarily small constant  $\epsilon > 0$ . The algorithm takes time polynomial in  $1/\epsilon$  and the number of tasks.

One can solve series-parallel task graphs in a similar fashion. Notice the "single" clause; clearly, if  $t_j(1)$  and  $t_j(P)$  are arbitrary, a single switch can be very bad. However, tree precedences are most common in divide and conquer problems, where problem sized decreases monotonically down any path from the root; thus for any P and  $j_1 \prec j_2$ , the efficiency of running  $j_1$  on P processors is at least that of  $j_2$ . Under this realistic condition it is easy to see that one switch makespan is within a factor of two of many switches.

While these worst-case factors are small, they may mask the benefits of investing in mixed parallelism in the first place. Therefore, we will need to supplement these worst-case results by simulations and actual implementation. This we do next.

## 3.4 Modeling and analysis

In this section we will obtain analytical and empirical results that indicate that the switched scheduling model achieves efficiency quite close to the best possible, in our context of divide-and-conquer problems. For this we will develop a model in two parts:

- 1. We propose a model for the efficiency of a data parallel module as a function of N/P, the "problem size per processor."
- 2. We model regular divide-and-conquer trees in terms of the out-degree d and the *shrink* factor c, the ratio of the problem size of a parent to that of a child.

#### 3.4.1 Efficiency of data parallelism

If all the data parallel modules j at the nodes of the task graph where ideally scalable, i.e., had  $t_j(\mu) = t_j(1)/\mu$  for  $\mu$  as large as P, the number of processors in the machine, then using mixed or switched parallelism is a non-issue. The gains from extending beyond pure data parallelism are a function of how poorly scalable the data parallel modules are. In this section we propose an efficiency model for a data parallel module.

We let e(N, P) be the parallel efficiency of solving a problem of size N on P processors. Informally, the "size" N of a problem or task is the total data volume, e.g., for  $n \times n$  matrix multiplication,  $N = \Theta(n^2)$ . If the serial running time is f(N), the parallel running time r(N, P) on P processors is  $r(N, P) = f(N)/(P \cdot e(N, P))$ . e(N, P) depends on the algorithm, and relative speeds of computation and communication. Despite e's possibly complex dependence on all these parameters, we will show that for a number of algorithms of interest, e(N, P) is accurately modeled by a simple two-parameter function of the problem size per processor, N/P.

By Amdahl's law, we expect e to be a decreasing function of P, with  $e(\cdot, 1) = 1$ . So our intuition is that e(N, P) should be an increasing function of N/P. We will let  $e_{\infty} \leq 1$ be its asymptotic value for large N/P. The next question is how e(N, P) approaches  $e_{\infty}$ . There are, of course, an infinite number of functions to model this, but we shall propose a simple model that we will empirically validate. Roughly speaking, the model captures programs having an area-to-volume relationship between communication and computation, which abounds in parallel scientific applications.

#### 3.4.1.1 A proposed model

One model we have found to agree very closely with experimental data is the following. The efficiency of a data parallel task of size N on P processors is modeled as

$$e(N,P) = \begin{cases} 1 & \text{if } P = 1\\ \frac{e_{\infty}}{1 + \sigma P/N} & \text{if } P > 1. \end{cases}$$
(3.1)

The parameter  $\sigma$  measures how fast the efficiency approaches its asymptotic value  $e_{\infty}$ . As shown in Figure 3.2 the efficiency reaches half its asymptotic value when  $N/P = \sigma$ . Thus, the smaller the value of  $\sigma$ , the more efficient the implementation is for a fixed problem size.

The parallel running time r(N, P) is

$$r(N,P) = \frac{f(N)}{e_{\infty}} \left(\frac{1}{P} + \frac{\sigma}{N}\right).$$
(3.2)

Equation (3.2) says that adding processors has diminishing returns, much like Amdahl's law. However, since no sequential and perfectly parallel components can be identified, this model is not identical to Amdahl's law. We used this model to estimate  $t_j(1)$  and  $t_j(P)$  for our switching algorithms, but any other estimator would also work fine.

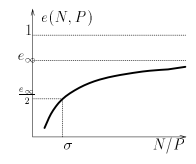


Figure 3.2: The proposed efficiency model for data parallelism within a single task.

#### **3.4.1.2** Experimental validation

We validated our model using experimental data. In Figure 3.3 on page 49, we consider three ScaLAPACK programs: LU, QR and Cholesky factorizations, and three machines: the Delta, Paragon and iPSC/860 [45]. Each graph plots performance in GFLOPS per processor versus N/P, including experimental data (the circles), as well as the prediction of the asymptotic model. The iPSC/860 experiments were run with 128 processors, and the Paragon and Delta experiments were run with both 128 and 512 processors. Each graph includes an estimate s\_inf (which is proportional to the asymptotic efficiency  $e_{\infty}$ ) of the per-processor GFLOPS as  $N/P \to \infty$  and an estimate of  $\sigma$  (sigma). The asymptotic model is a good fit for the actual efficiency profiles: the mean relative error is 6-11%.

Estimates of  $\sigma$  are important for performance analysis as well as runtime scheduling decisions. To this end, we collect values of  $\sigma$  for some parallel scientific libraries [45], using existing analytical performance models [40, 42]. For each of these routines, we have available the communication and computation time as functions of problem size, number

Machine	$\alpha$	$\beta$	M/P	$\sigma_{ m MM}$	$\sigma_{ m LU}$	$\sigma_{ m BS}$	$\sigma_{ m SF}$
Alpha+ATM1	$3.8  imes 10^5$	62	64	$1.3 \times 10^{4}$	$5.7 imes10^6$	$3.4  imes 10^6$	$2.7 imes10^6$
Alpha+ATM2	$3.8  imes 10^5$	15	64	6500	$5.6 imes10^6$	$3.4  imes 10^6$	$2.7 imes10^6$
Alpha+Ether	$3.8  imes 10^5$	960	64	$2.5 \times 10^5$	$6.9  imes 10^6$	$4.2  imes 10^6$	$3.4 imes10^6$
Alpha+FDDI	$3.8  imes 10^5$	213	64	$4.1 \times 10^4$	$5.9 imes10^6$	$3.6  imes 10^6$	$2.9  imes 10^6$
CM5	450	4	32	53	490	2234	3826
CM5+VU	$1.4 \times 10^4$	103	32	9100	$3.1  imes 10^5$	$1.9 \times 10^5$	$1.53 \times 10^5$
Delta	4650	87	16	7400	$1.5  imes 10^5$	$9.3  imes 10^4$	$7.2  imes 10^4$
HPAM (FDDI)	300	13	64	154	9300	5400	4250
iPSC/860	5486	74	16	5490	$1.5  imes 10^5$	$9.2 \times 10^4$	$7.3  imes 10^4$
Paragon	7800	9	16	633	$1.25  imes 10^5$	$7.7 \times 10^4$	$6 \times 10^4$
SP1	$2.8 \times 10^4$	50	64	4250	$4.8  imes 10^5$	$2.9 \times 10^5$	$2.4 \times 10^5$
T3D	$2.7  imes 10^4$	9	64	1544	$4.2  imes 10^5$	$2.5  imes 10^5$	$2 \times 10^5$

Table 3.1: Estimates of  $\sigma$  for different machines and problems in the asymptotic model. The Alphas use PVM as messaging software. ATM1 = current generation; ATM2 = projected next generation. HPAM = a cluster of HP workstations connected by FDDI with a prototype active message implementation. The programs are matrix multiplication (MM), LU factorization (LU), backsolve (BS), and sign function (SF, discussed later).  $e_{\infty} = 1$  for all problems in this table. Parameters  $\alpha$  (latency) and  $\beta$  (inverse bandwidth; transfer time per double) are normalized to a BLAS-3 FLOP, and the model is fit to data generated from analytical models [40, 42, 107]. The curves were fit for  $2 \leq P \leq 500$  and  $100 \leq n = N^{1/2} \leq 10000$ . An estimate of memory per processor in megabytes is given in the column marked M/P. Estimates for  $\alpha$  and  $\beta$  are in part from [107, 117, 88, 7].

of processors, network latency, and network bandwidth. Using these given functions, we first estimate the parallel running time r(N, P) for a given machine and problem, then fit Equation (3.1) to it using Matlab. The results are presented in Table 3.1.

#### 3.4.2 Regular task trees

The second part of our model has to address the task graph structure. The  $\mathcal{NP}$ -hardness of the special case of independent tasks *a fortiori* means that the scheduling problem (in either the switched or mixed model) is also  $\mathcal{NP}$ -hard. The best algorithms known are constant factor approximations, with worst case factors in the range 2–2.6 [111]. Unfortunately, a constant factor of this magnitude (which we will call "packing loss") may substantially mask the benefits which would otherwise be obtained from mixed parallelism. Furthermore, we know of no tighter analysis of this constant for a given graph, and the worst case constant may not be meaningful in the average instance.

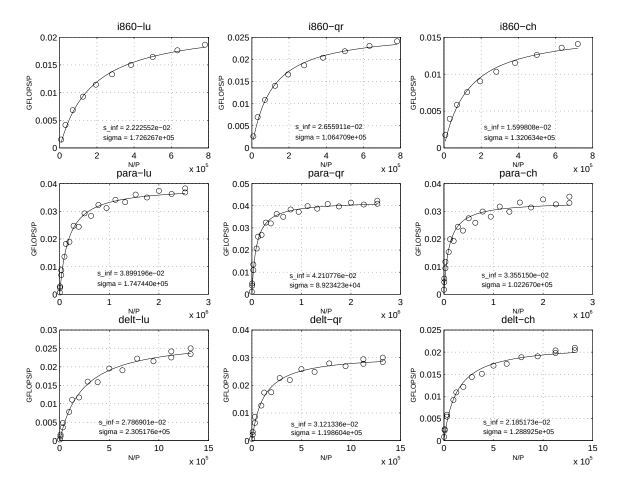


Figure 3.3: Validation of the asymptotic model using data from the ScaLAPACK implementations of LU, QR, and Cholesky (CH) factorization programs. The machines are iPSC/860 (i860), Paragon (para) and Delta (delt). In each graph, s\_inf is the per-processor GFLOPs as  $N/P \rightarrow \infty$ , which is proportional to the asymptotic efficiency  $e_{\infty}$ . sigma is the  $\sigma$  in the asymptotic model.

#### 3.4.2.1 Model

We therefore consider how mixed parallelism will perform in very favorable circumstances, namely in regular divide and conquer trees. Our assumptions are listed below.

- The task graph is a complete tree with branching factor  $d \ge 2$ .
- The d child tasks of a task of size N are all of size N/c, where c > 1.
- The work required to do a task of size N is  $f(N) = N^a$ , where  $a \ge 1$ .

We call such regular trees that have a root size of N as (N, a, c, d) trees. This is a very simple model, and so we need to understand the limits of its applicability. First, task communication is not accounted for. But for a > 1, task communication cost is generally of lower order than the node cost f(N) (e.g., O(N) vs.  $O(N^{3/2})$  in the case of many dense matrix operations). Thus, we expect our model to overestimate the benefits of mixed parallelism, provided problems are large enough. Therefore, a prediction of little benefit from mixed parallelism for a particular problem is likely to be trustworthy, while a prediction of great benefit from mixed parallelism must be further analyzed. Second, for evaluating efficiency gains with high accuracy, only regular trees could be considered. In spite of their simplicity, regular trees provide a good starting point for preliminary analysis [18].

#### 3.4.2.2 Examples

**Eigenvalue algorithms.** Several eigenvalue algorithms exhibit mixed parallelism because they use divide and conquer. One such algorithm was described in Example 3.2.1 on page 40. If the divide step on a matrix of size  $N = n^2$  is perfect, each child is of size  $\frac{n}{2} \times \frac{n}{2}$ , or N/4. Performing this separation requires  $O(N^{3/2})$  FLOPS. This successive separation process forms a binary tree with c = 4, d = 2, and a = 3/2. (We scale time so that the constant in  $O(N^{3/2})$  becomes one.)

For symmetric matrices, an algorithm similar in spirit is the beta-function technique of Bischof *et al* [17]. An eigenvalue algorithm of a different flavor, but still from the divide and conquer category, is Cuppen's method for symmetric tridiagonal matrices, where we can actually split the matrix exactly in half all the time [38, 99] (although the costs of the children are not so simple).

**Sparse Cholesky.** We consider the regular but important special case of the matrix arising from the 5-point Laplacian on a square grid, ordered using the nested dissection ordering [60]. In this case one may think of dividing the matrix into 4 independent subproblems, corresponding to dividing the square grid into 4 subsquares, each of half the perimeter. The work performed at a node which corresponds to an  $n \times n$  grid is  $O(n^3)$ ; most of this cost is a dense Cholesky of a small  $n \times n$  submatrix corresponding to the nodes on the boundaries of the subsquares. Thus  $N = n^2$ , a = 3/2, c = 4 and d = 4. We will also see that the results go over to matrices with planar graphs.

#### 3.4.3 Analysis

We make some simple observations about batches of independent tasks. For a single task with sequential running time f(N), the choice is only between 1 and P processors, and the running time is

$$f(N) \times \min\left\{\frac{1}{e_{\infty}}\left(\frac{1}{P} + \frac{\sigma}{N}\right), 1\right\}$$

For a batch of L independent tasks, each of size N, the sequential running time  $t_1$ of all L tasks is Lf(N). The data parallel running time  $t_D$ , where we run each task in data parallel fashion one after the other, is just L times the above expression. We let  $t_T$  denote the task parallel running time, where we assign one processor per task. Finally, we let  $t_M$  denote the mixed parallel running time, the optimal running time over all possible assignments of processors to tasks. Let  $e_D$ ,  $e_S$ , and  $e_M$  be the corresponding overall efficiencies.

By allocating only one processor per task, we can get parallel execution time  $\lfloor L/P \rfloor f(N)$ . Since the work lower bound is Lf(N)/P, pure task parallelism is optimal when  $L \ge P$  (modulo rounding effects, which we ignore here and elsewhere). Thus, we need only consider the case L < P. Also assume L divides P. We note the following easy observations.

- 1. When L divides P, the running time for L independent tasks of size N in the asymptotic model using optimal mixed parallelism is  $t_M = f(N) \times \min\left\{1, \frac{1}{e_{\infty}}\left(\frac{L}{P} + \frac{\sigma}{N}\right)\right\}$ ,
- 2. When L divides P, the running time using data parallelism is  $t_D = L f(N) \times \min\left\{1, \frac{1}{e_{\infty}}\left(\frac{1}{P} + \frac{\sigma}{N}\right)\right\}.$

3. When task parallelism is not optimal, the relative improvement of using mixed parallelism over pure data parallelism for the batch problem is

$$\frac{e_M}{e_D} = \frac{\frac{N}{\sigma P} + 1}{\frac{N}{\sigma P} + \frac{1}{L}}$$
(3.3)

**Example 3.4.1** We apply this analysis to complex matrix multiplication, which was reported as a benchmark for the Illinois Paradigm compiler [95]. The task is MM, the machine is the CM5 without vector units, and L = 4. From Figure 3.1 we obtain  $\sigma = 53$  and  $e_{\infty} = 1$  for this problem. To attain a relative improvement  $\epsilon$  of mixed over data parallelism, i.e.  $\epsilon_D/e_M < 1 - \epsilon$ , we need the problem size to be small. Specifically, if each MM involves  $n \times n$  matrices, we can substitute the numbers into (3.3) and see that n needs to be less than roughly  $\sqrt{53P(3-4\epsilon)/4\epsilon}$  for this improvement. E.g., if P = 64 and  $\epsilon = 0.5$ , then n < 42, a tiny problem indeed. It is interesting that the experiments reported in [95] for the CM5 use  $P \in \{64, 128\}$  processors and n = 64. On the Paragon with P = 512 and  $\sigma = 633$ , to ensure  $e_D/e_M < 0.5$  as above, we will need roughly that n < 569, which is still not large by the standard of many scientific applications: a matrix of this size fills only 0.03% of the Paragon's total memory.

The conclusion is that MM data parallelizes too well to benefit much from mixed parallelism on a machine as balanced (i.e. with as low a  $\beta$ ) as the CM5 without vectors units. Better hunting grounds for cases where mixed parallelism helps significantly are more unbalanced machines (high  $\beta$ ) and problems with less scalable data parallel components.

**Example 3.4.2** Changing the problem to BS (backsolve; solving an  $n \times n$  upper triangular system of linear equations) in the last example and changing the machine to a 16 processor SP1, we see that n < 1386 must hold for  $e_D/e_M \leq 0.5$ . This is a relatively realistic size. Mixed parallelism improves the efficiency from roughly 33% to 66%.

#### 3.4.4 Unbalanced batch problems

So far we have considered batches of tasks of identical size N. Here we argue that permitting tasks of different sizes makes data parallelism only closer in performance to optimal mixed parallelism, because, if there are a few large tasks that dominate the work content of the batch, the margin for improvement over pure data parallelism will be small. To prove this, suppose we have a batch of L > 1 tasks, the *i*-th task of size  $N_i$ . Suppose all tasks have the same efficiency profile  $e(N_i, P)$ , and that the sequential processing time for task *i* is  $f(N_i)$ , where  $f(x) = x^a$  as before. We will need the following theorem for our proof.

**Theorem 3.4.3 (Hölder's Inequality)** Let  $x_k, y_k > 0$  for  $1 \le k \le L$ , p > 1, and q be such that  $\frac{1}{p} + \frac{1}{q} = 1$ . Then

$$\sum_{k} x_k y_k \leq \left(\sum_{k} x_k^p\right)^{1/p} \left(\sum_{k} y_k^q\right)^{1/q}.$$

Let the pure data parallel running time be  $t_D$ , the optimal switched execution time be  $t_S$ , and the optimal mixed parallel running time be  $t_M$  ( $t_M \leq t_S \leq t_D$ ). We will show that, among all possible batches, a balanced batch poses the worst instance for data parallelism and thus provides the greatest potential for improvement through mixed parallelism. Of course, certain quantities have to remain invariant over the space of maximization. Two invariants are possible:

- 1. The total work  $\sum_{i} f(N_i) = F$ , a constant.
- 2. The total size  $\sum_i N_i = N$ , a constant.

We will consider both variations.

**Lemma 3.4.4** With  $t_D$ ,  $t_M$ , and f defined as above,

$$\frac{e_M}{e_D} = \frac{t_D}{t_M} \le \frac{1}{e_\infty} + \frac{\sigma P}{e_\infty} \left( \frac{\sum_k f(N_k) / N_k}{\sum_k f(N_k)} \right)$$

**Proof.** Immediate, using  $t_M \geq \frac{1}{P} \sum_k f(N_k)$ .

The remaining exercise is to bound from above the parenthesized term in the above RHS.

**Theorem 3.4.5** Subject to either invariant,

$$\frac{e_M}{e_D} \leq \frac{1}{e_\infty} \left( 1 + \frac{\sigma PL}{N} \right).$$

**Proof.** Here we will do a continuous analysis, assuming  $N_i$ 's are real, rather than integers. Also assume  $N_i > 1$  to avoid problems near zero.

For constant F, the quantity in parentheses is maximized when all  $f(N_i) = F/L$ . This follows since  $x/f^{-1}(x) = x^{1-1/a}$  which is convex for a > 1.

For constant N, using p = a/(a-1) and q = a in Theorem 3.4.3, we obtain

$$\sum_{k} N_k^{a-1} \cdot 1 \leq \left(\sum_{k} N_k^a\right)^{1-\frac{1}{a}} \cdot L^{1/a}.$$

Since  $\sum_k N_k^a \ge L(N/L)^a = N^a/L^{a-1}$ , we have

$$\frac{\sum_k N_k^{a-1}}{\sum_k N_k^a} \leq \frac{L^{1/a}}{\left(\sum_k N_k^a\right)^{1/a}} \leq \frac{L}{N}$$

This value is achieved when all  $N_k$  are set to N/L.

It can be verified that the claim also holds for functions of the form  $f(x) = x \log x$ , etc., so the claim is quite broadly applicable. We can trivially get similar bounds on the efficiency gap for irregular task graphs by throwing away the dependency information.

**Corollary 3.4.6** The maximum benefit from mixed parallelism for a task graph G with L vertices such that the sum of problem sizes is N can be bounded as

$$\frac{e_M}{e_S} \leq \frac{e_M}{e_D} \leq \frac{1}{e_\infty} \left(1 + \frac{\sigma PL}{N}\right),$$

in the asymptotic model using P processors.

We can also derive some heuristic bounds to the performance gap in some irregular graphs. E.g., Gilbert and Tarjan study nested dissection algorithms to solve sparse systems on planar graphs [62], where a problem of size N is divided into d = 2 subproblems, where each part is no bigger than 2N/3. No matter what strategy we use in the upper levels, we only need to go down roughly  $\ell(\epsilon) = \frac{\lg(\epsilon/P)}{\lg(2/3)} \approx 1.71(1 - \frac{\log \epsilon}{\log P}) \lg P$  levels before the largest leaf is of size at most  $\epsilon N/P$ . At this point task packing is at most  $(1 + \epsilon)$  times optimal. Given that there is not much need to go below this level,  $L \leq P^{1.71}$ , so the maximum benefit cannot be much larger than  $t_S/t_M \leq t_D/t_M \leq 1 + \sigma P^{2.72}/N$ .

#### 3.4.5 Simulations

As we mentioned earlier, to get accurate estimates of the performance gains from mixed or switched parallelism, we use simulations based on the data parallel and task graph models.

Our space of optimal schedules includes those that assign arbitrary subsets of processors to the tasks. This involves the following runtime support:

- It must be possible to create and destroy subsets of the set of physical processor and rename them in a virtual processor space. These are called *contexts* in the Message Passing Interface (MPI) jargon [44]. Note that every message has to be indirectly addressed to a virtual destination processor, and if many contexts are defined hierarchically, many table-lookups are needed for each message.
- Contexts need to provide synchronization and collective communication primitives, so that they can be gang-scheduled. These can be potentially expensive. The CM-5, for example, provides a 5  $\mu$ s hardware barrier over the physical partition, but the best software barrier over a processor subset takes over 50  $\mu$ s.

In our simulations, we will ignore these overheads, and thus get an upper bound to the best possible efficiency, and compare our switched performance against this upper bound. We will also compare switching with pure data parallelism as in HPF.

**Results.** The space of programs, machines, and problem sizes is too large to examine completely; therefore we take some slices through this space that give insight into the benefits of mixed parallelism for typical current architectures and our suite of scientific programs. The following graphs are shown.

- 1. We fix P = 128,  $e_{\infty} = 1$ , a = 3/2, and c = d = 4 (as in sparse Cholesky), and plot  $e_D/e_M$  and  $e_S/e_M$  against  $\sigma$  (log-scale) in Figure 3.4. The memory per node is assumed to be 64 MBytes, and the four plots correspond to problem sizes that fill 25, 50, 75 and 100% of the memory. For typical values of  $\sigma$  for various machines see Table 3.1.
- 2. For the same sparse Cholesky problem, we consider four machines. In each case, the x-axis is P. N is such that the memory is completely filled. We plot  $e_M$ ,  $e_S$ , and  $e_D$  against lg P in Figure 3.5.
- 3. The setting is as above, except that typical values of P are chosen for each machine and the x-axis is  $n = N^{1/2}$  (log-scale). See Figure 3.6.
- 4. The setting is as in item (3), but the problem is the sign function program (c = 4, d = 2). See Figure 3.7. For each task, as a reasonable estimate, there are 15 LU's, 15 BS's and 8 MM's. For this compound data parallel task, estimates of  $\sigma$  for various

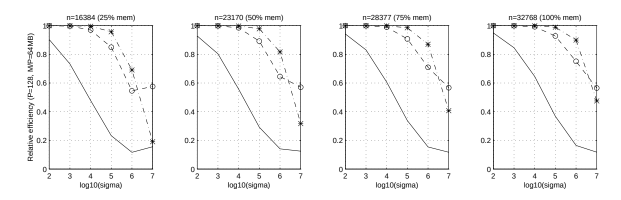


Figure 3.4: Cholesky. The line with stars is absolute efficiency of mixed parallelism  $(e_M)$ . The dashed line with circles is the relative efficiency  $e_S/e_M$  of switched parallelism. The solid line is the relative efficiency  $e_D/e_M$  of data parallelism.

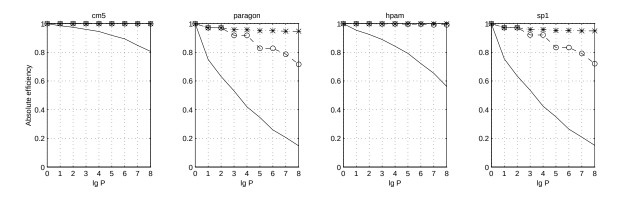


Figure 3.5: four machines using mixed, switched, and data parallelism.  $e_M$  (starred),  $e_S/e_M$  (circled), and  $e_D/e_M$  (solid) are plotted against lg P, always choosing N to fill all memory.

machines are shown in column SF of Table 3.1 ( $e_{\infty} = 1$ ). If Cuppen's eigenvalue algorithm is used, and the effect of "deflation" is small [38], the task tree has the same parameters as the sign function example above, although  $\sigma$  is different.

**Comments.** From Table 3.1, typical values of  $\sigma$  are all in the  $10^2$  to  $10^6$  range. Throughout this range, switched parallelism appears to make up for much of the deficit in data parallel performance. The non-monotonicity in Figure 3.4 occurs because after  $\sigma$  becomes absurdly large (>  $10^6$ ), parallelism is no longer effective. In general, for fine-grain MPP-class machines, mixed parallelism has little marginal benefit, while for more coarse-grain networks of workstations, switched parallelism is adequate. The choice

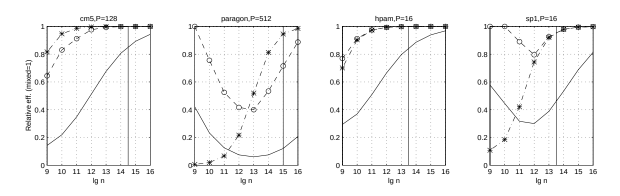


Figure 3.6: except that typical values of P are chosen for each machine and the x-axis is  $n = N^{1/2}$  (log-scale). Maximum size limit is shown by the vertical bar, where memory per node is from Table 3.1. The line with stars is absolute efficiency of mixed parallelism  $(e_M)$ . The dashed line with circles is the relative efficiency  $e_S/e_M$  of switched parallelism. The solid line is the relative efficiency  $e_D/e_M$  of data parallelism.

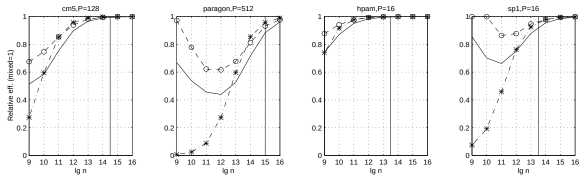


Figure 3.7: The setting is as in Figure 3.6, except that the problem is changed to sign function, with parameters c = 4, d = 2, and a different value of  $\sigma$ .

of strategy is dictated not only by  $\sigma$ , N and P, but also the size reduction factor c and branching degree d. This is seen in figures 3.6 and 3.7: mixed parallelism gives less marginal benefit over switched or data for small values of d and large values of c (and vice versa). Figures 3.6 and 3.7 exhibit troughs because at the lower end of problem sizes, absolute efficiency of all the strategies are very small but close to each other.

# 3.5 Experiments

Encouraged by the promising simulation and analysis, we implemented the switched parallel scheduler as a new module in the ScaLAPACK scalable scientific software library [45]. In particular, it was used to schedule the divide and conquer tree generated by the sign function example in §3.2. The ScaLAPACK library is structured as follows. The lowermost layer called BLACS (Basic Linear Algebra Communication Subroutines) is built on top of a message passing layer such as NX (on the Paragon), MPI/MPL (Message Passing Interface, on the IBM SP's), or CMMD (on the CM-5). For us BLACS provides two types of functions:

- Routines to define *contexts*: these are subsets of processors that form a virtual machine and can participate in collective communication.
- Routines to transfer array sections between contexts.

On top of BLACS is built PBLAS (Parallel Basic Linear Algebra Subroutines): these handle elementary matrix-vector and matrix-matrix operations with block-cyclic layouts. ScaLAPACK itself is the topmost layer providing higher functionality like Cholesky and LU factorization.

#### 3.5.1 Software structure

Our implementation of the library has the following components.

- PBLAS/ScaLAPACK subroutines are modified to export the FLOP count of relevant subroutines symbolically in terms of the input problem size. Currently we implement this by hand using header files.
- The machine can be queried for its message latency and bandwidth. This is a set of functions similar to query functions for machine epsilon and overflow.

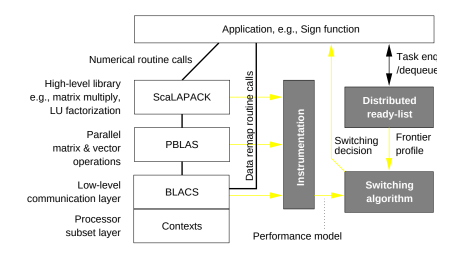


Figure 3.8: The software architecture of the dynamic scheduling module in ScaLAPACK.

- Our (σ, e<sub>∞</sub>) model is constructed from the above information statically at configuration time. This is later used to estimate t<sub>j</sub>(1) and t<sub>j</sub>(P) for tasks j on the ready-list at runtime.
- There is a distributed data structure for the ready-list. Before the switch-point, this is essentially a replicated work-list kept consistent across all processors. The scheduling computation is currently replicated on all processors as well. When the switching decision is made, several things happen. First, new singleton contexts are generated in each processor. Second, a preliminary task packing is done to decide which task goes where. Third, a global data remap phase moves arrays distributed over all processors to the appropriate singleton context. At this point the work-lists of different processors are different. Finally, the ready-list behaves as a task-parallel distributed queue with work-stealing [19].

#### 3.5.2 Results

We measured performance on two machines: a 25-processor Paragon and a 36-processor SP-2. The results are shown in Figure 3.9. Each graph plots performance (MFLOPS) against problem size. The problem size  $\sqrt{N}$  is the side of the matrix at the root of the divide and conquer tree. The bottom line shows "data-parallel" performance. This is not merely letting pure data parallelism take its course on the entire machine right

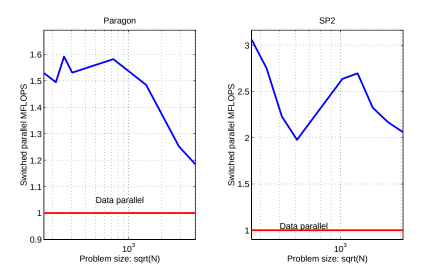


Figure 3.9: Performance improvements of switched parallelism over data-parallel execution on the Paragon and SP2, with the latter being scaled to 1. The data parallel baseline scheduler switches when the largest task j in the ready-list has  $t_j(1) < t_j(P)$ ; the switched scheduler applies the prefix-suffix heuristic. The x-axis is problem size n, the side of the matrix at the root of the divide-and-conquer tree, and the y-axis is MFLOPS normalized to the purely data-parallel version (higher is better).

down to leaves (of size  $1 \times 1$  or  $2 \times 2$ ). Instead, we look at the largest task j in the ready-list, and if  $t_j(1) < t_j(P)$ , where P is the total number of physical processors, we switch. This is what the most reasonable HPF runtime may do, although we certainly cannot claim even production compiler or runtime systems do this.

The upper line is the performance of our prefix-suffix heuristic relative to data-parallel as above; higher is better. On the Paragon, which has a weak CPU and relatively strong network (at least for coarse-grain communication) the gains are already up to 50%; on the more recent SP-2 with a significantly worse communication to computation cost, the gains are between 200 and 300%. In both cases, as  $N \to \infty$  no optimization is needed; the performance of data and switched parallelism tend to the same limit (1, for this problem). While not quite infinite, our problem range  $\sqrt{N}$  goes into several thousands, and therefore the performance gains are much more impressive and meaningful than the benefits on static problems typically of size  $\sqrt{N} = 64 \dots 128$ , on P = 64 processors [110, 95].

We have also measured some typical numbers for the overhead of the data remap step in the switching algorithm. They are quite low; a few percent. See Figure 3.10. But the

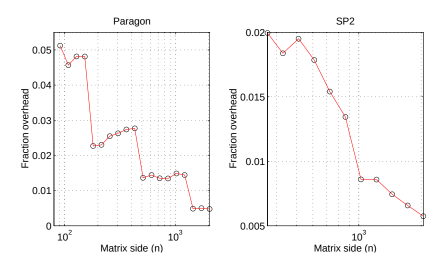


Figure 3.10: More statistics. The x-axis is the problem size as before. The y-axis gives the fraction of total time spent in parallel by all processors doing data remapping during switching.

task graphs expanded typically have ten or more data-parallel nodes; if we had implemented general mixed parallelism with arbitrary processor subsets, a data remap would be needed before and after executing each data-parallel task, blowing up this overhead to perhaps tens of percents. Thus switched parallelism is not only appealing for its simplicity, but also for locality reasons.

# **3.6** Related work

As noted before, the idea of supplementing data parallelism with task parallelism where available is quite obvious and has been explored earlier. In adaptive mesh refinement (AMR) algorithms, there is task parallelism between meshes and data-parallelism within a mesh [11]. In computing eigenvalues of nonsymmetric matrices, the sign function algorithm does divide and conquer with matrix factorizations at each division [12]. In timing-level circuit simulation there is parallelism between separate subcircuits and parallelism within the model evaluation of each subcircuit [116]. In sparse matrix factorization, multi-frontal algorithms expose task parallelism between separate dense sub-matrices and data parallelism within those dense matrices [85]. In global climate modeling [89], there are large data parallel computations performed on grids representing the earth's atmosphere and oceans, and task parallelism from the different physical processes being modeled.

Several researchers have proposed support to take advantage of this mixed parallelism. In the theory area, the best known on-line scheduling algorithm for mixed parallelism is 2.62-optimal [16, 52], and the best off-line algorithm is 2-optimal [113, 87]. But these are worst-case guarantees. In the systems area, the Paradigm compiler [95], iWarp compiler [110], and NESL compiler [32] all support forms of mixed task and data parallelism, and there are plans to merge data Fortran D with Fortran M [56] and pC++ with CC++ [84] to support mixed parallelism. The compiler efforts all depend on static task graph and profile data and perform extensive optimizations to allocate processors. We have already noted how dynamic task parallelism is more promising to exploit.

A problem domain quite different from scientific computing where mixed parallelism has been used is in parallel database query scheduling. In a relational database, each query is roughly speaking an in-tree of parallelizable tasks; where each task is an operator like a join or select. Each operator has partitioned parallelism similar to data-parallelism. A chain of operators may also show *pipelined* parallelism, but we can regard those nodes to be collapsed into a single node with data parallelism. There is also task parallelism between unordered nodes. Recently algorithms have been designed for trading between locality and load balance in this scenario [33]. We will come back to similar problems in Chapter 4.

The switching technique has been independently discovered *after* our paper [26] was published in a different context: that of scheduling tasks with penalties. Every task has a running time, and a penalty for rejection; the goal is to minimize the sum of the makespan of accepted tasks and the penalty of rejected tasks [15]. While their main result is an on-line algorithm for independent tasks, we also give an off-line algorithm for trees (this can be generalized to series-parallel graphs).

# 3.7 Discussion

We have shown in this chapter that by exploiting dynamic task parallel to supplement the diminishing returns of data parallelism in mixed parallel divide-and-conquer applications, and by designing simple heuristics for switched scheduling, significant performance improvements are possible. We conclude the chapter with a few comments.

What happens if the leaf tasks are different from the interior tasks? It is usually the case that the leaf tasks are more efficient in terms of CPU operations, but they may be less

parallelizable. This in fact happens in the non-symmetric eigenvalue code. The prefix-suffix heuristic can be modified so that the task packing is done with the leaf estimates. This works well in practice (the profit margin is even higher than 300%), but is not interesting to report on from the modeling and analysis point of view.

What happens if the task tree folds back, as in a conquer phase? In our application domain, the sizes are known during the unfolding, so the folding back will be a static scheduling problem. In terms of the runtime strategy, processors can do one of these two things:

- rendezvous at the first phase switch point, compute the second phase reverse switch frontier, and go into a task parallel phase in between, OR
- go into the first phase task parallel mode as before, rendezvous before the folding back, compute the reverse switch frontier, and proceed as above.

Can we extend the analysis to make global statements about the switching heuristic, in particular, qualify the empirical intuition that since problem sizes decrease down any path, a switching decision cannot be far wrong? This will depend on the out-degree and the size shrink factor.

What if data transfer cost at the switching point/s are significant? This will be particularly important for arbitrary task graphs with arbitrary problem sizes at the nodes, where many switches may be needed. In that case we will need to minimize number or cost of switches, or trade that against the costs of data parallel inefficiency and task parallel packing loss.

# Chapter 4

# Scheduling resource constrained jobs

# 4.1 Introduction

So far in this thesis we have been concerned mostly with optimizing the performance of a single parallel program. On a typical multiprocessing system several such programs run simultaneously, contending for CPU time and other resources. While classical scheduling theory has been largely concerned with processor scheduling, memory, network and IO bandwidth are other resources which dictate how best to schedule programs with diverse needs. Moreover, classical scheduling results are predominantly for minimizing makespan (the finish time of a fixed set of programs); in real life, system utilization, fairness, and honoring priority are also important. In this chapter we will develop new techniques for scheduling such workloads.

To use terminology consistent with scheduling literature, we will refer to what independent users submit as *programs*. Each program is a collection of *jobs*. (In Chapter 3 there was only one program, and we called the jobs parallelizable or data-parallel tasks.) Every job specifies some resources that it needs to reserve during execution and perhaps some other jobs that need to finish before it can start.

Judicious job scheduling is crucial to obtaining fast response and effective utilization. Consequently, algorithms for scheduling have been extensively researched since the 60's, both in theory and in practice. In the past decade, there has been increasing interest in scheduling specifically for parallel systems. There is increased awareness of general resource scheduling problems, rather than just processor scheduling [50].

Owing to the vastly different nature and applicability of sequential and parallel computing systems, the scheduling problems of practical interest are rather different in the two settings. For example, scheduling general-purpose jobs is essential for a uniprocessor operating systems, while it is hardly an issue yet for existing parallel systems, none of which run the average workstation workload; similarly, scheduling threads for massive game tree searches is a reality for parallel systems while it is not an issue for uniprocessors.

In this chapter, we extend our scope from optimizing single programs using compiler and runtime approaches to that of scheduling multi-user parallel workloads. To propose a model, we focus on applications where parallelism has been effectively exploited. We isolate parallel databases and scientific computing as two such areas. In both these areas the computation is well-structured so regular subproblems can be solved in parallel, and the resource requirements of jobs can be estimated in advance. For these reasons, parallel computing has proven highly successful in these two areas. Here, we carefully model the resource scheduling problems in these applications and study them.

In the model we abstract from these applications, many users submit *programs* to the system over time; each program is a directed acyclic graph of *jobs*; the graph is assumed to be known when the program is submitted. A job is characterized by a set of resource requirements, which are also known at submission time. Resources are of various different types. Some resources, like processors, can be traded for time gracefully, while others, like memory, are not flexible in such a continuous fashion. The acyclic graph represents precedence constraints; each job specifies a set of predecessors that must complete before it starts.

We initiate a study of such resource scheduling models. Two features distinguish our goal from many classical results. First, our jobs are themselves parallelizable, like the data-parallel tasks in the applications in Chapter 3. Second, rather than minimizing the finish time of a fixed set of jobs, we wish to accept programs over time; these programs have priorities and we minimize the *sum* of finish time weighted by the respective priorities (this is called a *minsum* metric, as compared to makespan, which is a *minmax* metric). The results in this chapter represent initial work towards extending scheduling algorithms to take into account features of real-life multiprocessing workload, and we expect a number of possible directions for extending them.

#### 4.1.1 Model and problem statement

We describe some motivating scenarios in parallel databases and scientific computation later in §4.2. The model and problems we abstract from there are as follows.

**Model.** The multiprocessor consists of m identical processors and s types of other resources not including the processors. It is not necessary to maintain the identity of programs separate from jobs: the arrival of a program is equivalent to the arrival of some number of jobs, specifying some precedence graph between them. Thus we will consider the unit of computation to be a job. A job j will need several kinds of resources. A resource is malleable if there is a range of trade-offs between the amount of that resource allocated to the job and its running time. It is non-malleable if the job has to reserve a fixed amount of the resource for as long as it runs, no matter what that time is.

A job arrives with the following information:

• The trade-off between processor and time. This is expressed by a running time function,  $t_j(\mu)$ , on  $1 \leq \mu \leq m$  processors, where m is the number of processors in the system. In general, the running time  $t_j$  would be a function of *all* malleable resources. Our case studies seem to indicate that only one resource is really malleable: processors. Once the processor allocation is fixed, we shall see that all other resources essentially become non-malleable.

We shall study various forms of the trade-off function. In the most general setting we will assume that  $t_j()$  are arbitrary computable functions; in this case we say the processor resource is *malleable*. In a simpler setting, we will assume  $t_j(\mu) = t_j m_j/\mu$ , where j shows linear speedup up to  $m_j$  processors, on which its running time is  $t_j$ ; in this case we shall say the processor resource is *perfectly malleable*, up to a limit. Some programs are not written to be adaptive to any number of processors. For jobs generated by such programs, processors are a *non-malleable* resource. Job j will run on exactly  $m_j$  processors for time  $t_j$ . We will round  $t_j$ 's to powers of 2 for convenience, and also denote  $T = \max_j \{t_j\}/\min_j \{t_j\}$ . We will overload  $t_j$  to represent both a fixed number and a function; the meaning will be clear from context.

• Precedence constraints: a set of jobs (that have already been submitted to the scheduler) that must complete before j can start. This induces a partial order  $\prec$ 

on the jobs. Note that if jobs arrive on-line, then  $\prec$  may be revealed to the scheduler gradually.

- Apart from the processors requirement, jobs may also specify a vector of other non-malleable resources. We will let there be s distinct types of resources, where we can assume s = O(1). E.g., if we are modeling processors, memory and IO bandwidth as resources, s = 2. The resource vector will be denoted r<sub>j</sub> = (r<sub>j1</sub>,...,r<sub>jk</sub>,...,r<sub>js</sub>), of the fractions r<sub>jk</sub> of resource of type k demanded by job j, k = 1,...,s. For notational simplicity we scale the total available resource to be 1 unit for each type. For the reasons above we keep the processor resource separate from all others.
- Job j specifies a positive priority  $w_j$ , also called its weight. The higher this number, the more profitable it is to finish it early.

**Problem.** A scheduler is an algorithm that specifies for each processor and time slot, a portion of at most one job to be executed, so that all jobs get executed subject to all the constraints above. Consider a set of n jobs. Let a scheduler complete job j at time  $C_j$ . The two performance metrics that we seek to minimize are the makespan,  $\max_j C_j$ and the weighted average completion time (WACT),  $\frac{1}{n} \sum_j w_j C_j$ . Other possible metrics are mentioned in §4.6, some of those are provably much harder than either makespan or WACT.

We impose further constraints on the schedulers. First, preemption is not allowed. Time-slicing and preemption of space-shared resources is very expensive because (1) the state has to be evicted to slower layers of the memory hierarchy, (2) processes have to synchronize and switch across protection domains, and (3) in-flight messages have to be flushed and reinjected [50, page 6]. Anecdotal evidence suggests that many practitioners switch off time sharing for production runs on parallel machines that do not permit creating dedicated space partitions. To recover from pathological cases, some machines (like the Cray T3D) have an expensive roll-in/out mechanism, but it is important to minimize its deployment. Second, the number of *types* of non-malleable resources, *s*, can be assumed to be a small constant. Resources are not private to processors; they are equally accessible by all jobs. Typically the set of resources will be memory, disk and network bandwidth (i.e., s = 3). IO and network bandwidth are often centralized from the user's perspective. With respect to memory, our model is closer to SMP's than distributed memory machines. Third, it suffices for our scheduler to be a sequential algorithm since in both databases and operating systems these decisions are typically made by a front-end processor connected to clients. Fourth, the precedence graph may be thought of as a collection of trees or series parallel graphs only; these are the most prevalent instances in applications (see §4.2). Finally, the scheduler can collect jobs over some time window but cannot wait for all jobs to be submitted, as jobs arrive on-line over time. Moreover, each arriving job specifies only its predecessors at the time of its arrival, but not its successors.

#### 4.1.2 Discussion of results

Many special cases of our resource scheduling problem are strongly  $\mathcal{NP}$ -hard, even for makespan. Thus the goal is to find approximations for the worst case and heuristics in practical settings.

#### 4.1.2.1 Makespan lower bound

We first show that the resource scheduling problem has somewhat different character from classical makespan results for job graphs. In fact, theoretical results show that if precedence is combined with non-malleable resource constraints, there exist workloads that make common scheduling techniques perform arbitrarily badly. Feldmann *et al* show that if  $t_j$  is not known before a job completes, no on-line algorithm can give a makespan within a factor better than n, the number of jobs, of the off-line optimal [51]. Even if  $t_j$  is known upon arrival, Garey and Graham showed that any greedy list-schedule has worst case makespan performance ratio at least nT/(n+T), which is roughly T when  $T \ll n$  and n when  $T \gg n$  [57, Theorem 1]. No algorithm is needed to achieve these bounds. The status of clairvoyant non-greedy algorithms was previously unknown.

Essentially all known job graph scheduling results depend on two bulk parameters of the input graph: the volume and the critical path. To simplify the discussion suppose there is only one non-malleable resource. The volume V is the sum, over all jobs in the graph, of their resource-time product; the critical path  $\Pi$  is the earliest possible completion time assuming unlimited resources. Clearly  $\Omega(V + \Pi)$  is then a lower bound to makespan, and in most existing settings this can be achieved to a constant factor. In our problem, some parameter other than V and  $\Pi$  is at work. Specifically, we give instances of our problem where no schedule can achieve a makespan smaller than  $\Omega(V + \Pi \log T)$ .

Reference	#	Job	Malleable	Non-malleable	Precedence	On-line/	Makespan
	$\operatorname{procs}$	type	processor	processor	$\prec$	off-line	/ WACT
Garey et al [57]	N/A	N/A	×	$\checkmark$	Ø	on	makespan
Feldmann et al [51]	m	$\mathbf{par}$	$\checkmark$	×	any	on	makespan
Hall et al [70]	1, m	seq	×	×	any	off	both
	1, m	$\mathbf{seq}$	×	×	Ø	on	both
This chapter [29, 28]	m	par	$\checkmark$	×	any	on	both
	m	$\mathbf{par}$	×	$\checkmark$	Ø	on	both
	m	$\mathbf{par}$	$\checkmark$	$\checkmark$	any	on	makespan
	m	$\mathbf{par}$	$\checkmark$	$\checkmark$	any	$_{ m off}$	WACT
	m	$\mathbf{par}$	$\checkmark$	$\checkmark$	forest/SPG	on	both

Table 4.1: Comparison of results. N/A=not applicable, seq=sequential, par=parallel, SPG=seriesparallel graph. A SPG is expressed recursively as follows: every SPG is a single node or a DAG with a source and sink; an SPG is either two SPG's with an edge from the sink of one to the source of the other; or it comprises a new source, with edges to the sources of two SPG's, and edges from their sinks to a new sink.

#### 4.1.2.2 Makespan upper bound

Certain special cases of the resource scheduling problem had previously known approximation algorithms; these are discussed below and summarized in Table 4.1.

No precedence. If the precedence graph is empty, (i.e., jobs are independent) then a number of approaches are known to get approximation bounds [57, 87, 113, 112, 111]. In the database scenario, it is possible to collapse each query, consisting of several jobs with a precedence relation among them, into one job, i.e., allocate maximum resources over all the jobs in the query, and then apply the results for independent jobs [119]. This has a serious drawback in that some obvious, critical co-scheduling may be lost. For example, a CPU-bound job from one query and the IO-bound job of another can be co-scheduled and it is highly desirable to do so [73]; this cannot be done after collapsing the query.

**Only a malleable resource.** If there are no non-malleable resources but only a malleable resource, then precedence can be handled (in the sense of approximating makespan) even by a scheduler that does not know  $t_j$  before a job finishes [51].

**Small resource demand.** Another possible restriction is to allow non-malleable resources, but require each job to reserve no more than  $\lambda$  fraction of the non-malleable resources, where  $\lambda$  is small. That is, the maximum fraction requested by a job is at most  $\lambda$ . In that case naive approaches will work well. The approximation ratio for greedy scheduling

can be easily shown to be  $1 + \frac{1}{1-\lambda}$ . This is however useless even if one, or a *few* jobs need a large fraction, or equivalently, if  $\lambda$  approaches 1. In fact,  $\lambda = 1 - O(1/m)$  suffices to render the greedy schedule useless. Intuitively, a few resource-intensive jobs can delay a convoy of tiny jobs.

We show that we can attain the existential lower bound on the makespan, for the above choice of parameters  $(V, \Pi \text{ and } T)$ . We give a simple approximation algorithm that matches the  $O(V + \Pi \log T)$  makespan bound. In contrast to our algorithm, most known practical solutions use some variant of greedy list- or queue-type scheduling [50, 59, 90]. Jobs on arrival are placed in a list ordered by some heuristic (often FIFO). The scheduler dispatches the first ready job on the list when enough resources become available. List scheduling and its variants are appealingly simple to implement, but they can be notoriously bad, which is not surprising given the negative results mentioned above.

#### 4.1.2.3 Weighted average completion time

We extend the makespan upper bound to the weighted average completion time metric. For this we use a recent elegant framework by Hall *et al* for optimizing WACT, which we also show to be nearly optimal for makespan. The framework needs two subroutines: a knapsack-type routine for picking a set of "high-returns" jobs to run next, and a makespan algorithm to run them. By designing these subroutines, we give the first logarithmic approximation algorithm for our resource scheduling problem using the WACT metric. For several slightly simpler variants, the logarithmic factor can be reduced to a small constant. Many of these on-line results are the first constant approximation algorithms for the corresponding problems, on-line or off-line, while others are on-line algorithms whose performance is close to the best known off-line results. These ideas can be adapted to yield the first known (off-line or on-line) constant-approximation algorithms for minsum open shop scheduling and minsum job shop scheduling with a fixed number of machines.

Previous work on the WACT metric gave either non-clairvoyant, preemptive solutions for sequential jobs or jobs that used a perfectly malleable processor resource, with job precedence [41]; or clairvoyant, non-preemptive solutions for independent jobs with only non-malleable resource, if any [92, 112]. Apart from algorithm design, we believe it is important to point out some differences between existing scheduling literature and features needed by schedulers in parallel computing systems. We also remark that although our problem is different from existing theoretical settings, our solution borrows from various existing techniques [57, 103, 29, 70].

In §4.2 we study database and scientific computing scenarios to justify our model. In §4.3 and §4.4 we give the makespan lower and upper bounds. In §4.5 we show how to extend the makespan algorithm to a WACT algorithm. In §4.6 we pose some unresolved problems.

# 4.2 Motivation

#### 4.2.1 Databases

Query scheduling in parallel databases is a topic of active research [23, 90, 119, 66, 59]. Queries arrive from many users to a front-end manager process. A query is an in-tree, where the internal vertices are operations like sort, merge, select, join etc., which we call jobs. (We think of pipelines as collapsed into single vertices.) The leaves are relations stored on disk. Edges represent data transfer; the source vertex sends output to disk, which the target vertex later reads. Queries may have a priority associated with them, e.g., an interactive transaction has high priority and statistics collection has low priority.

Databases keep certain access statistics along with the relations, which are used to predict the size of the result of a join or a select and how many CPU instructions will be required to compute these results. The tools are standard in database literature [101]. For parallel databases, one can also estimate for each operation the maximum number of processors that can be employed for near-linear speedup [73]. Thus  $t_j$  and  $m_j$  can be estimated when a job arrives. Estimates of sizes of intermediate results can be used to estimate the memory and disk bandwidth resource vector  $\vec{r_j}$ .

The running time of a job is roughly inversely proportional to the number of processors in the range  $[1, m_j]$ , but not the total available memory. For example, a hash-join between two relations  $R_1$  and  $R_2$  with, say,  $R_1$  being smaller, takes time roughly proportional to  $\lceil \log_r |R_1| \rceil$ , where r is the memory allocated; typically the query planner picks  $r = |R_1|$  or  $r = |R_1|^{1/2}$ , independent of other queries [90]. Once processor and memory allocation are fixed, the disk bandwidth requirement can be estimated from the total IO volume and job running time.

This model is best suited to shared memory databases running on symmetric

multiprocessors (SMP) with shared access to disk [73]. They currently scale to 30–40 processors. There is growing consensus that SMP's and scalable multiprocessors will converge to networked clusters of SMP nodes [66]. Since communication costs across clusters is much more expensive than shared access within a cluster, the expectation is that most queries will be parallelized within an SMP node.

#### 4.2.2 Scientific applications

Multiprocessor installations are shared by many users submitting programs to manager processes running on the front-end that schedules them. Examples of front-end schedulers are DJM (distributed job manager) on the CM5, NQS (network queuing system) on the Paragon, POE (parallel operating environment) on the SP2. Users may submit a script to the manager. A script file has a sequence of invocations of executables, each line typically specifying a priority, the number of processors, estimated memory and running time. Notice that although there may be some flexibility in the amount of memory needed, the user typically specifies a fixed choice to the scheduler, which has to regard it as inflexible. To improve utilization, system support has been designed to express jobs at a finer level inside an application and convey the information to the resource manager by annotating the parallel executable [65, 48, 91].

The common precedence graphs are chains for scripts, series-parallel graphs (defined in Table 4.1) for structured programs, forests for database queries, and trees for divide-and-conquer and branch-and-bound algorithms.

#### 4.2.3 Fidelity

The most general representation of a job is its running time as a function of the resources allocated to it. It is difficult to find this function [59], and unclear if it is simple enough to be used by the optimizer. We group the resources into two types: malleable and non-malleable. We handle only one malleable resource that affects the running time. It may be of interest to evaluate more elaborate alternatives. We assume that  $t_j$  is known (constant or function), which is reasonable for our target domain, but for less predictable applications, some type of adaptive scheduling is needed. Even though virtual memory appears to make memory a malleable resource, most parallel systems that support paging (including the SP2 or the Paragon, and excluding the CM5) strongly encourage users to

write programs that fit in physical memory. Paging non-cooperatively on a multiprocessor can be disastrous for performance.

We have tried to avoid both unwarranted simplifications and gratuitous generality. For example, we have not assumed for simplicity that all resources are malleable. Memory is clearly not malleable, even for programs that are adaptive in limited ways to the amount of allocated memory. On the other hand, we have not modeled the details of the inter-processor network.

### 4.3 Makespan lower bound

Many results in DAG scheduling rely on two bulk parameters of the workload. One, which we call the *volume* parameter, is the sum of resource-time products over jobs in the graph, denoted V. For example, consider jobs with only one resource type: processors. If each job runs on only one processor, as in list-scheduling [64], then the fractional resource every job occupies is  $r_j = 1/m$ , where there are m processors in all. If the processor resource is perfectly malleable up to limit  $m_j$ , or non-malleable, fixed at  $m_j$  for job j, then the fractional resource job j occupies is  $r_j = m_j/m$ . In all cases we define  $V = \sum_j t_j r_j$ . The other parameter is the critical path  $\Pi$  of the graph.  $\Pi = \max_j \Pi_j$ , where  $\Pi_j$  is the minimum time at which j can complete, if infinite resources were available.

 $\Omega(V + \Pi)$  is a lower bound to makespan. If the problem is relaxed to remove either precedence or non-malleable resources, then this lower bound can be achieved to a constant factor:  $\Pi + \frac{1}{m} \sum_j t_j = \Pi + V$  for sequential jobs [64], and  $\Pi + (\frac{\sqrt{5}-1}{3-\sqrt{5}}) \frac{1}{m} \sum_j t_j m_j = \Pi + O(V)$ for malleable jobs [51]. These depend on arguments of the form: "if a critical path is being ignored, most of the resources are being utilized."

In this section, we show that  $O(V + \Pi)$  makespan is not always possible with both non-malleable resources and precedence constraints. Thus this differentiates our problem from list-scheduling [64, 57] and malleable job scheduling [51]. Our formulation is different in flavor because jobs may have to wait even when resource utilization is very low because their specific requirement of the non-malleable resources are not met. In contrast, jobs can proceed with a smaller amount of a malleable resource with proportionate slowdown. Thus some parameter other than V and  $\Pi$  is at work.

Claim 4.3.1 With precedence and non-malleable resources, the optimal makespan can be

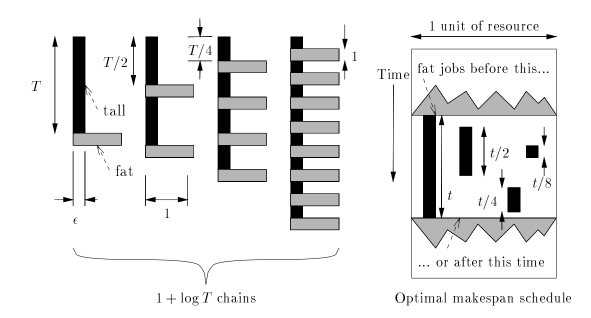


Figure 4.1: Illustration of the makespan lower bound.

as large as  $\Omega(V + \Pi \log T)$ , where  $T = \max_{i} \{t_i\} / \min_{i} \{t_i\}$ .

**Proof.** Consider the following instance with one non-malleable resource dimension. There are  $1 + \log T$  independent job chains. Chain *i* has  $2^i$  sequential compositions of the following two-job chain: the *tall* predecessor has  $t_j = T/2^i$  and  $r_j = \epsilon$  (which is a small number we will pick shortly) and the *fat* successor has  $t_j = r_j = 1$ . Then it can be verified that  $\Pi = O(T)$ , and that  $V = \sum_i 2^i (\epsilon T/2^i + 1) \le \epsilon T \log T + 2T = O(T)$ , when we choose  $\epsilon \le 1/\log T$ .

Note that only tall jobs can run concurrently; fat jobs cannot run concurrently with each other or with tall jobs. Because of the dependencies, the optimal schedule can run at most one job from each chain in parallel. Also note that the total length of tall jobs is  $\Theta(T \log T)$ . We will account for the makespan of an optimal schedule by iteratively picking the currently largest job j in the schedule, say of length  $t_j = T/2^x$ . No larger jobs are present at this point. Thus this job only overlaps with smaller jobs, but at most one of each distinct length. Remove j and all these jobs from consideration. This reduces the makespan by  $T/2^x$ and the total remaining tall job length by at most  $T(1/2^x + 1/2^{x+1} + \cdots + 1) \leq 2T/2^x$ . Note that the fat jobs need to be only over  $1 - 1/\log T$  wide in the resource dimension.

We shall use shop-scheduling techniques in the next section to get a logarithmic approximation for this problem, but we know of no similar lower bound instance for shop

problems. Specifically, a shop problem instance has some number of jobs; each job j consists of some *operations*; the *i*-th operation must run for time  $t_{ji}$  on a specified machine  $m_{ji}$ ; for each job, no two of its operations may run concurrently. Let  $\Pi = \max_j \{\sum_i t_{ji}\}$  and  $V = \max_m \{\sum_{m_{ji}=m} t_{ji}\}$ , then  $\max\{\Pi, V\}$  is the only known lower bound on the shop makespan based on bulk properties of the instance; this seems to indicate that the resource scheduling problem is somewhat different from shop-scheduling.

Claim 4.3.1 also implies that a recent elegant constant factor WACT approximation technique due to Chekuri *et al*, which converts uniprocessor schedules to multiprocessor schedules [34], will not generalize to handle non-malleable resources. Their technique is to start with a uniprocessor schedule, where job j completes at time  $C_j^1$ , and derive an m-machine schedule with  $C_j^m = O(C_j^1/m + \Pi_j)$ . In their model all jobs are sequential and the jobs need no other resources. If we additionally add jobs that request non-malleable resources, their conversion does not go through, and with good reason: if it did, a schedule with  $O(\Pi)$  additive term, rather than the  $\Omega(\Pi \log T)$  term that we showed above, will hold, thus violating the lower bound.

## 4.4 Makespan upper bound

In this section we will give an algorithm Makespan that is polynomial in s, T and n that will achieve a makespan of  $O(Vs + \Pi \log T)$  for an input set of jobs J with the bulk parameters V,  $\Pi$  and T defined before, where there is one malleable resource and there are s types of non-malleable resources. Let  $\vec{v}_j = t_j \vec{r}_j$ , and  $V = \max\{\frac{1}{m} \sum_j m_j t_j, \|\sum_j \vec{v}_j\|_{\infty}\}$ . Also let  $\Pi_j$  be the critical path length from a root through job j, and  $\Pi = \max_j \{\Pi_j\}$ .

The makespan algorithm first invokes the malleable scheduling of Feldmann et al [51] to allocate processors to jobs in J, and to assign (infeasible) preliminary execution intervals to these jobs that still violate other resource constraints. Then we partition the jobs into a sequence of *layers* with jobs within a layer being independent of each other. Finally we schedule these layers one by one using bin-packing.

Step 1. Let  $\frac{1}{2} < \gamma < 1$  be a free parameter to be set later. Compute a greedy schedule for J ignoring all non-malleable resource requirements, as follows. Whenever there are more than  $\gamma m$  free processors, schedule any job j in J (whose predecessors have all completed) on the minimum of  $m_j$  and the number of free processors [51].

Denote by  $\mu_j$  the number of processors allocated to job j. After processor allocation let the modified job times be  $t'_j = t_j m_j / \mu_j$ , and modified critical path lengths be  $\Pi'_j$ . At this stage the non-malleable resource requirements may not be satisfied.

Step 2. Round all job times to powers of two: first scale up the time axis by a factor of two, then round down each job to a power of two. Do not shift the job start times, so that all precedences are still satisfied. We will show that after this step, for all  $j, t'_j = O(t_j)$  and  $\Pi'_j = O(\Pi_j)$ . For notational convenience we will continue to refer to the modified quantities as  $t_j$  and  $\Pi_j$ , and assume that the modified time  $t'_j$  is a power of two, with  $\min_j t'_j = 1$  and  $\max_j t'_j = T$ , all this affecting only constants.

Step 3. Partition the jobs by Dividing their earliest possible start times into blocks of length T; i.e., let  $B_{\tau} = \{j : \tau T \leq \Pi_j - t_j < (\tau + 1)T\}$ . Each block of length T will be expanded to a sequence of layers of total length  $O(T \log T)$ . Specifically, remove from  $B_{\tau}$  all jobs of length T and schedule them in a layer of length T. This is the last layer. Divide  $[\tau T, (\tau + 1)T)$  into  $[\tau T, (\tau + \frac{1}{2})T)$  and  $[(\tau + \frac{1}{2})T, (\tau + 1)T)$  and recurse, placing the generated layers in pre-order [103]. The total length of the schedule, which may still violate non-malleable resource limits, will be  $O(\Pi \log T)$ .

Step 4. Schedule each layer separately in time order. Consider each layer to be an instance of a generalized s-dimensional bin-packing problem<sup>1</sup> In this problem, studied by Garey and Graham [57], and Garey, Graham, Johnson and Yao [58], there are items to pack into bins; each item e is an s-dimensional vector  $\vec{r_e}$  with components in (0, 1), and the bin is an s-dimensional with all components set to one. A set E of items fits in a bin if  $\sum_{e \in E} \vec{r_e} \leq \vec{1}$ , where  $\leq$  is elementwise. A first-fit (FF) bin packing of each layer suffices for our purpose.

**Lemma 4.4.1** After Step 2, the modified times and critical paths obey  $t'_j = O(t_j)$  and  $\Pi'_j = O(\Pi_j)$ . Furthermore, the length of the schedule is  $O(\frac{1}{m}\sum_j m_j t_j + \Pi)$ .

**Proof.** (Sketch) The former claim follows because each job j is assigned either  $m_j$  machines, in which case  $t'_j = t_j$ , or at least  $\gamma m$  machines, in which case  $t'_j \leq t_j/\gamma$ . Picking  $\gamma$  such that  $(1 - \gamma)(1 + \frac{1}{\gamma}) = 1$ , the length of the (invalid) schedule is at most  $\Pi + \frac{1}{m\gamma} \sum_j m_j t_j$ , similar to [51].

<sup>&</sup>lt;sup>1</sup>However, this problem is not one where items are solid blocks with volume and the bin is a hollow unit cube.

**Lemma 4.4.2** Jobs assigned to a particular layer I are independent; the layer ordering is consistent with job precedence  $\prec$  and in any layer I,  $\sum_{j \in I} \mu_j \leq m$ .

**Proof.** At every stage in the recursion, consider the jobs of length t removed from the block of length t (and put in a separate layer of length t). Any pair of such jobs must have overlapping "execution" intervals after the first step. They must therefore have been independent, and the sum of processors assigned to them was at most m in the first step. A job of length t starting in a block of length t can only have (smaller than t) predecessors starting in the same block and no successors starting in the same block. When the block is bisected, these predecessors are all completed before any of the t-long jobs in this block starts.

Note moreover that every job j placed in a layer I of length t(I) has  $t_j = \Omega(t(I))$ . We will need the following observation which follows from a pigeon-hole argument.

**Lemma 4.4.3** For any set  $\{\vec{v}\}$  of s-dimensional vectors,  $\|\Sigma\vec{v}\|_{\infty} \geq \frac{1}{s}\Sigma \|\vec{v}\|_{\infty}$ .

**Theorem 4.4.4** For s resource dimensions the above algorithm obtains a makespan of  $O(Vs + \Pi \log T)$ .

**Proof.** Consider a layer I that is t(I)-long in time, and generates f(I) + 1 bins using FF. Then there is at most one bin that is less than half-full in all s dimensions. Each of the other f(I) bins are at least half-full in at least one dimension. Call these bins  $F_{1/2}(I)$ . For a bin b define  $\vec{v}_b = \sum_{j \in b} \vec{v}_j$ . Then for all  $b \in F_{1/2}(I)$ ,  $\|\vec{v}_b\|_{\infty} \geq \frac{1}{4}t(I)$ . Hence we obtain  $V \geq \|\sum_b \vec{v}_b\|_{\infty} \geq \frac{1}{s} \sum_b \|\vec{v}_b\|_{\infty} \geq \frac{1}{s} \sum_I \sum_{b \in F_{1/2}(I)} \|\vec{v}_b\|_{\infty} \geq \frac{1}{4s} \sum_I f(I)t(I)$ . The total length of the schedule is thus at most  $\sum_I t(I)(f(I) + 1) \leq 4Vs + O(\Pi \log T)$ .

Observe that the makespan routine is polynomial in n, T, and s, and works for any precedence. The above method also gives an alternative algorithm and much simpler analysis (weaker only in a constant) for the (s + 1)-approximate resource constrained scheduling result of [57]. Their (s + 1)-approximation is for  $\prec = \emptyset$ . In this case  $\Pi = T$ , and we allocate log T layers with  $t(I) \in \{1, 2, 4, \ldots, T\}$ . Then  $\sum_{I} t(I) < 2T$ , giving an O(s)approximation.

**Corollary 4.4.5** If  $\prec = \emptyset$ , the above analysis (with a trivial Step 3) gives a schedule of length O(Vs + T).

We do not know the status of the gap between the upper and lower bound w.r.t. the factor of s.

# 4.5 Weighted average completion time

In this section we will describe how to extend the makespan algorithm developed earlier to the weighted average completion time (WACT) metric. We first justify why this metric is of interest. Makespan algorithms were designed for batch operations, which means that only the maximum completion time  $\max_j \{C_j\}$  was of concern. In parallel computer installations, jobs are submitted by different parties over time, are collected over some windows and then scheduled with some priorities. One good metric is *response time*, which over an arbitrary set of test jobs can be denoted  $\sum_j (C_j - a_j)$ , where  $a_j$  is when job j is submitted and  $C_j$  is the completion time. Unfortunately this is a rather intractable metric in the worst case, but in practice the following strategy works well: collect jobs in suitable time windows, assign priorities to jobs based on how long they have been waiting, and then run a priority-based batch scheduling algorithm to select which jobs to do next. The Operating Systems research literature has many proposals for choosing the priority function to favor starved or short jobs [54].

We will thus be interested in designing subroutines that input a set of jobs that have arrived but not been scheduled yet, and constructs a schedule with near-minimal  $\sum_j w_j C_j$ , where  $w_j$  is the priority assigned to job j and  $C_j$  the completion time of job jmeasured from the current time. This metric is called weighted average completion time (WACT). For a single processor and no other resources, "shortest remaining processing time" is optimal for WACT, given independent jobs. In some sense, we show how to generalize this intuition to multiple processors, multiple resource types, and jobs with precedence.

#### 4.5.1 The bicriteria framework

For a single processor, any optimal WACT schedule has optimal makespan. Surprisingly, this does not hold for unrelated parallel machines, on which a job can run at various speeds on various machines, and the speed of a machine is not independent of the job. For unrelated machines, there are instances where any optimal WACT schedule for n jobs has makespan  $\Omega(\log n/\log \log n)$  times the optimal makespan [74].

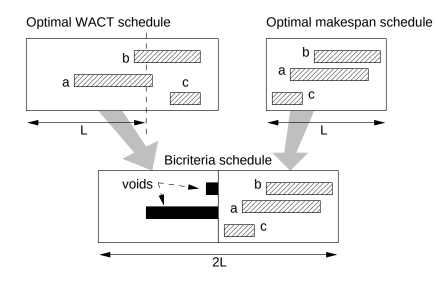


Figure 4.2: Illustration of existence of schedules that are good in both the makespan and WACT metric.

By relaxing the requirement on WACT from optimal to a constant factor, we show in Theorem 4.5.2 that there are schedules optimal for both WACT and makespan to within constant factors [29], even for unrelated machines and jobs with precedence. The basic idea is to truncate the optimal WACT schedule at the optimal makespan point, collect the unfinished jobs and schedule them in a second round of length at most the optimal makespan; we omit the proof and instead show it in a picture (Figure 4.2).

**Definition 4.5.1** Given a set of jobs, an  $(a_1, a_2)$  schedule is one that has makespan at most  $a_1$  times the optimal makespan, and WACT at most  $a_2$  times the optimal WACT, of the given set. Note that the two optimal schedules are in general different.

**Theorem 4.5.2** For any scheduling problem, there exists a (2, 2)-schedule.

The constants have since been improved to (2, 1.8) [108]. However, note that the proof is not constructive, since we need the optimal schedules for makespan and WACT for the construction. How can we get constructive polynomial-time computable  $(a_1, a_2)$ -schedules for small constants  $a_1$  and  $a_2$ ?

Hall *et al* suggest a framework to achieve small  $a_2$  [70]; their framework is actually also good for makespan [29]. First divide time into geometrically increasing intervals. That

is, define  $\tau_0 = 1$ ,  $\tau_\ell = 2^{\ell-1}$ ,  $\ell = 1, \ldots$  In what follows, consider the  $\ell$ th interval in time, namely,  $(\tau_{\ell-1}, \tau_{\ell}]$ ; other intervals are processed similarly.

Step 1. Let  $J_{\ell}$  be the set of jobs that have arrived within time  $[1, \tau_{\ell}]$  but have not already been scheduled. We remove from consideration any job which cannot be scheduled within the  $\ell$ -th interval because of dependence and consequent critical path length. Compute the earliest possible finish time  $\Pi_j$  of each job j based on critical path (assuming unlimited resources and processors). Any job j for which  $\Pi_j > \tau_{\ell}$  is removed from  $J_{\ell}$  and deferred to a later interval.

**Step 2.** In this step, we choose a suitable subset of  $J_{\ell}$  to be scheduled, carrying a possibly empty remainder forward to the subsequent intervals.

The subset of  $J_{\ell}$  chosen for scheduling is described as follows. From  $J_{\ell}$  we have to pick a subset  $J_{\ell}^*$  such that  $\sum_{j \in J_{\ell}^*} t_j m_j \leq m \tau_{\ell}, \sum_{j \in J_{\ell}^*} t_j \vec{r_j} \leq \tau_{\ell} \vec{1}$ , for any jobs  $j_1, j_2, j_1 \prec j_2$ and  $j_2 \in J_{\ell}^*$  implies  $j_1 \in J_{\ell}^*$ , and the objective  $w(J_{\ell}^*) = \sum_{j \in J_{\ell}^*} w_j$  is maximized. Suppose the optimal value of the objective above is  $W_{\ell}^*$ . Since the above problem is  $\mathcal{NP}$ -hard, we will instead obtain a subset  $J_{\ell}'$  with value at least  $W_{\ell}^*$  closed under  $\prec$ , and satisfying  $\sum_{j \in J_{\ell}'} t_j m_j = \pi m \tau_{\ell}$ , and  $\sum_{j \in J_{\ell}'} t_j \vec{r_j} \leq \pi \tau_{\ell} \vec{1}$ , for some factor  $\pi$ . We will describe this  $\pi$ -approximate procedure<sup>2</sup>, called **DualPack**, in §4.5.2. We postpone the jobs in  $J_{\ell} \setminus J_{\ell}'$  to a later interval.

Step 3. In this step we schedule the jobs in  $J'_{\ell}$  using a suitable  $\kappa$ -approximate Makespan subroutine which will find a schedule of length  $\kappa \tau_{\ell}$ . We schedule the output of Makespan in the interval  $[\kappa \pi \tau_{\ell}, \kappa \pi \tau_{\ell+1})$ .

The following analysis of the geometric series framework for WACT was proposed by Hall *et al* [70]. We repeat this for later reference, and give a proof that it is also good for makespan.

**Theorem 4.5.3** Given a  $\kappa$ -approximation for the makespan problem and a  $\pi$ -approximation to the knapsack problem, one can constructively and deterministically find a  $(4\kappa\pi, 4\kappa\pi)$ -schedule.

<sup>&</sup>lt;sup>2</sup>Notation: a  $\rho$ -approximate minimization algorithm returns an objective function that is at most  $\rho$  times optimal in the worst case.

**Proof.** (Sketch) Consider  $J_{\ell}$  and  $J'_{\ell}$  as in the algorithm. Let  $w(J'_{\ell})$  be weight of the jobs in  $J'_{\ell}$ . Say an optimal schedule completes within time  $\tau_L$ , finishing job j at time  $C^*_j$ , and let the total weight of jobs completing in  $(\tau_{\ell-1}, \tau_{\ell}]$  be  $W^*_{\ell}$  in that optimal schedule. Notice that for all  $\ell = 1, \ldots, L, \sum_{i=1}^{\ell} w(J'_i) \ge \sum_{i=1}^{\ell} W^*_i$ . This follows from the design of **DualPack**. Now observe that because of this dominance property, the above algorithm too finishes within round L, and also that  $\sum_{\ell} w(J'_{\ell}) = \sum_{\ell} W^*_{\ell}$ . Finally, the cost of the the schedule it determines is at most  $\kappa \pi \sum_{\ell=1}^{L} \tau_{\ell+1} w(J'_{\ell}) = 4\kappa \pi \sum_{\ell=1}^{L} \tau_{\ell-1} w(J'_{\ell}) \le 4\kappa \pi \sum_{\ell=1}^{L} \tau_{\ell-1} W^*_{\ell}$ , which is at most  $4\kappa \pi \sum_j w_j C^*_j$ . Let the optimal makespan be  $C^*_{\max}$ , and  $B_{\max} = 2^K$  be the start of the interval containing  $C^*_{\max}$ . By the design of **DualPack**, all jobs will be scheduled in iteration K + 1, and will complete within time  $\kappa \pi 2^{K+2}$ .

The best expected constants are obtained by scaling the geometric series by a random variable [29]. Specifically, instead of  $\tau_{\ell} = 2^{\ell}$ , one picks  $\tau_{\ell} = 2^{-X+\ell}$ , where X is a random variable uniformly distributed in (0, 1].

**Theorem 4.5.4** Given a  $\kappa$ -approximation for the makespan problem and a  $\pi$ -approximation to the knapsack problem, one can find an expected  $\left(\frac{2\kappa\pi}{\ln 2}, \frac{2\kappa\pi}{\ln 2}\right)$  approximation for makespan and WACT.

**Proof.** (Sketch) Fix an optimal WACT schedule in which job j completes at  $C_j^*$ . Let  $B_j$  be the start of the interval  $[\tau, 2\tau)$  in which j completes. Since the intervals have random endpoints,  $B_j$  is a random variable. Its expected value is  $E[B_j] = C_j^* \int_0^1 2^{-x} dx = \frac{1}{2 \ln 2} C_j^*$ . As in Theorem 4.5.3 this means that the approximate schedule has  $E[\sum_j w_j C_j] \leq 4\kappa\pi \sum_j w_j E[B_j] \leq \frac{4\kappa\pi}{2 \ln 2} \sum_j w_j C_j^*$ . The makespan argument extends similarly.

#### 4.5.2 DualPack and applications

Using the basic framework, we propose new WACT algorithms for a variety of scheduling scenarios where jobs may have malleable and/or non-malleable resource constraints. The crux of these algorithms is the design of the two subroutines: DualPack and Makespan. The most general version of Makespan described in §4.4 will suffice in all the problems we shall study. Therefore in this section we will focus on DualPack.

Our implementation of  $\mathsf{DualPack}(J, D)$  has the same outline in each of our applications. The inputs are J, the candidate jobs for the next iteration from which we pick a subset, and D, the amount of time allocated for the next iteration. First,

we prune from J the jobs that are impossible to complete within D time units, either because of precedence constraints or because their processing time is too large. In the exposition, we shall restrict our attention to precedence constraints  $\prec$  that are out-trees; in- or out-forests and series-parallel graphs can all be handled similarly. To prune out-trees based on precedence constraints, define PathToRoot $(j) = t_j$  if j is a root, and PathToRoot $(j) = PathToRoot(parent(j)) + t_j$  otherwise. This quantity can be computed in linear time, and then jobs j with PathToRoot(j) > D can then be pruned. Second, we use a auxiliary routine Knapsack to solve an appropriate knapsack problem over the remaining jobs in order to find a set of jobs of sufficiently large weight to schedule in the next iteration.

The input to the routine Knapsack (J, S) consists of a set J of n items (jobs), where item j has weight  $w_j$  and size  $s_j$ , and a knapsack of size S; in addition, we might be given precedence constraints on the items; if  $j_1 \prec j_2$  in the precedence ordering then we forbid the packing of  $j_2$  into the knapsack unless  $j_1$  is also packed. The knapsack problem is to find the maximum weight set that can be packed into the knapsack; let the optimal value be denoted  $W^*$ . The routine Knapsack(J, S) finds a set of total weight at least  $W^*$  that has total size at most  $(1 + \epsilon)S$ , where  $\epsilon > 0$  is an arbitrarily small constant (we used  $\pi = 1 + \epsilon$ in the previous section). To achieve this, we round down each  $s_j$  by units of  $\epsilon S/n$  and then use dynamic programming as in [75].

In each of the following subsections, we will give an implementation of DualPack for a specific problem; throughout, we denote the deadline by D and the set of jobs from which we choose by J. All of these are results for both makespan and WACT; we report only the WACT result and omit proofs for brevity.

#### 4.5.2.1 Malleable jobs

We give an algorithm for malleable parallelizable jobs without precedence constraints. The best off-line performance guarantee known for the non-malleable special case, where all jobs arrive at time zero, is 8.53, due to Turek *et al* [112]. Using an idea of Ludwig *et al* [87], this can be extended to the malleable case. Our WACT algorithm handles malleable jobs, on-line job arrival and has a performance guarantee of  $12 + \epsilon$ , and if we allow randomization, a nearly identical (expected) guarantee of 8.67.

Recall that each job j specifies a running time function  $t_j(\mu)$ . We implement  $\mathsf{DualPack}(J,D)$  as follows: for each job j, we find the value of  $\mu$  such that  $t_j(\mu) \leq D$ 

for which  $\mu t_j(\mu)$  (i.e., the processor-time product) is minimized. Jobs for which there is no such  $\mu$  are removed from J; otherwise, let  $m_j$  be the value for which this minimum is attained. If job j is scheduled by **DualPack**, it will be run on  $m_j$  machines. We set the weight and size of job  $j \in J$  to be  $w_j$  and  $m_j p_j$ , respectively, and then call **Knapsack**, returning J' = Knapsack(J, mD). Finally, we use the list scheduling algorithm of Garey and Graham [57] to schedule J'.

**Theorem 4.5.5** The above DualPack routine is a  $(3 + \epsilon)$ -approximation algorithm for the maximum scheduled weight problem. This gives a deterministic on-line  $(12 + \epsilon)$ approximation algorithm for scheduling malleable jobs on parallel machines, and a randomized on-line algorithm with expected performance within 8.67 of optimal.

#### 4.5.2.2 Perfectly malleable jobs with precedence

We shall consider perfectly malleable jobs, as in Feldmann *et al* [51], and precedence constraints that are forests or series parallel graphs. Our **DualPack** routine is as follows. We remove from J any j with PathToRoot(j) > D, set  $J' = \text{Knapsack}(J_{\ell}, mD)$ , and list schedule J' as in [51]: let  $\phi = (\sqrt{5} - 1)/2$  be the golden ratio; whenever there is a job j with all of its predecessors completed and the number of busy processors is less than  $\phi m$ , schedule the job on the minimum of  $m_j$  and the number of free processors.

**Theorem 4.5.6** The above DualPack routine is a dual  $(2 + \phi + \epsilon)$ -approximation algorithm for the maximum scheduled weight problem. This gives a deterministic on-line 10.48-approximation minsum algorithm for perfectly malleable jobs with forest precedence constraints, and a randomized algorithm with expected performance within 7.58 of optimal.

#### 4.5.2.3 **Resource and precedence constraints**

After handling the special cases of independent jobs and perfectly malleable jobs, we study the general model set up at the beginning of this chapter where jobs use both malleable and non-malleable resources and have precedence constraints. We point out that throughout the previous applications, we had  $\kappa$ , the approximation factor for **Makespan**, to be a small constant, and  $\pi$ , the approximation factor for **Knapsack**, to be  $1 + \epsilon$  for  $\epsilon > 0$ arbitrarily small. As we saw in §4.3, if we can only use bulk parameters like volume and path in the knapsack routine (this is all we can do at the moment), then  $\kappa = \Theta(\log T)$  for the case where we have both non-malleable resources and precedence constraints. Thus we can use the same **DualPack** routine as for perfectly malleable jobs, but cannot use list-scheduling for the makespan routine. We instead use the algorithm in §4.4, obtaining the following.

**Theorem 4.5.7** There is an algorithm polynomial in T and n which gives an  $O(\log T)$  approximation for both makespan and WACT with on-line job arrival, assuming s = O(1).

Recall that throughout we have assumed s = O(1). In general, the algorithm above can be proved to be an  $O(s + \log T)$  approximation.

If we are only interested in off-line schedules, we can compute  $J'_{\ell}$  via rounding an integer program similar to [70], obviating the need for the DualPack routine. We omit the details of the following claim.

**Theorem 4.5.8** There is an off-line algorithm, polynomial in s, T and n, that approximates makespan and WACT to an  $O(s + \log T)$  multiplicative factor.

If there is no precedence  $(\prec = \emptyset)$ , we can use Corollary 4.4.5 to obtain the following generalization of Turek *et al*'s result [112] to many non-malleable resources.

**Corollary 4.5.9** There is an off-line algorithm, polynomial in s, T and n, that approximates makespan and WACT to an O(s) multiplicative factor.

# 4.6 Extensions

Finally, we raise several questions regarding extensions of the model and algorithms in this chapter.

**Tighter bounds.** V,  $\Pi$  and T are not the best characterization of an instance, since the optimal makespan is  $\Omega(\Pi \log T)$  for some but not all instances. A better characterization of the lower bound and improved algorithms are needed. It is not clear how such a characterization can improve the WACT algorithm.

**Imperfect malleability.** In reality the processor resource is not perfectly malleable, neither are other resources perfectly non-malleable. How important is it to model and optimize for complicated intermediate forms of malleability?

**Persistent resources.** A job may allocate memory mid-way through execution. We cannot model this by a chain of two jobs, since the memory the job was already holding is not released. How can jobs with such persistent resource needs be scheduled?

Non-clairvoyance and preemption. For the motivating applications, reasonable estimates of job running time are possible. More general purpose schedulers must be non-clairvoyant, i.e., work without knowledge of  $t_j$  before j completes [105, 92]. To handle this, recourse to job preemption or cancellation is needed, whose large cost has to be factored into the algorithm.

Arbitrary DAGs. While we have handled hierarchical job graphs such as forests or seriesparallel graphs, the general DAG case is open. It is known that precedence-constrained knapsack with general precedence is strongly  $\mathcal{NP}$ -hard [75], unlike forests. We show that settling the approximability issue will be challenging [4]. This shows that the framework of [70] may need modification to handle DAG's, not necessarily that the scheduling problem is difficult.

**Claim 4.6.1** There is an approximation preserving reduction from Expansion, the problem of estimating the vertex expansion of a bipartite graph, to P.O.K., the partial order knapsack problem, even when all item costs and profits are restricted to  $\{0, 1\}$ .

**Proof.** Given G = (L, R, E), suppose we need the vertex expansion of R. Construct a two layer partial order  $\prec$ : the upper layer contains a job j for every  $u \in L$ , with c(j) = 1, p(j) = 0. The lower layer contains a job j for every  $v \in R$ , with c(j) = 0, p(j) = 1. E is directed from L to R. Run the routine for P.O.K. n times, with target profits  $P \in \{1, \ldots, n\}$ . Return  $\min_P\{C(P)/P\}$ , where C(P) is the cost returned for target profit P.

**Flowtime.** In this chapter we consider  $\max_j C_j$  and  $\sum_j w_j C_j$ ; some more ambitious objective functions are  $\sum_j (C_j - a_j)$  and  $\sum_j w_j (C_j - a_j)$ , commonly called *flowtime*. Unfortunately, even with a single machine, off-line problem instance, and no resource or precedence constraints, it is  $\mathcal{NP}$ -hard to approximate non-preemptive flowtime better than about a factor of  $\Omega(\sqrt{n})$  [80]. In a practical implementation of our algorithm, one might artificially increase  $w_i$  for jobs waiting for a long time.

# Chapter 5

# Dynamic scheduling using parallel random allocation

# 5.1 Introduction

Thus far in the thesis we have developed scheduling algorithms for problems with one benign feature: when a job arrives at the scheduler, its resource requirements and running time can be estimated. For the scientific problem domain addressed so far, this is a reasonable assumption. However, in more dynamic and irregular programs, the scheduler may not have any knowledge of the running time of jobs or tasks prior to completion. Schedulers that produce schedules without this knowledge are called *non-clairvoyant*. There are a few techniques to handle non-clairvoyant scenarios, depending on the cost model and the optimization objective.

In the simplest non-clairvoyant model, the goal is to minimize the finish time of a dynamically growing task graph, and there is a central pool in which all ready jobs are placed. Assuming that shared access to the central pool takes zero time, Graham analyzed the makespan of the resulting schedule (called a *list*-schedule) and showed that it is less than a factor of two worse than the optimal makespan [64].

This solution has a communication bottleneck problem: all processors access the central pool. The overhead can be especially large for fine-grain tasks. One technique to reduce the overhead is to let each processor have a local task pool from which it removes and executes some task, if any. New tasks are sent to the pool of a processor chosen

uniformly at random. Karp and Zhang analyzed the performance of this strategy for unit-time tasks. In §5.2 we extend the analysis to handle arbitrary unknown task times without preemption, i.e., tasks run to completion once they are started. (If arbitrarily frequent preemption is permitted, the unknown time case is no different from the unit-time case. We mention in passing that for the weighted average completion time metric, the only known non-clairvoyant approximation algorithms use preemption every time-step.)

The chapter continues in §5.3 with the study of an extension of random allocation in a distributed setting. In the previous model, each task was assigned a random destination only once, independent of where other tasks were sent. Roughly speaking, a single round of random allocation gives a logarithmic smoothing of load. As a more precise example, when n balls are thrown independently and uniformly at random into n bins, the number of balls in the most heavily loaded bin is  $\Theta(\log n/\log \log n)$ , with probability over 1 - O(1/n). Note that balls do not need to communicate with each other, and only one round of random throwing is needed. We show how the load imbalance can be decreased as we invest more rounds of random throws (which implicitly propagates the load information).

Apart from the usual applications in distributed job scheduling, such multi-round load distribution protocols may find other uses. For example, in a serverless distributed file system, newly created data blocks have to be assigned storage on the disk at one of the workstations using the file system. To distribute blocks evenly among the disks, we may send each block to a random disk, or maintain global information about the number of free blocks on each disk. Our analysis explores the trade-off between these two extremes.

# 5.2 Random allocation for dynamic task graphs

We analyze the performance of random allocation schemes applied to irregular and dynamic task-parallel programs. Execution of the program defines a job precedence graph with vertices representing jobs and directed edges representing precedence constraints. The precedence graph is revealed on-line and is irregular in shape, and the processing times of jobs are diverse and unknown before job completion. The objective function to minimize is makespan, the maximum completion time of a job. Although the exact problem is  $\mathcal{NP}$ -hard, good approximations are possible if the algorithm assigning jobs to processors is centralized, and thus has perfect global knowledge. For example, Graham's list-scheduling algorithm [64] will result in a finish time at most 2 - 1/P times optimal, where P is the number of processors. However, this may lead to a severe communication bottleneck at the processor where the pool of jobs resides, especially for fine-grained tasks. Our goal, therefore, is to study decentralized allocation which avoids such bottlenecks.

The bottleneck can be relieved in a variety of ways, each of which reduces communication cost by sacrificing global load information and thus risking some load imbalance. We study *work sharing*, where busy processors forward jobs to random processors. Some other techniques are *work stealing*, where idle processors ask for work [19], and *diffusion*, where neighbors exchange local load information and then move some jobs from busy to lazy processors [61].

#### 5.2.1 Models and notation

The input comprises a set J of jobs presented to the algorithm in a distributed and on-line fashion. Job j has running time  $t_j$ , also referred to as its "weight" (not to be confused with "priority" as in Chapter 4). We assume these are powers of 2; this will affect the results only in constant factors. We assume that  $t_j$  can be known only when job j completes. The total work or weight in a job set J is denoted  $t(J) = \sum_{j \in J} t_j$ . The number of jobs in Jis denoted n(J). The average job weight is  $\overline{t}(J) = t(J)/n(J)$ . Let  $t_{\max}(J) = \max_{j \in J} \{t_j\}$ and  $t_{\min}(J) = \min_{j \in J} \{t_j\}$ , and let  $T(J) = t_{\max}(J)/t_{\min}(J)$ . Equivalently we scale jobs so that  $t_{\min} = 1$  and  $t_{\max} = T$ . This is in keeping with recent analyses of on-line load balancing algorithms [10], and is more broadly applicable than results with assumptions about distribution or variance.

J will have an associated acyclic precedence relation  $\prec \subset J \times J$ . Let  $\Pi_j = \max_{j' \prec j} \{\Pi_{j'}\} + t_j$  be the earliest time at which j can finish given infinitely many processors (the definition is not circular since the precedence graph is acyclic). Let  $\Pi(J) = \max_{j \in J} \{\Pi_j\}$  be the longest critical path. We assume there is a unique root job. The number of edges on a path from j to the root is denoted h(j); the path will be clear from context. Also let h(J) be the maximum number of edges on any precedence path in J.

J will be omitted when clear from context. We assume that job times and the job graph are oblivious of the decisions made by the scheduler. In exhaustive traversal, J is finite and the goal is to execute all jobs in J in any order obeying  $\prec$ . In heuristic search or branch and bound, the job graph J provided may be very large or even infinite. Each job j has an associated cost c(j), with the requirements that  $j_1 \prec j_2 \Rightarrow c(j_1) < c(j_2)$  and all costs are distinct without loss of generality. The goal of the execution is to start at the root and execute jobs obeying  $\prec$ , until the leaf node with minimum cost  $c^*$  is identified. It is not necessary to generate and execute all jobs in J.

Sequential algorithm. A common sequential strategy is the "best first" traversal. All available jobs with completed predecessors are maintained in a priority queue. While the queue is non-empty, the job j with least c(j) is removed and executed. The jobs executed are  $\tilde{J} = \{j \in J : c(j) < c^*\} \subseteq J$ . In the special case of complete traversal,  $\tilde{J} = J$ . For all models, we assume that operations on a priority queue for job selection take negligible time compared to job execution time. In this model, the sequential algorithm takes time  $t(\tilde{J})$ . We assume  $|J| \geq |\tilde{J}| \geq P$ . The interesting case for us is when  $|J| \gg |\tilde{J}|$ , so that work stealing is not an option.

**Parallel algorithm.** Parallel execution starts with the root job in one processor. A job can be started when all predecessors have been completed. When a processor completes executing a job j, all successors of j become available to that processor. Processors can negotiate to transfer available jobs among themselves.

There is no coordinated global communication for load balancing purposes. We study the setting with a local priority queue of jobs in the memory of each processor as in [78]. Thus, priority is preserved within each local queue but not across processors. An idle processor non-preemptively executes the best job from its local queue, if any. Any newly available job with completed predecessors is enqueued into the priority queue of a processor chosen uniformly at random. The destination processor is not interrupted.

**Communication.** The machine model consists of processors with individual local memory connected by a communication network. We ignore the topology of the interconnect as in the LogP model [37]. Communicating a job takes unit time at the two processors involved in the transfer.

**Pruning and termination.** In parallel branch and bound, each processor has to periodically propagate the cost of the least cost leaf it has expanded, so that all processors know the cost of the global best cost leaf in order to use it for pruning. Also, barrier synchronizations are required to detect situations where all local queues are empty so

that the processors can terminate. We note that these can be done infrequently with low overhead, so they do not affect the time bounds we derive.

#### 5.2.2 Discussion of results

For exhaustive search  $\tilde{J} = J$ , and greedy centralized schedules give a simple bound on makespan in terms of variables defined above:  $\Theta(t/P+\Pi)$ , the average work per processor plus the critical path length. We show that the situation changes somewhat in the branch and bound setting:  $\Omega(\frac{t(\tilde{J})}{P} + h(\tilde{J}) \cdot T(J))$  may be necessary even for an ideal centralized scheduler with no communication cost. Then we give an analysis of parallel branch and bound with  $\tilde{J} \neq J$  in a complete network: we show that with probability at least  $1 - \epsilon$ , the makespan is  $O(\frac{t}{P} + hT \log hT + T \log \frac{n}{\epsilon})$ , where  $t = t(\tilde{J})$ ,  $h = h(\tilde{J})$ ,  $n = n(\tilde{J})$ , and T = T(J). We also report on experience with some irregular programs. This is necessary for two reasons. First, our analysis is probabilistic and asymptotic; in practice, constant factors would be important. Second, although the above result establishes near-optimal load balance, our model does not reflect the gains from avoiding communication bottlenecks.

Other results of the diffusion type are based on occasionally matching busy and idle processors and transferring jobs [61, 98]. These are not appropriate for relatively fine-grain jobs which is our focus. Notice also that diversity in job execution times makes coordination even harder unless a processor can suspend long jobs and participate in global communication.

Work stealing is the strategy of least communication for the particular case where the job graph J is an out-tree, all jobs must be executed, and the relative order of execution is immaterial (provided it obeys  $\prec$ ). In work stealing, the graph is expanded depth-first locally in each processor, and idle processors steal jobs nearest to the root [122, 19]. In many application such as parallel search or branch and bound, the total work done is very sensitive to the job order, and one wishes to deviate from the best sequential order as little as possible [47, 8].

#### 5.2.3 Weighted occupancy

At first we consider the weighted occupancy problem, where there is no precedence among jobs. There are n weighted balls (jobs), ball j having weight  $t_j$ . These balls are thrown uniformly at random into P bins (processors). We want to bound the weight of the heaviest bin.

**Lemma 5.2.1** For random allocation of weighted balls to bins, with probability at least  $1-\epsilon$ , each bin has  $O(\frac{t}{P} + T(\log \log T + \log \frac{P}{\epsilon}))$  weight.

**Proof.** Classify the balls into weights 1, 2, ..., T, where there are  $n_i$  balls of weight  $2^i$ . Fix one bin. The probability that there are at least  $m_i$  balls of weight  $2^i$  in this bin is at most  $\binom{n_i}{m_i}P^{-m_i} \leq (\frac{en_i}{Pm_i})^{m_i}$ , which is less that  $\frac{\epsilon}{P\log T}$  for  $m_i = O(\frac{n_i}{P} + \log\log T + \log\frac{P}{\epsilon})$ . Adding over i and all P bins gives the result.

#### 5.2.4 Delay sequence

Next, we show an upper bound for random allocation. Unlike in the previous section, occupancy results cannot be used directly, since unlike a batch, a DAG schedule cannot be composed from arbitrary task subsets. Further, in analyses related to global task pools as above, arguments depend significantly on statements to the effect that during certain intervals of time, most processors do useful work. We can no longer say this when each processor has a local task pool: processors can remain idle even though there are tasks yet to expand, because they can be in the queues of other processors. We handle this using a delay sequence argument. The following lemma is similar to Ranade's construction [96].

**Lemma 5.2.2** Suppose the execution finishes at time  $\tau$ . Then the following 4-tuple  $(s, Q, R, \Pi)$  exists:

- s is a job that finished no earlier than  $\tau$ . Let  $S = (s_1, \ldots, s_{h(s)})$  be a path of "special" jobs from the root of the DAG to job  $s = s_{h(s)}$ .
- $Q = (q_1, \ldots, q_{h(s)})$  is an ordered list, where  $q_{\ell}$  is the processor that executed  $s_{\ell}$ , for  $1 \leq \ell \leq h(s)$ .
- $R \subset \tilde{J}$  is a set of jobs, and
- $\Pi_1, \ldots, \Pi_{h(s)}$  is a partition of  $[1, \tau]$  such that
  - $R \cap \{s_1, \ldots, s_{h(s)}\} = \emptyset.$
  - Each job in R become "ready" and arrives into  $q_j$  during interval  $\Pi_j$ , for some  $j, 1 \leq j \leq h(s)$ .

$$- t(R) \ge \tau - \Pi(\tilde{J}) - h(\tilde{J})T(J).$$

**Proof.** Label s as  $s_{h(s)}$  and starting at task s, move up towards the root. If a task  $s_{\ell}$  has more than one parents, go to the one that completed last of all parents, and call it  $s_{\ell-1}$ . Thus trace a path root  $= s_1, \ldots, s_{h(s)} = s$ .

To obtain R, work backwards from the time  $\tau$  and consider the latest time instant  $\tau' < \tau$  such that  $q_{h(s)}$  was empty at time  $\tau' - 1$  (we refer interchangeably to a processor and its local work pool). This means that all tasks executed by  $q_{h(s)}$  during the interval  $[\tau', \tau]$  also arrived there during this interval. Call these tasks  $R_{h(s)}$ . Include  $R_{h(s)}$  into R, and set  $\Pi_{h(s)} = [\tau', \tau]$ . Then continue the construction from  $\tau' - 1$  in an iterative manner.

From Figure 5.1 (b), it can be seen that the processing times of nodes in R must cover all of  $[1, \tau]$ , except for the time spent in processing nodes  $s_1, \ldots, s_{h(s)}$ , which is at most  $\Pi(\tilde{J})$ , and the time spent finishing jobs in progress when  $s_\ell$  arrives at  $q_\ell$ , which accounts for at most  $h(\tilde{J})T(J)$ . Thus let  $R = \bigcup_{\ell} R_\ell$  and observe that  $t(R) \ge \tau - \Pi(\tilde{J}) - h(\tilde{J})T(J)$ . We have also constructed a ordered partition  $\Pi = (\Pi_1, \ldots, \Pi_{h(s)})$  of  $[1, \tau]$ .

We also note the following fact, which follows, e.g., from considering the two cases  $b \log 2a \le a$  and  $b \log 2a > a$ .

**Lemma 5.2.3**  $x > a + b \log x$  holds for  $x > \Omega(a + b \log b)$ , where a, b > 0,

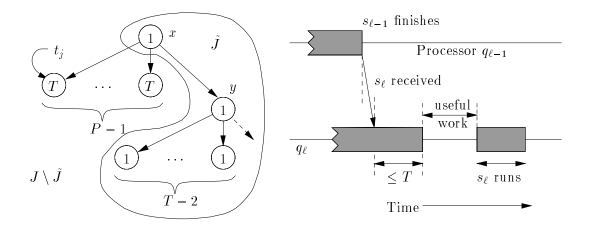


Figure 5.1: (a) A bad instance for the central scheduler. (b) Accounting for time in the delay sequence.

**Theorem 5.2.4** With probability at least  $1 - \epsilon$ , the execution time for branch and bound is  $O(\frac{t}{P} + hT \log hT + T \log \frac{n}{\epsilon})$ , where  $t = t(\tilde{J})$ ,  $h = h(\tilde{J})$ , n = n(J), and T = T(J).

**Proof.** We will bound the probability over all s, Q,  $\Pi$  and R that a 4-tuple as above will occur. Given fixed values for s, Q,  $\Pi$ , R and  $\tau$ , a conforming execution happens with probability at most  $P^{-(h(s)+|R|)}$ . Thus our target expression is  $\sum_{s,Q,\Pi,R} P^{-(h(s)+|R|)} = \sum_{s,\Pi,R} P^{-|R|}$ , since, given s, the number of choices for Q is just  $P^{h(s)}$ . In the sum  $\sum_{s,\Pi,R} P^{-|R|}$ , s can be chosen in n ways. The number of ways to pick  $\Pi$  is at most  $\binom{\tau+h}{h} \leq (6\tau)^h$ . It only remains to evaluate  $\sum_R P^{-|R|}$ , where the sum is over all R such that  $t(R) \geq \tau - \Pi(\tilde{J}) - h(\tilde{J})T(J)$ .  $\sum_R P^{-|R|}$  is the probability, over all  $R_1, \ldots, R_{h(s)}$ , that  $R_\ell$  got assigned to  $q_\ell$ ,  $1 \leq j \leq h(s)$ . This is the same as the probability that some bin gets a weight of at least  $\tau - \Pi(\tilde{J}) - h(\tilde{J})T(J)$  when n - h(s) balls were randomly assigned to P bins. This can be bounded using Lemma 5.2.1.

Specifically, setting  $\tau - \Pi - hT = \Omega(\frac{t}{P} + T \log \frac{P \log T}{\delta})$  bounds the probability of makespan being  $\tau$  to at most  $n \cdot (6\tau)^h \cdot \delta$ . By picking  $\tau$  to be suitably large,  $\delta$  can be made small. Fortunately,  $\tau$  goes up only as  $\log \frac{1}{\delta}$ , so we will be able to drive the above product down to a small probability  $\epsilon$ . We need  $n \cdot (6\tau)^h \cdot \delta < \epsilon$ , which means we need  $\frac{1}{\delta} > \frac{n(6\tau)^h}{\epsilon}$ . We can therefore use a value of  $\tau$  such that

$$\tau \geq \Omega\left(\frac{t}{P} + hT + T\log\frac{nP(6\tau)^{h}\log T}{\epsilon}\right)$$
$$= \Omega\left(\frac{t}{P} + hT + T\log\frac{n}{\epsilon} + hT\log\tau\right),$$
(5.1)

from which the result follows by using Lemma 5.2.3.

For branch and bound, this is not far away from the best possible makespan, even with a central pool and free communication.

Claim 5.2.5 With a centralized scheduler, the execution time for branch and bound is  $\Theta\left(\frac{t(\tilde{J})}{P} + h(\tilde{J})T(J)\right).$ 

**Proof.** For the lower bound we will produce a J and  $\tilde{J} \subset J$  with  $t(\tilde{J}) = o(\Pi(\tilde{J}))$  such that even a centralized scheduler will need  $\Omega(h(\tilde{J})T(J))$  time. The instance is shown in Figure 5.1(a). In the first time-step, one processor expands the root job, generating P children that all P processors start expanding at the second time-step. P-1 of these are jobs in  $J \setminus \tilde{J}$  with  $t_j = T$ , meant to keep P-1 processors busy for time T, so the last

processor is left alone to expand part of  $\tilde{J}$  as shown. In the figure, job x has P children and job y has T - 1, so that when the new set of P nodes are generated, the P - 1 processors just freed grab the new decoys. This can be arbitrarily repeated.

For the upper bound, suppose the makespan is  $\tau$  and s is a job finishing at time  $\tau$ . Label s as  $s_{h(s)}$  and starting at s, move up towards the root. If a task  $s_{\ell}$  has more than one parent, go to the one that completed last of all parents, and call it  $s_{\ell-1}$ . Thus trace a path root  $= s_1, \ldots, s_{h(s)} = s$ . Note that  $\sum_{\ell} t_{s_{\ell}} \leq \Pi(s)$ . Suppose  $s_{\ell}$  runs in the interval  $[B_{\ell}, E_{\ell}]$ . Note that  $s_{\ell}$  appears in the central job pool at time  $E_{\ell-1} + 1$ , and in the interval  $[E_{\ell-1} + T, B_{\ell}]$ , if non-empty, all processors are executing jobs inside  $\tilde{J}$  since they all picked some other jobs j' with  $c(j') < c(s_{\ell})$ , meaning  $j' \in \tilde{J}$ . Thus  $P(\tau - \Pi(\tilde{J}) - T \cdot h(\tilde{J})) \leq t$ , which proves the claim since  $\Pi = O(hT)$ .

#### 5.2.5 Empirical evaluation

Unlike in analyses of centralized schemes, our results are probabilistic and hide constants at several places. It is therefore interesting to evaluate the cost and benefit of decentralization in practical settings. We report on experiments with two applications. The first is a parallel divide and conquer algorithm to solve a symmetric tridiagonal eigenvalue problem [43]. The second is a parallel symbolic multivariate polynomial equation solver which uses a procedure similar to branch and bound [31]. Both our applications lead to tree-shaped precedence between jobs, and the job times are diverse. In both cases, most of shared data can be replicated at small communication cost, so random allocation is feasible. Random allocation with diverse times have also been use in N-body simulation [86] and integer linear programming [47].

For each application, we added instrumentation to the sequential program to emit the task tree with task times, and input this tree to a simulator that simulated the parallel execution of the randomized load balancing algorithm, as well as Graham's list schedule. By an idealized simulation without communication cost and other overheads, we first isolate and study only the loss in load balance owing to random allocation. In Figure 5.2, the eigenproblem instance has  $t = 108247480\mu$ s, n = 2999, h = 20,  $T = 184377\mu$ s÷ $4486\mu$ s≈ 41,  $\Pi = 237917\mu$ s, and  $t/\Pi \approx 455$ . The symbolic equation instance has  $t = 11053339\mu$ s, n = 142, h = 11,  $T = 174860\mu$ s÷ $1184\mu$ s≈ 148,  $\Pi = 474880\mu$ s, and  $t/\Pi \approx 23$ .

In the graphs above we have presented the speedup without explicitly measuring

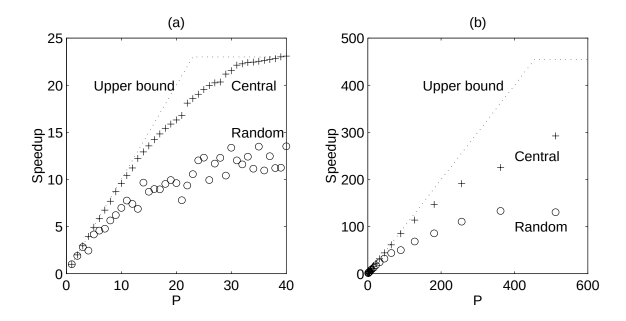


Figure 5.2: Comparison of speedup between Graham's list schedule and random allocation with zero communication cost. (a) Symbolic algebra, (b) Eigensolver.

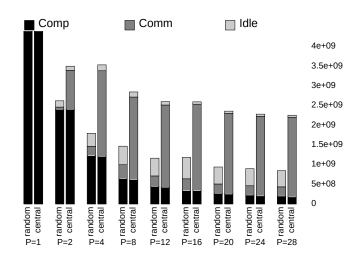


Figure 5.3: Performance comparison of Graham's list schedule and random allocation on the CM5. The y-axis represents time, broken down into computation, accessing the job pool, and idle time.

communication costs. We also measured actual speedup on the CM5 multiprocessor. Comparing the speedup curves enabled us to judge the closeness of the simulation to reality. In Figure 5.3, we present a break-up analysis of parallel running time for the eigensolver, comparing a centralized task queue with the random distributed allocation. Although load imbalance is larger for random allocation, the benefit due do decentralization is overwhelming.

#### 5.2.6 Discussions

Several questions arise from this and related results. It would be interesting to tighten the makespan estimates in this section, as well as to provide randomized lower bounds. The performance of multiple round strategies [9, 1] for dynamic job graphs remains open, as is their effect on further reducing the load imbalance. Finally, extending the results to capture network contention as in the atomic message model [86] seems like a natural goal.

### 5.3 Multi-round load balancing protocols

The goal of this section is to generalize single round random allocations to multiple rounds, aided by spreading load information, to facilitate better load balance. It is well known that when n balls are thrown independently and uniformly at random into n bins, with high probability (by which we shall mean  $1 - O(\frac{1}{n})$ ) the maximum number of balls received by any bin is  $\Theta(\frac{\log n}{\log \log n})$  [93]. Recently, an important extension of this result was proven by Azar *et al* [9]. Suppose we place the balls sequentially, one at a time; for each ball, we choose two bins independently and uniformly at random, and place the ball in the less full bin. When all the balls have been placed, the fullest bin contains only  $\Theta(\log \log n)$  balls with high probability, an exponential improvement over the simple randomized approach. Notice that in both these cases, the *expected* number of balls in a bin is one, but the *maximum* occupancy grows with n, with the obvious implication that any technique for reducing the gap between the expected and maximum occupancy may be useful for load balancing parallel jobs. For example, such logarithmic terms appeared in the analysis in §5.2.

We mention a few scenarios that motivate a detailed study of the trade-off between the investment in communication rounds and the resulting load balance. The first motivation is directly from the analysis in  $\S5.2$ , where the randomized upper bounds had logarithmic multiplicative factors over the lower bound (Theorem 5.2.4) in the context of allocating processors to fine-grain tasks in a scalable multiprocessor where the overhead of central load information is prohibitive. Here is another setting: modern computer networks often have decentralized compute-servers (bins) and client workstations issuing jobs (balls). A distributed load-balancing strategy has to assign jobs to servers. Clients are ignorant of the intention of other clients to submit jobs; contention is known only from server load. Servers are ignorant of jobs from clients that have not communicated with them. It is also prohibitively expensive for clients to globally coordinate job submissions. The primary objectives are to minimize the maximum load achieved as well as the number of communication rounds required. Reducing the number of rounds is an important goal since, in a network setting, the time to complete a round is determined by network latency, which is generally orders of magnitude higher than CPU cycle times.

Description	Rounds	Max Load	
Occupancy problem	1	$\Theta\left(\frac{\log n}{\log\log n}\right)$	
Azar et al	n	$\Theta(\log \log n)$	

The single round and *n*-round policies are extremes on a continuum, as shown in the above table. Unfortunately, the new method requires the resting place of the balls to be determined sequentially [9]. This limits its applicability in parallel and distributed settings, a major drawback when compared to the simple approach, which has one parallel round of random assignment (we will give a precise definition of *rounds* later). In this section we will explore the middle ground, which enables load-balancing algorithms to adjust their communication to the relative cost of communication and computation on the system.

### 5.3.1 The model

We describe our model in terms of balls and bins, with the understanding that "balls" refer to jobs or tasks, and "bins" refer to processors. Each of m balls is to be placed in one of n bins. (For simplicity, we shall concentrate on the case m = n. Extension to general values of m and n are analogous.) Each ball begins by choosing d bins as prospective destinations, each choice being made independently and uniformly at random (with replacement) from all possible bins. The balls decide on their final destinations using r rounds of communication, where each *round* consists of two stages. In the first stage each ball is able to send, in parallel, messages to any prospective bin, and in the second stage

each bin is able to send, in parallel, messages to any ball from which it has ever received a message. In the final round, the balls commit to one of the prospective bins and the process terminates. Messages are assumed to be of size polylog(n,m). The goal is to minimize the maximum load, which is defined to be the maximum number of balls in any bin upon completion.

#### 5.3.2 Summary of results

Our main goal is to develop and analyze a multiple round protocol for assigning balls to bins. Informally, for each round, each bin sets a *threshold* T, and accepts up to T balls in that round. Excess balls are thrown to a random destination again. Clearly, increasing T will complete the protocol in a few communication steps, but the maximum bin load will be large. On the other hand, reducing T limits the maximum load at the expense of many rounds. We will quantify this trade-off in §5.3.3.

Our analysis exploits a basic tool that expresses the relation between occupancy distributions and Poisson distributions [63, 1]. Using this tool, we show that after a fixed number of rounds r, the final maximum load is  $O\left(\sqrt[r]{\frac{\log n}{\log \log n}}\right)$  with high probability.

We also show via lower bounds that no better load balance can be achieved by a large class of load balancing algorithms. The class that we restrict our attention to are *non-adaptive*, in that the possible destinations are chosen before any communication takes place. We will also restrict our discussion to strategies that are *symmetric*, in the sense that all balls and bins perform the same underlying algorithm and all possible destinations are chosen independently and uniformly at random. We believe that these restrictions have practical merit, as an algorithm with these properties would be easier to implement. We provide a lower bound for non-adaptive and symmetric strategies. For any fixed number rof rounds of communication and any fixed number d of choices for each ball, we show that with constant probability the maximum load is at least  $\Omega\left(\sqrt[r]{\frac{\log n}{\log \log n}}\right)$ .

#### 5.3.3 Threshold protocol

Balls and bins execute the following protocol, which we call  $\mathsf{Threshold}(T)$ .

- 1. While there exists a ball that has not been assigned to a bin do the following round of steps.
- 2. In parallel, each unaccepted ball chooses a random bin and sends it a request.

3. In parallel, each bin chooses up to T requests from the current round, sends acceptances to these requesting balls, and sends rejects to the other balls of the same round.

Notice that the protocol is asynchronous: each ball maintains a count of its current round in the messages that it sends to bins. As long as a request includes the number of its current round as part of the message, messages from distinct rounds can be handled simultaneously. Moreover, balls send and receive at most one message per round. Finally, we shall show that this method demonstrates a potentially useful tradeoff between the maximum load and the number of rounds.

#### 5.3.3.1 Poisson approximation

In this section we show a relationship between occupancy and Poisson distributions. Suppose  $\ell$  balls are thrown into n bins independently and uniformly at random, and let  $X_i^{(\ell)}$  be the number of balls in the *i*-th bin, where  $1 \leq i \leq n$ . Also let  $Y_1^{(m)}, \ldots, Y_n^{(m)}$  be independent Poisson random variables with mean  $\frac{m}{n}$ . We will omit the superscript when clear from context. In this section we will derive some relations between these two sets of random variables. We note the following lemma that follows from routine manipulations.

**Lemma 5.3.1** For non-negative integers  $x_1, \ldots, x_n$ ,

$$\Pr\left[Y_1^{(m)} = x_1, \dots, Y_n^{(m)} = x_n\right] = \Pr\left[X_1^{(\Sigma_i x_i)} = x_1, \dots, X_n^{(\Sigma_i x_i)} = x_n\right].$$
(5.2)

This means in particular that for any non-negative f,

$$\mathbb{E}\left[f(Y_1^{(m)}, \dots, Y_n^{(m)}) \middle| \sum_i Y_i^{(m)} = k\right] = \mathbb{E}\left[f(X_1^{(k)}, \dots, Y_n^{(k)})\right]$$
(5.3)

Note that m disappears from the right hand side. We will also need the following facts.

**Lemma 5.3.2** For a Poisson distributed random variable Y with mean  $\lambda \geq 1$ , where  $\lambda$  takes integer values,  $\Pr[Y \leq \lambda]$  and  $\Pr[Y \geq \lambda]$  are both bounded strictly between zero and one. Specifically,  $\frac{1}{2} \leq \Pr[Y \leq \lambda] \leq \frac{3}{4}$ .

**Proof.** Let  $p_{\lambda}(k) = \Pr[Y = k] = \frac{e^{-\lambda_{\lambda}k}}{k!}$ . We need to show that  $p_{\lambda} = \sum_{k \leq \lambda} p_{\lambda}(k)$  lies between  $\frac{1}{2}$  and  $\frac{3}{4}$ . First,  $p_1 \approx 0.74$ . Next, we need to verify that  $p_{\lambda+1} \leq p_{\lambda}$ .

$$p_{\lambda} - p_{\lambda+1} = \sum_{k \le \lambda} \frac{e^{-\lambda} \lambda^k}{k!} - \sum_{k \le \lambda+1} \frac{e^{-(\lambda+1)} (\lambda+1)^k}{k!}$$

$$= e^{-(\lambda+1)} \left[ \sum_{k \leq \lambda} \frac{e^{\lambda k} - (\lambda+1)^k}{k!} - \frac{(\lambda+1)^{\lambda}}{\lambda!} \right]$$
  

$$\geq \frac{e^{-(\lambda+1)}}{\lambda!} \left[ \sum_{k \leq \lambda} \left( e^{\lambda k} - (\lambda+1)^k \right) - (\lambda+1)^{\lambda} \right]$$
  

$$\geq \frac{e^{-(\lambda+1)}}{\lambda(\lambda-1)\lambda!} \left[ e^{\lambda+2} - \lambda(\lambda+1)^{\lambda} - (e-1)\lambda - 1 \right].$$

The expression in brackets is at least

$$\lambda^{\lambda} \left( e\lambda^{2} - \lambda(1 + \frac{1}{\lambda})^{\lambda} \right) - (e - 1)\lambda - 1 \geq e\lambda^{\lambda+1}(\lambda - 1) - (e - 1)\lambda - 1,$$

which is positive for all  $\lambda \geq 2$ .

Finally, we apply the Central Limit Theorem [53] to observe that  $\lim_{\lambda\to\infty} p_{\lambda} = \frac{1}{2}$ .

Let  $\{X_i\}$  be a sequence of *n* iid random variables. Suppose the expectation  $\mu = \mathbb{E}[X_i]$  and variance  $\sigma^2 = \mathbb{V}[X_i]$  exist and let  $S_n = \sum_i X_i$ . Then for every fixed  $\beta$ ,

$$\lim_{n \to \infty} \Pr\left[\frac{S_n - n\mu}{\sigma\sqrt{n}} < \beta\right] = \Phi(\beta),$$

where  $\Phi(x) = \Pr[X \le x]$ , and X is a normally distributed random variable with E[X] = 0 and V[X] = 1.

As an immediate corollary, let there be  $n = \lambda$  Poisson distributed variables each with  $\mu = \sigma = 1$ . Then  $S_n$  is distributed Poisson with mean and variance  $\lambda$ , i.e.,  $S_n \sim Y$ . Now letting  $\beta = 0$  gives the result.

**Theorem 5.3.3** If  $f(x_1, \ldots, x_n)$  be a non-negative function, then

$$E\left[f(X_1^{(m)}, \dots, X_n^{(m)})\right] \leq \sqrt{2\pi m} E\left[f(Y_1^{(m)}, \dots, Y_n^{(m)})\right].$$
 (5.4)

Further if  $E\left[f(X_1^{(k)}, \ldots, X_n^{(k)})\right]$  is monotonically increasing with k, then

$$\mathbb{E}[f(X_1,\ldots,X_n)] \leq 4 \cdot \mathbb{E}[f(Y_1,\ldots,Y_n)].$$
(5.5)

Similarly if  $E\left[f(X_1^{(k)}, \ldots, X_n^{(k)})\right]$  is monotonically decreasing with k, then

$$\mathbf{E}[f(X_1,\ldots,X_n)] \leq 2 \cdot \mathbf{E}[f(Y_1,\ldots,Y_n)].$$
(5.6)

**Proof.** We have that

$$E \Big[ f(Y_1^{(m)}, \dots, Y_n^{(m)}) \Big] = \sum_{k \ge 0} E \Big[ f(Y_1^{(m)}, \dots, Y_n^{(m)}) \Big| \sum_i Y_i^{(m)} = k \Big] \cdot \Pr \Big[ \sum_i Y_i^{(m)} = k \Big]$$
  
$$= \sum_{k \ge 0} E \Big[ f(X_1^{(k)}, \dots, X_n^{(k)}) \Big] \cdot \Pr \Big[ \sum_i Y_i = k \Big], \quad \text{using (5.3)}$$
  
$$\ge E \Big[ f(X_1^{(m)}, \dots, X_n^{(m)}) \Big] \cdot \Pr \Big[ \sum_i Y_i = m \Big].$$

Equation (5.4) now follows by noting that  $\Pr[\Sigma_i Y_i = m] = \frac{m^m e^{-m}}{m!}$  and applying Stirling's approximation.

If 
$$\operatorname{E}\left[f(X_{1}^{(k)},\ldots,X_{n}^{(k)})\right]$$
 increases with  $k$ , then by a similar argument we have  

$$\operatorname{E}\left[f(Y_{1}^{(m)},\ldots,Y_{n}^{(m)})\right] = \sum_{k\geq 0} \operatorname{E}\left[f(X_{1}^{(k)},\ldots,X_{n}^{(k)})\right] \cdot \operatorname{Pr}\left[\sum_{i}Y_{i}=k\right], \quad \text{using (5.3)}$$

$$\geq \operatorname{E}\left[f(X_{1}^{(m)},\ldots,X_{n}^{(m)})\right] \cdot \operatorname{Pr}\left[\sum_{i}Y_{i}\geq m\right]$$

Since  $E[\Sigma_i Y_i] = m$ , Lemma 5.3.2 says that  $Pr[\Sigma_i Y_i \ge m] \ge \frac{1}{4}$ , and (5.5) follows. The derivation of (5.6) is similar.

In what follows, we will use "[c]" to stand for the indicator  $I_c$  which has value one if condition c is true, zero otherwise.

**Example 5.3.4** Suppose we want to show that the number of bins with at least  $\theta$  balls is not likely to be much larger than B, for appropriate  $\theta$  and B. Define f as follows:

$$f(x_1,\ldots,x_n) = \left[\sum_i [x_i \ge \theta] > B\right].$$

Since  $E\left[f(X_1^{(k)}, \ldots, X_n^{(k)})\right]$  increases with k, we can upper bound the probability that more than B bins have at least  $\theta$  balls each, by the probability that  $I = \sum_i I_i > B$ , where  $I_i = \left[Y_i^{(m)} \ge \theta\right]$ , where the  $Y_i$ 's are iid Poisson with mean  $\frac{m}{n}$ . For  $m \le n, \theta > 2$  will suffice to infer that  $\Pr[Y_i \ge \theta] \le 2\Pr[Y_i = \theta] \le \frac{2}{\theta!}$ . Note also that  $\Pr[Y_i \ge \theta] \ge \Pr[Y_i = \theta]$ .  $\Box$ 

#### 5.3.3.2 Upper bound for Threshold

**Theorem 5.3.5** For m = n balls, if r is fixed independent of n, then  $\mathsf{Threshold}(T)$  terminates after r rounds with high probability, where  $T = O\left(\sqrt[r]{\frac{r \log n}{\log \log n}}\right)$ .

**Proof.** Let  $k_j$  be the number of balls to be (re)thrown after j rounds ( $k_0 = n$ ). We will show by induction that

$$k_j \leq n \left(\frac{4\log n}{T!}\right)^{\frac{T^j-1}{T-1}} \tag{5.7}$$

with high probability (by which we mean  $1 - O(\frac{1}{n^a})$  for a suitably large constant a). The case j = 0 is readily verified. Now consider the situation when  $k_j$  balls are thrown into n bins in the (j + 1)-st round. For large enough  $n, k_j/n \leq 1 < T$  for all  $j \in \{0, \ldots, r\}$ .

Moving freely between the occupancy and the Poisson case by virtue of Theorem 5.3.3, we can now proceed as in Example 5.3.4 to obtain that

$$\frac{e^{-k_j/n}(k_j/n)^T}{T!} \leq \Pr[I_i = 1] \leq \frac{2e^{-k_j/n}(k_j/n)^T}{T!}.$$
(5.8)

the expected number  $I_i$ 's that are ones is at most  $\frac{2ne^{-k_j/n}(k_j/n)^T}{T!}$ . Now we use the simple version of Chernoff bound, which says that for n Bernoulli trials with success probability p, and  $0 < \epsilon \leq 1$ , the number of successes  $S_n$  obeys  $\Pr[S_n \geq (1+\epsilon)np] \leq e^{-\epsilon^2 np/3}$ . In our case  $p = \Pr[I_i = 1]$ . By setting  $\epsilon = 1$  and verifying that  $np \geq a \log n$  for a suitably large constant a (this suffices since r is constant too), we get that with high probability, after the (j+1)-st round, w.h.p. the number of bins with more than T balls is at most  $\frac{4ne^{-k_j/n}(k_j/n)^T}{T!}$ . We can make the conservative upper bound assumption that with probability exponentially close to one, none of these over-full bins have more than  $\log n$  balls in it, so that w.h.p.,

$$k_{j+1} \leq \frac{4ne^{-k_j/n}\log n}{T!} \left(\frac{4\log n}{T!}\right)^{\frac{T(T^{j}-1)}{T-1}} \leq n \left(\frac{4\log n}{T!}\right)^{1+\frac{T(T^{j}-1)}{T-1}} = n \left(\frac{4\log n}{T!}\right)^{\frac{T(T^{j+1}-1)}{T-1}}.$$
(5.9)

The induction follows. It only remains to verify that for constants r and  $0 < \epsilon < 1$ , and suitably large n,  $T = O\left(\sqrt[r]{\frac{r \log n}{\log \log n}}\right)$  suffices to reduce  $k_{r-1}$  to less than  $n^{1-\epsilon}$ , at which point one more round suffices. This immediately implies a maximum load of O(rT), which, for fixed r, is  $O(\sqrt[r]{\log n}/\log \log n)$ .

Some additional bounds can be derived for non-constant r. In particular, for T = 1, one can show that  $r = \Theta(\log \log n)$  w.h.p.

#### 5.3.3.3 Lower bound

In this section we will show a matching lower bound for fixed r. We can show directly that the **Threshold** protocol will need  $\Omega(r)$  rounds to complete given a choice of Tonly slightly smaller than the above choice in the upper bound case. More generally, we can show this property for all symmetric, non-adaptive protocols.

Symmetric, non-adaptive protocols can be modeled using an *edge orientation* setting. Consider the case of n balls and n bins. Each ball is an vertex in an n-vertex graph. Initially, n random undirected edges are added to the graph, where by "random" we

mean that for each edge, independent of others, the two endpoints are chosen uniformly at random (the issue of self-loops does not affect the asymptotic analysis). Choosing a final destination is equivalent to choosing an orientation for the edge. The goal of the algorithm is to minimize the maximum indegree over all vertices of the graph. In the case where there are n balls and n bins, the corresponding graph is a random graph from  $\mathcal{G}_{n,n}$ , the set of all graphs with n vertices and n edges.

**Definition 5.3.6** Let N(x) be the edges incident with vertex x. The (r, x)-neighborhood of an edge e = (x, y), denoted by  $N_{r,x}(e)$ , is defined inductively by:  $N_{1,x}(e) = N(x) - \{e\}$ ,  $N_{r,x}(e) = N(N_{r-1,x}(e)) - \{e\}$ .

A ball (edge) (x, y) gets *confused* about which way to orient if large neighborhoods of both the endpoints are isomorphic. Specifically, suppose we invest r rounds of communication. In each round of communication, a ball discovers a little more about the graph. Specifically, since we want lower bounds, we may assume that the bins transport all available information about the balls whenever possible. In r rounds, a ball e = (x, y)knows everything about the balls in  $N_{r,x}(e) \cup N_{r,y}(e)$ , and no more, before it must commit to a bin; this follows from a simple induction argument. Suppose  $N_{r,x}(e)$  and  $N_{r,y}(e)$  are isomorphic trees rooted at x and y. Then the ball has no reason to prefer one bin over another, and must essentially choose randomly between bins x and y.

Obvious tie-breaking schemes do not affect the lower bound. For instance, if the balls are ordered at the bins, either by random I.D. numbers or by a random permutation, and then choose a bin according to their rank, the balls are essentially choosing a bin at random. The proof can easily be modified also for the case where the balls are ranked at the bins by some fixed ordering, by using the symmetry of the destination choices of the balls.

If many confused balls are incident on one bin, with constant probability, over half of them will opt for the bin, which will become overloaded. We thus need to show that for r and T suitably related to n, a random graph  $G \in \mathcal{G}_{n,n}$  has, with high probability, a *near isolated* (T, r)-tree, defined as follows (see Figure 5.4).

1. It is a rooted balanced tree of depth r. All the leaves are at depth r.

2. The root has out-degree T. Internal nodes have one parent and T-1 children.

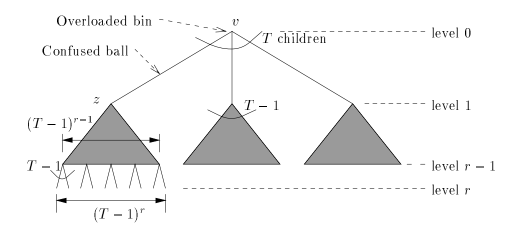


Figure 5.4: A (T, r)-tree with confused balls.

3. The tree is near-isolated in the sense that no edge of G other than the tree edges are incident to any internal node of the tree.

**Theorem 5.3.7** With constant probability, a random graph from  $\mathcal{G}_{n,n}$  contains a nearisolated (T,r)-tree for fixed r and  $T = \Omega(\sqrt[r]{\log \log n})$ . Assuming confused balls pick each bin with probability  $\frac{1}{2}$ , some bin will have load  $\Omega(T)$  with constant probability.

**Proof.** A (T, r)-tree has

$$\begin{aligned} X_{(T,r)} &= 1 + T \sum_{0 \le k < r} (T-1)^k = 1 + T \frac{(T-1)^r - 1}{T-2} \approx T^r & \text{vertices}, \\ Y_{(T,r)} &= 1 + T \sum_{0 \le k < r-1} (T-1)^k = 1 + T \frac{(T-1)^{r-1} - 1}{T-2} \approx T^{r-1} \end{aligned}$$

vertices incident only on internal edges, and  $Z_{(T,r)} = X_{(T,r)} - 1$  internal edges. We will drop the subscript (T, r) from X, Y, and Z when clear from context.

Let  $\vec{v} = (v_1, \ldots, v_X)$  be a sequence of  $X_{(T,r)}$  vertices, and let  $I_{\vec{v}}$  be the indicator for " $\vec{v}$  is a near-isolated (T,r)-tree," with  $v_1$  being the root,  $v_2, \ldots, v_{T+1}$  being the first level children, and so on. Let  $I = \sum_{\vec{v}} I_{\vec{v}}$ . Note that I is not an indicator; it is a non-negative integer random variable. For such variables it is routine to verify that  $\Pr[I=0] \le 1 - \mathbb{E}[I]^2 / \mathbb{E}[I^2]$ . We will use this inequality to show that  $\Pr[I=0]$  is strictly less than one; for this we will bound E[I] from below and  $E[I^2]$  from above by appropriate constants in the next two lemmas.

**Lemma 5.3.8** E[I] is at least a constant for fixed r and  $T = \Omega(\sqrt[r]{\log \log n})$ .

**Proof.** Given  $\vec{v}$ , the probability that  $\vec{v}$  represents a (T, r)-tree is

$$\frac{\binom{n-Y}{2}^{n-Z}\binom{n}{Z}Z!}{\binom{n}{2}^{n}}.$$
(5.10)

The denominator is all possible ways of picking n edges in an n-vertex graph. For the numerator, we multiply the number of ways in which Z tree edges can be chosen and permuted, and the number of ways in which the remaining n - Z edges can be assigned endpoints from the n - Y external or excluded nodes. The number of ways of picking  $\vec{v}$  is

$$\binom{n}{1;T;\underbrace{T-1;\ldots;T-1}_{\frac{Z-T}{T-1}}} = \frac{n!}{T((T-1)!)^{(Z-1)/(T-1)}(n-X)!},$$
(5.11)

where the multinomial  $\binom{n}{a;b}$  means  $\frac{n!}{a!b!(n-a-b)!}$ . Multiplying (5.10) and (5.11) and simplifying we get

$$\mathbf{E}[I] = \frac{n2^{Z} \left(1 + o(1)\right)}{e^{2Y} T \left((T-1)!\right)^{(Z-1)/(T-1)}},$$

from which the result follows for the given choice of T.

**Lemma 5.3.9** For fixed r and  $T = \Omega(\sqrt[r]{\log n / \log \log n})$ ,  $E[I^2] = (E[I] + E[I]^2)(1 + o(1))$ .

**Proof.** Since  $E[I^2] = E[I] + \sum_{\vec{v} \neq \vec{w}} I_{\vec{v}} I_{\vec{w}}$ , we need to find only the second term in the sum. There are two cases: the trees represented by  $\vec{v}$  and  $\vec{w}$  do not intersect, or they do. In the former case, it is routine to check that

$$\sum_{\vec{v}\cap\vec{w}=\emptyset} \mathbb{E}[I_{\vec{v}}I_{\vec{w}}] \leq \frac{\binom{n-2Y}{2}^{n-2Z}\binom{n}{2Z}(2Z)!}{\binom{n}{2}^{n}} \times \frac{n! (n-Y)!}{(n-X)! (n-X-Y)! T^2((T-1)!)^{\frac{2(Z-1)}{T-1}}}.$$

In the latter case, the only way  $\vec{v}$  and  $\vec{w}$  can intersect at an internal node is when the root of  $\vec{w}$  is an internal node in  $\vec{v}$  (without loss of generality). The greater the overlap between  $\vec{v}$  and  $\vec{w}$ , the more likely it is that these will be subgraphs of a random graph G. Let z be a fixed child of the root of  $\vec{v}$ . By the above argument, we can bound

$$\sum_{\vec{v} \cap \vec{w} \neq \emptyset, \vec{v}, \vec{w}} \mathrm{E}[I_{\vec{v}}I_{\vec{w}}] \leq Y \cdot \mathrm{E}[I_{\vec{v}}] \cdot \Pr[z \text{ is a root of a } (T, r) \text{-tree, given } I_{\vec{v}} = 1].$$

Let  $p_z$  be the last  $\Pr[\cdots]$ . The number of new edges that must be added to  $\vec{v}$  so that z can be the root of a (T,r)-tree is  $Z' = (T-1)^r$ . If we can show  $p_z = o(1/Y)$ , we will be done. The denominator of  $p_z$  is  $\binom{n-Y}{2}^{n-Z}$ , the number of ways in which n-Z additional edges can be added to the graph without touching any of the Y internal nodes of  $\vec{v}$ . The numerator has several factors:

- The number of ways in which additional vertices can chosen to complete the (T, r)-tree rooted at z is  $(\underbrace{T-1;...;T-1}_{Z'/(T-1)}) \frac{1}{(T-1)!} \binom{T-1}{1;...;1} \frac{Z'/(T-1)}{Z'/(T-1)}$ .
- The number of ways of picking Z' edges from n Z edges to complete the (T, r)-tree rooted at z, which is at most  $\binom{n-Z}{Z'}Z'!$ .
- The number of ways of adding n Z Z' remaining edges while avoiding our trees of interest; this is  $\binom{n-Y-\frac{Z'}{T-1}}{2}^{n-Z-Z'}$ .

Using Stirling's approximation,  $p_z$  can be simplified to roughly  $\frac{(2m/n)^{Z'} \exp(-\frac{2mZ'}{nT})}{T!^{Z'/T}}$ . Summing over all cases completes the proof. We omit the manipulations.

The techniques carry over directly to the case where there are m balls and n bins, and each ball sends requests to d > 2 bins, by generalizing each edge to a d-vertex hyperedge. The details can be found in [2].

#### 5.3.4 Other protocols

Other parallelizations of the Azar *et al*  $\Omega(n)$ -round protocol are possible. For example, we can run the following protocol, called Sibling(2):

- 1. In parallel each ball picks two random bins and sends them requests.
- 2. In parallel each bin sends the total number of requests that it has received to each requesting ball.
- 3. Each ball commits to the bin that reported fewer requests.

Notice that this, too, is a symmetric, non-adaptive protocol and therefore the lower bound of §5.3.3.3 applies with r = 2. Intuitively, the "noise" in the request information, as against previous commits as in Azar *et al*'s protocol, makes it impossible to achieve an  $O(\log \log n)$ 

ſ	Balls	Simple	Azar	Sibling(2)	Threshold(T)		
	n	Random	$et \ al$	d=2	r = 2	r = 3	r = 5
Γ	$10^{6}$	8-11	4	5-6	5-6	4-5	4
	$5 \cdot 10^6$	9 - 12	4	5-6	5-6	4-5	4
	$10^{7}$	9 - 12	4	5-6	5-6	4-5	4
	$5 \cdot 10^7$	9-12	4	5-6	6	5	4

Table 5.1: Simulation results. d is the number of bins that a ball sends initial request to; this is set to two. r is the number of rounds.

load balance. We can show a corresponding upper bound [1]; we omit the details. Notice that this is a *synchronous* protocol; bins need to know when all balls are done sending requests before sending out load information.

**Theorem 5.3.10** The Sibling(2) protocol achieves an  $O\left(\sqrt{\frac{\log n}{\log \log n}}\right)$  load balance.

#### 5.3.5 Concluding remarks

For simplicity, we limited the number of bins chosen in each round by each ball to one or two. The results above can be extended to *d*-way choices by the balls, where *d* is constant or grows sufficiently slowly with *n*. The details can be found in Adler *et al*'s paper [1]. Also, the general case of  $m \ge n$  balls and *n* bins can be handled readily.

Since the maximum load grows very slowly with n for all these schemes, and our analysis is asymptotic and probabilistic, we were interested in performance seen in actual simulations. We set the number of balls and bins to be equal, over the range of values shown in Table 5.1. The numbers given in the table represent the ranges for the maximum load found after between fifty and one hundred trials for each strategy. As expected, both Sibling and Threshold perform somewhere between simple random allocation and Azar *et al.* Also, we note that T was not optimized for Threshold; in practice one might want to take care to optimize the threshold for a given number of balls.

## Chapter 6

## Survey of related work

Scheduling and resource allocation problems have been studied extensively in all the forms discussed in this thesis. In this chapter we review some of the major results in the various areas. Most of the exact problems are  $\mathcal{NP}$ -hard, sometimes in the strong sense, and therefore the thrust is on developing good approximation algorithms. Particular technical references and their relation with this thesis are given in specific chapters.

## 6.1 Communication scheduling

The results in Chapter 2 follow (and generalize) a large number of preliminary results in communication optimization, such as message vectorizing, coalescing, and redundancy elimination [72, 123, 20, 5, 21, 68]. Message vectorization is the technique of hoisting communication of array elements or sections out of loops to produce a single large message. Typically this works locally on single loop-nests [123, 72, 67, 83, 68]. As compilers were applied to more complex code, eliminating redundant communication beyond single loop-nests and even across procedures became essential. This was typically done by tracing the uses of an array (section) to a defining statement, placing all the communication entries after that def, and canceling subsumed communication entries [69].

Communication can also be regarded as invoking a long-lasting asynchronous operation on the network "functional unit." To minimize the latency penalty for fine-grain communication, it helps to detect writes to shared memory that can be reordered or overlapped without violating the programmer's view of consistency of program variables. An early result in this direction finds cycles of conflicting reads and writes and inserts minimal synchronization to break all cycles [102]. This technique has recently been improved for single program multiple data (SPMD) sources [82]. Another related area is instruction scheduling, which has been intensively studied [94]. The performance impact of these scheduling algorithms has grown with the advent of superscalar RISC architectures which have multiple functional units permitting out-of-order execution.

## 6.2 Multiprocessor scheduling

Job scheduling in multiprocessors has been extensively researched in both Operations Research and Computer Science [71]. Two decades later there is significant interest in the Computer Science community on scheduling *parallel* jobs [111, 41]. For independent jobs that need a fixed given number of processors, various rectangle packing algorithms naturally model the problem: every job can be regarded as a rectangle of width equal to the number of processors, and of height representing the running time [14, 13].

It is more realistic to assume that processors are a malleable resource: i.e., they can be traded for time. This trade-off can be specified as an arbitrary function for the running time on any given number of processors. For jobs with this model of trade-off, various constant-factor approximations for makespan are known for the special case where there is no precedence relation; i.e., the jobs are independent [16, 87]. Our results in Chapters 3 and 4 extend these results.

In a slightly simpler but still reasonably realistic model of malleability, every job specifies a maximum number of processor up to which it will give perfect speedup. For this model, constant factor makespan approximations are known that handle precedence between jobs exposed on-line. Furthermore, the algorithm is *non-clairvoyant*; i.e., it does not need to know the running time of a job before completion [51]. These and similar algorithms [15, 34] are essentially based on online gambling techniques.

## 6.3 Minsum metrics

Most makespan results are relatively straight-forward in that the approximation algorithms are compared against a very simple lower bound: the sum of average work per processor and critical path length [64, 51]. Also, makespan results are relevant for a batch of jobs in isolation; in fact, on-line arrival of the batch make little difference [105] (also see Chapter 5). In contrast, for a parallel computer installation, other measures that emphasize notions of fairness or resource utilization are considered more appropriate. In such scenarios, on-line arrivals may make problems harder.

Some such measures of recent interest are weighted average completion time (WACT) and weighted flow time (WFT), defined as follows. Suppose job j arrived at  $a_j$  and is completed at time  $C_j$ . It also specified a weight, or priority,  $w_j$ . Then the WACT is  $\sum_j w_j C_j$  and WFT is  $\sum_j w_j (C_j - a_j)$ . Another dimension of the problem is whether the jobs arrive on-line or are all known off-line; in the former model the scheduler does not know of job j before time  $a_j$ .

For independent non-malleable jobs in the off-line setting, several WACT results have been known [111]. Recently, Hall etal suggested two important general frameworks for off-line and on-line WACT optimization [70]. Our results in Chapter 4 build upon these results.

### 6.4 Resource scheduling

Garey *et al* [57] study a generalization of non-malleable jobs where there are *s* distinct types of resources. At any time a total of at most one unit of each resource can be allocated. Job *j* specifies an *s*-dimensional resource vector  $\vec{r_j} = (r_{jk})$ , where  $r_{jk} \leq 1$  is the fraction of resource of type *k* that *j* keeps allocated for the duration it runs. Their positive (makespan) results are for independent jobs [57] and for dependent jobs all of equal duration [58].

We showed in Chapter 4 that for the general case with arbitrary times and precedence, even with only one resource type, the simple lower bound: the maximum of average resource-time product  $V = \sum_j t_j r_j$  and critical path II, cannot always be matched to a constant factor by even an optimal makespan algorithm. (Here  $\Pi_j$  is the earliest time job j can finish ignoring resource needs and only considering the total time of its longest predecessor chain.) Not surprisingly, Garey *et al* [57] observe that list scheduling algorithms can perform very poorly (give a makespan that is larger than optimal by a factor equal to the number of jobs) on such problems.

## 6.5 Distributed load balancing

The above settings all assumed a centralized scheduler. For fine-grain jobs, such as in parallel programs with light-weight task parallelism, an important concern it to make scheduling decisions in a decentralized way, so that there is no central bottleneck where processor allocation is done.

The bottleneck can be relieved in a variety of ways, each of which reduces communication cost by sacrificing global load information and thus risking some load imbalance. We studied *work sharing*, where busy processors forward jobs to random processors. This is similar to Karp and Zhang's model [78]. Work sharing is good at dispersing hot-spots.

Another technique that has been designed and analyzed lately is *work stealing*, where idle processors ask for work [19]. The theoretical bounds are only applicable to divide and conquer, where a finite, fixed task tree has to be completely executed. The basic idea, dating back to Kung and Wu's work, is for each processor to follow depth-first tree expansion order locally, while an idle processor that "steals" work from others gets a shallowest unexpanded node [122]. Work stealing is good at reducing communication, since many jobs run on the processors where they were generated. They do not handle more general cases like branch and bound where the total work to find the solution may depend on the schedule.

Both of the above assume an amorphous network. For general network topologies, another technique that has been studied a great deal recently is *diffusion*, where neighbors exchange local load information and then move some jobs from busy to lazy processors [98, 61].

Our application of random allocation to load balancing scenarios is related to results on random hashing and PRAM emulation [22, 77]. In those problems, reusing hash functions and economizing on randomness are important issues; we have used randomness generously, our main concern being communication and load balance.

### 6.6 Open ends

Except in Chapter 5, we have only considered *clairvoyant* scheduling, where the resources and time required by a job are specified when the job arrives. This is reasonable

for the applications we have targeted. In the design of schedulers for general-purpose workstations that run a mix of diverse jobs for which no *a priori* estimates of time and resource can be made, *non-clairvoyance* is of great importance. As might be suspected, non-clairvoyant schedulers are remarkably handicapped compared to clairvoyant ones [92], even when permitted the (essential) power of preemption. These results are for sequential jobs; some of them have been extended to the perfectly malleable job model [41].

A further refinement of the scheduler performance metric is weighted flow time, defined as  $\sum_{j} w_{j}(C_{j} - a_{j})$ , where  $w_{j}$  is the priority of job j, and  $a_{j}$  and  $C_{j}$  are its arrival and completion time respectively. The results reported above show that polynomial time constant factor approximations for weighted average completion time are possible for a variety of situations, including non-clairvoyant ones (the latter with preemption). In sharp contrast, a remarkable recent result shows that for non-preemptive weighted flow time, approximating better than about a factor of  $\sqrt{n}$  is  $\mathcal{NP}$ -hard [80] even in the simplest model of sequential jobs on a single processor, with an off-line clairvoyant scheduler. This lower bound holds a fortiori for our generalized job models. It is perhaps unreasonable in practice to demand near-optimal weighted flow time without preemption. Here an interesting distinction arises between one and many processors. For a uniprocessor, the least remaining processing time or LRPT heuristic is easily seen to be optimal for total flow time  $\sum_{j} (C_j - a_j)$ , while for any fixed number of processors  $m \ge 2$ , Du *et al* [46] show that obtaining a preemptive schedule to minimize total flow time is  $\mathcal{NP}$ -hard. In fact, C. Phillips (personal communication) has shown that LRPT is off by a factor of  $\Omega(\log n)$  from optimal for just two machines.

## Chapter 7

# Conclusion

In this thesis we have formulated and solved several resource allocation and scheduling problems relevant to parallel computation, in the context of compilers, runtime systems, operating systems, and applications. The specific contributions of the thesis are summarized below.

- By judicious compile time communication code scheduling through global array dataflow analysis, we optimized network access while also performing redundant communication elimination; this reduced communication cost by factors of two to three in data-parallel programs.
- By extending the pure data-parallel model to support dynamic task parallelism, and designing runtime scheduling heuristics, we extended the class of scientific applications that can be efficiently solved. Two to three times the MFLOPS of pure data-parallelism was obtained.
- We initiated a study of generalized resource scheduling that is intended to model realistic extensions of processor scheduling scenarios. We designed new algorithms that handle inflexible resources such as memory, precedence between jobs, and job priorities.
- For irregular, dynamic problems, we gave a precise characterization of the trade-off between load balance and communication cost. These results give valuable guidelines for distributed load-balancing systems. We also showed that randomness is useful to exploit in fine-grain task allocation.

It seems that promising areas of parallel computing research will be guided by the following goals.

- Reasonable and cost-effective speedups must be obtained for essentially all applications on low-end personal desktop installations.
- Remarkable speedups and scaleups must be obtained on perhaps expensive high-end installations shared by many users.

Multithreaded architectures and aggressive compilation are prominent means to attain the first objective. Moreover, as transistor density increases, it becomes more attractive to design multiple processors on a chip. Such machines can either be used to improve utilization over different unrelated jobs, or parallelize a single job. For the former goal, compilers have to solve the problem of scheduling threads which are, roughly speaking, a sequence of operations on different functional units. This appears related to flow-shop problems in Operations Research. For the latter goal, two different problems arise. The first problem is to detect parallelism; since our goal here is general-purpose parallelism, this detection must be automatic. For popular languages like C or C++, detecting parallelism in presence of aliases, pointers, and linked data structures is very difficult. Several projects such as Suif, Olden, and Jade, as well as commutativity and synchronization analyses have made initial progress in these directions [24, 118, 97]. The second problem is for the compiler to explicitly orchestrate communication between different register files or functional units. The latter goal appears to guide multiprocessor system design in the following direction.

- They must be built out of commodity processors with a stable instruction set and interface between the processor and the network, so that a processor upgrade can be directly integrated into an existing multiprocessor.
- The network interface and software must have performance high enough for asynchronous, small messages, to enable efficient execution of challenging irregular and dynamic codes.
- Since such installations are likely to be expensive, system support at the OS level has to be provided so that apart from running one application well, the system allocates resources to multiple jobs effectively. The work in the latter half of the thesis represents some advance in that direction.

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