PRACTICAL PARALLEL EXTERNAL MEMORY ALGORITHMS
VIA SIMULATION OF PARALLEL ALGORITHMS

by

David E. Robillard

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Abstract

This thesis introduces PEMS2, an improvement to PEMS (Parallel External Memory System). PEMS executes Bulk-Synchronous Parallel (BSP) algorithms in an External Memory (EM) context, enabling computation with very large data sets which exceed the size of main memory. Many parallel algorithms have been designed and implemented for Bulk-Synchronous Parallel models of computation. Such algorithms generally assume that the entire data set is stored in main memory at once. PEMS overcomes this limitation without requiring any modification to the algorithm by using disk space as memory for additional “virtual processors”. Previous work has shown this to be a promising approach which scales well as computational resources (i.e. processors and disks) are added. However, the technique incurs significant overhead when compared with purpose-built EM algorithms. PEMS2 introduces refinements to the simulation process intended to reduce this overhead as well as the amount of disk space required to run the simulation. New functionality is also introduced, including asynchronous I/O and support for multi-core processors. Experimental results show that these changes significantly improve the runtime of the simulation. PEMS2 narrows the performance gap between simulated BSP algorithms and their hand-crafted EM counterparts, providing a practical system for using BSP algorithms with data sets which exceed the size of RAM.
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PEMS2 could not exist without Mohammad Nikseresht, whom I thank for the PEMS1 implementation and setting up a development site for the project.

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Chapter 1

Introduction

1.1 Background and Motivation

External Memory (EM) algorithms are designed to work with data sets much larger than main memory. Though strictly defined in more general terms, EM models typically consider a 2-tier memory hierarchy: “main memory” (RAM) and “external memory” (disk). Algorithms designed for these models explicitly transfer blocks of data between these levels of memory, attempting to minimize the number of transfers between them. In addition to minimizing data transfer, EM algorithms may also be designed to access external memory (e.g. disk) in an efficient pattern to minimize expensive disk seeking.

Unfortunately, most algorithms are designed for Random Access Memory (RAM) models rather than EM. RAM algorithms work with a single level of memory, and assume a read or write at any location has a fixed constant cost. Because RAM algorithms do not consider locality of reference a performance factor, translating a RAM algorithm into an EM algorithm with acceptable performance is not simple or automatable in the general case. There are, however, certain classes of algorithms which can work well in an EM context despite not being designed with EM specifically in mind.

The goal of this thesis is to enable the practical use of such algorithms on problems that exceed the size of RAM, allowing an algorithm to scale beyond the limits of main memory without requiring a complete rewrite. Parallel algorithms are particularly desirable in this context since very large problems may exceed the resources of a single machine and sequential computation with data of this magnitude in reasonable time is generally not feasible.
1.2 Computational Models

1.2.1 Parallel Disk Model (PDM)

The multiple disk model originally proposed by Vitter and Shriver [19][20], usually referred to as the PDM model, is commonly used for designing disk-based algorithms. In PDM, an algorithm has access to an internal random access memory of size $M$, and $D$ disks which transfer in blocks of size $B$. I/O is fully parallel and blocked, i.e. a transfer of size $BD$ (to $D$ disks) is considered a single I/O operation. The complexity of an algorithm is measured exclusively in terms of the number of such I/O operations, ignoring other factors such as computation time. This reflects the reality that disk access is orders of magnitude more expensive than RAM access. Computation time of an algorithm may also be given for algorithms with especially high computational requirements, though typically I/O time dwarfs computation time by a large enough margin that computation does not significantly contribute to the total run time.

1.2.2 Bulk Synchronous Parallel (BSP) and Related Models

The BSP model [18] was proposed as an abstract “bridging model for parallel computation”. BSP serves as a common model for both system/hardware and algorithm/software designers which allows for accurate performance analysis on a wide range of parallel computers. BSP considers a set of processors each with independent local memory that communicate by sending messages between each other. Computation proceeds in a series of synchronised “supersteps”, each of which consists of a “computation superstep” followed by a “communication superstep”. The total runtime of an algorithm is thus the sum of the computation time, communication time, and synchronisation time.

A superstep where each processor sends and receives $O(h)$ data is called an “$h$-relation”. BSP* [3] and Coarse Grained Multicomputer (CGM) [8], other common models of parallel computation, are special cases of BSP with restrictions on $h$ to ensure a more coarse grained computation. In practical terms BSP* and CGM algorithms proceed in an identical fashion to BSP algorithms, i.e. in a series of supersteps. Accordingly, the three are considered equivalent for much of this thesis and collectively referred to as “BSP-like”. 
Though all BSP-like models function similarly, the performance characteristics of restricted models have important implications when used with EM. CGM requires that each processor works with $O(\frac{N}{v})$ local data (i.e. $h = O(\frac{N}{v})$). This ensures balanced computation and communication with coarse granularity. Synchronisation overhead is thus minimized, while processor and disk parallelism is exploited efficiently. Since communication and synchronisation in PEMS is relatively expensive due to disk I/O, these characteristics are especially desirable in this context. The applications presented in Chapter 8 are CGM algorithms.

BSP-like algorithms are useful on a wide variety of configurations, particularly the common and inexpensive “cluster” style of parallel computer composed of several commodity machines connected by a switched Ethernet network.

1.2.3 EM-BSP Models

The parallel and distributed memory nature of BSP-like algorithms is advantageous from an EM perspective since, as in PDM, a collection of parallel disks can perform I/O much faster than a single disk. The EM-BSP, EM-BSP*, and EM-CGM models \cite{1,3,6,7} augment the corresponding BSP-like model by adding local disk(s) to each machine. Such a configuration is shown in Fig. 1.2.

Computation proceeds in supersteps as in BSP, except each processor may access local disk as necessary during the computation superstep. Thus, the EM-BSP models
can be considered a hybrid of the BSP-like models and the PDM model: synchronisation and communication is inherited from BSP, and I/O from PDM.

1.3 Previous Work

1.3.1 STXXL

STXXL is a C++ library for EM algorithms. STXXL is composed of many layers, as shown in Fig. 1.3 (reproduced from [10]). Higher level layers in STXXL make use of the lower level layers, though user applications may directly use any layer, bypassing higher level functionality if desired.

The lower level Block Management and Asynchronous I/O Primitive layers provide generic functionality useful to EM algorithms, such as asynchronous I/O and transparent parallel disk access.

The STL User Layer provides an implementation of the C++ Standard Template Library (STL), the algorithms and data structures component of the C++ standard library. This layer can be used to write C++ code in the standard style that functions as an EM algorithm, or simplify the porting of existing C++ RAM algorithms to EM.

The Streaming Layer provides additional functionality that does not fit within the confines of the STL API. EM-specific techniques such as pipelining and I/O optimal scanning are implemented in this layer.
STXXL provides a rich suite of EM code, making it simple to write advanced EM algorithms at a relatively high level. Notably, all layers above and including the Block Management Layer transparently support parallel disks. Thus, applications built with STXXL can take advantage of parallel disk performance without any specific effort required on behalf of the application developer. The Asynchronous I/O Primitives layer provides a simple, low-level, and portable interface to asynchronous I/O. Since the asynchronous I/O interface of operating systems is typically more complex and varies between systems, this layer is useful to applications that require asynchronous I/O but not the higher level functionality of STXXL.

1.3.2 Cache-Oblivious Algorithms

Cache-Oblivious algorithms [12] are designed with I/O efficiency in mind (unlike RAM algorithms), but without any explicit block size parameters (unlike EM algorithms). For example, traditional EM algorithms explicitly transfer blocks of some size $B$ between disk and main memory. The algorithm implementation must know the value of $B$ at run time. In contrast, a cache-oblivious algorithm is unaware of (or oblivious...
to) any such parameter, and may transfer data with arbitrary size and alignment much like a RAM algorithm. However, unlike most RAM algorithms, cache-oblivious algorithms are analysed in terms of memory transfers of an arbitrary size $B$, and aim to minimize the number of transfers much like an EM algorithm\(^1\). Thus, an efficient cache-oblivious algorithm is efficient for any $B$ and does not require modification to perform well on various systems.

This approach is particularly useful in the presence of cache hierarchies, where many levels of cache are in use at one time, each with a different block size. I/O efficiency is an increasingly important performance factor, even for algorithms that work only with internal memory (RAM). On modern systems, a cache miss can be several hundred times slower than a cache hit [11]. Cache-oblivious literature often presents this problem in the context of a modern processor’s cache and memory hierarchy, though the block size independent nature naturally applies where the lowest level of the memory hierarchy is disk. This suggests cache-oblivious algorithms are a promising strategy for the design of algorithms that show good performance across a very wide range of problems sizes.

1.3.3 MPI

MPI (Message Passing Interface) [11] is an Application Programming Interface (API) for distributed memory parallel programming. MPI provides communication and synchronisation functions useful for many types of parallel program. Most relevant to this thesis are the “collective communication” MPI functions, since these can be used to implement BSP-like algorithms.

Collective communication functions in MPI synchronise all processors, then perform communication. There are many different styles of communication available, such as MPI\_Gather (each processor sends a message to a single processor) or MPI\_Alltoall (each processor sends a message to every other processor)\(^2\).

In a BSP-like program implemented with MPI, a call to a collective communication represents a communication superstep and subsequent superstep barrier. In this way,

\(^{1}\)Cache-oblivious literature typically uses $L$ (for “line”), rather than $B$. This thesis consistently uses $B$ and “block” regardless of whether cache or disk is being discussed.

\(^{2}\)A more detailed description of the collective communication functions described here can be found in Chapter 7.
a BSP-like algorithm can be implemented as a series of MPI collective communication calls interleaved with computation code.

MPI is a widely used interface for distributed memory parallel programming with many implementations for a variety of systems.

### 1.3.4 EM-BSP Simulation

Many parallel algorithms intended to work with large data sets have been designed for BSP-like models. Though these algorithms scale to larger data sets than single processor RAM algorithms by exploiting the memory available to several machines, unfortunately they do not generally make use of disk and are thus limited to problems that fit entirely within main memory.

Fortunately, it is possible to use these existing algorithms with data larger than main memory via simulation in the EM-BSP models\(^3\). The basic idea is to simulate a number of "virtual processors", each with memory small enough to fit into "real processor" main memory. A subset of these virtual processors is executed at once, while the (virtual) memories of others are swapped out to disk. Thus it is possible to run a bulk-synchronous algorithm with total memory size exceeding that of real main memory, limited only by the amount of available disk space.

To illustrate, consider a BSP-like algorithm that requires 128 processors, each with 1 GiB of RAM. If these resources are available, the algorithm may be executed directly. However, this is not the case if only 32 processors are available. Nevertheless, the algorithm may be executed using these limited resources via simulation as follows: for each superstep, rather than run 128 processes in parallel, run 32 processes in parallel, storing any generated messages on disk. Then, another round of 32 processes is executed in a similar fashion, and so on until all 128 processes have been executed. At the end of this process, all computation for the superstep has been completed and all communication is stored on disk, so the next superstep may begin.

In practice, this strategy can be implemented as a library which provides communication functions for use by BSP-like applications. In particular, no special operating system level support is required. All details pertaining to external memory can be managed by this library; the application code need not be changed.

\(^3\)This idea was introduced with the original presentation of the EM-BSP models [13][7][6]
1.3.5 PEMS

PEMS1 [15] (Parallel External Memory System) is an implementation of the EM-BSP simulation technique which provides an API similar to that of MPI. Fig. 1.4 shows an overview of the PEMS1 design.

![PEMS1 Design Diagram](image)

Figure 1.4: PEMS1 Design

Significant modifications to PEMS1 have been made as a part of this thesis. Where the distinction is necessary the previous implementation is referred to as “PEMS1”, and this improved version as “PEMS2”. Both are collectively referred to as “PEMS” where appropriate.

PEMS1 is implemented as a library which transparently handles virtual processor swapping, synchronisation, memory allocation, and communication. The user program is an MPI-like program, but communication may be deferred to disk to allow the simulation of more processors than are actually available.

The interface to PEMS1 is, with a few exceptions, semantically identical to a
subset of MPI, though functions names have a different prefix to avoid conflicts\textsuperscript{4}. When the application calls collective communication functions, PEMS\textsubscript{1} internally performs the necessary network or I/O operations, swapping virtual processors in and out as required. Though much occurs “behind the scenes”, from the application’s point of view the collective communication operation has been completed exactly as if it had been performed directly by MPI.

Internally, the system’s MPI library is used to perform communication between virtual processors on separate real processors. I/O is performed using the operating system’s I/O interface; specifically that of POSIX, the standard common to all UNIX-like systems such as GNU/Linux, Solaris, or Mac OS X.

Thread support in PEMS\textsubscript{1} is handled via the GNU Pth library, which implements user-space threads. This is advantageous for single-core processors since thread switching does not incur the overhead of a kernel-level context switch. However, user-space threads do not allow for true thread concurrency on multi-core machines.

PEMS\textsubscript{1} has been shown to scale well in practice on sorting and list ranking problems significantly larger than the total amount of available RAM \cite{16}. The distinguishing characteristic of PEMS is that existing algorithms not explicitly designed as EM algorithms may be efficiently used with external memory. In addition to the large number of suitable (BSP-like) existing algorithms, a considerable advantage of this approach is the ability to exploit distributed memory parallel computers. While it is possible to implement distributed memory algorithms using STXXL or a cache-oblivious approach, the algorithm must be deliberately designed to have this ability – a significantly more difficult task than designing a sequential EM algorithm. Algorithms designed to support both distributed memory parallelism and external memory are relatively rare. In contrast, all algorithms that work with PEMS are inherently capable of executing on a distributed memory parallel computer.

Because of this ability, PEMS can easily scale to extremely large problem sizes without requiring any modification to the algorithm. If, for example, one wanted to use a straightforward STXXL application on a problem too large to feasibly handle with a single computer, the algorithm may require a significant redesign in order to scale further. A PEMS application, however, can easily scale to very large problem

\textsuperscript{4}This has been resolved in PEMS\textsubscript{2}, see §1.4
sizes by adding disk and/or processor resources as necessary.

Similarly, if a given problem takes an unacceptable amount of time, processor and/or disk resources may be added to improve the run time. Experiments in Chapter 8 and previous work on PEMSl [15] show that, though the STXXL sort is faster than PEMSl given equivalent computational resources, computation resources can be added until PEMSl out-performs the STXXL sort.

1.4 Summary of Contributions

This thesis presents PEMSl2, an enhanced version of PEMSl with new functionality and improved performance and usability. These enhancements include fundamental changes to the simulation process, such as new I/O drivers and multi-core support; as well as new communication primitives with improved performance characteristics.

While PEMSl supported parallelism across several machines in a cluster configuration, SMP (or “multi-core”) on each of these machines was not explicitly supported. Though at the time, most commodity machines were single core, recently multi-core has become ubiquitous. PEMSl2 introduces support for multi-core machines, allowing the simulation to take advantage of multiple cores with less overhead than simply running several local MPI processes. The computation performed by the simulated algorithm is executed in parallel across many cores, allowing for speedup in computation heavy algorithms. However, even for algorithms that are I/O bound, many of the improved communication primitives achieve an I/O reduction proportional to the number of local cores available.

PEMSl used explicit, blocking, aligned I/O operations exclusively. While the improvements presented here can also work in the same fashion, the implementation has been redesigned to allow simple switching between various I/O “drivers”. In conjunction with other changes, this allows for the use of asynchronous I/O, or memory-mapped I/O, both of which have significantly different performance characteristics to traditional blocking I/O. In particular, memory-mapped I/O is interesting because a superstep does not necessarily incur a swap of the entire context as with explicit I/O. With memory mapped I/O the memory access characteristics of the simulated algorithm dictate the I/O performed during simulation, allowing algorithms to take advantage of desirable memory access characteristics when used with PEMSl.
Experiments show that these I/O strategies are beneficial in some cases, but not always an improvement depending on the nature of the simulated algorithm.

The most powerful communication primitive in PEMS1, *ALLTOALLV*, has been redesigned to use a new message delivery strategy which avoids the need for an area on disk reserved for delivery. The new algorithm thus requires less I/O and disk space to perform the same task. In practice this also eases configuration since the user no longer needs to calculate the message volume of a given algorithm in order to allocate disk space to virtual processors.

Several common collective communication primitives are merely restricted cases of *Alltoallv*, including all of the primitives implemented in PEMS1. However, these can often be implemented much more efficiently than the equivalent call to *Alltoallv*. These primitives, as well as *Alltoallv* itself, have been optimised to eliminate any unnecessary swapping. In particular, with the introduction of multi-core support, “rooted” communication primitives can be implemented more efficiently using appropriate synchronisation techniques. “Rooted” communication primitives are those which send to or receive from some root virtual processor, as opposed to *Alltoallv* in which all processors communicate as equals. This thesis introduces a small set of thread synchronisation primitives that handle swapping in such cases, to ensure a minimal amount of I/O is performed.

Also introduced in PEMS2 is a new type of collective communication function that performs communication as well as computation, unlike those implemented in PEMS1 which perform communication alone. *REDUCE*, and similar methods, are beneficial to certain algorithms since the system can perform the combined communication and computation more efficiently than a user program could by using communication primitives alone. These operations are defined by MPI and used in many BSP algorithms, expanding the useful scope of the implementation.

In addition to these fundamental changes, the implementation has been thoroughly rewritten with the intention of being straightforward to use with existing or new MPI programs on any appropriate system. MPI programs can be compiled against PEMS2 without modification\(^5\), making it straightforward to simulate any existing MPI algorithm for problems sizes vastly exceeding the amount of available main

\(^5\)Assuming, of course, that the program is restricted to the set of calls implemented by PEMS2
memory. All parameters of PEMS2 can be passed at run-time to the program through command line arguments, simplifying automated or manual experimentation. An integrated benchmarking system can record the overall run time of a simulation or a fine-grained breakdown of run-time at each superstep. Benchmark results are written to a gnuplot compatible file which can be used to generate plots like those in this thesis. A comprehensive test suite adapted and augmented from several existing MPI test suites ensures correctness at the application level for any configuration. PEMS2 is freely available on the web [1] under an Open Source license and is straightforward to compile and use on any UNIX system.

1.5 Thesis Outline

The notation, terminology, and variables used throughout are described in Appendix B.

To more thoroughly introduce the reader to the context of this thesis, Chapter 2 describes in detail the approach to EM-BSP simulation taken in PEMSl, and limitations which PEMS2 aims to improve. Subsequent chapters describe how these limitations are addressed in PEMS2: Chapter 3 gives a brief overview of the architecture of PEMS2. Chapter 4 describes the modifications necessary to allow the simulation to take advantage of multi-core processors, including the synchronisation primitives referenced in later chapters. The various styles of I/O available in PEMS2 are described in Chapter 5. The choice of I/O style does not affect the implementation of communication algorithms, but may affect analysis; the consequences of this choice are also discussed in Chapter 5. Chapter 6 describes a new message delivery strategy which differs significantly from that used in PEMSl, using the ALLTOALLV operation as an example.

The communication algorithms presented in Chapter 7 make use of the material in preceding chapters to implement several new communication methods with improved performance characteristics. Chapter 8 then presents several applications built using these methods along with theoretical and experimental performance analysis. This data shows the claims of improvement in previous chapters translate into “real-world” application scenarios.

Finally, Chapter 9 discusses conclusions that can be drawn from PEMS2 results, and suggests potential directions for future work.
Chapter 2

Overview of PEMS1

2.1 Overview

For clarity this overview describes the case where a single real processor is used; the strategy for simulation with multiple real processors is similar, and addressed in detail in later sections.

PEMS1 implements EM-BSP simulation by assigning a thread to each simulated virtual processor. Since there are \( v \) virtual processors, \( v \) threads exist simultaneously. However, only a single thread executes at a given time.

The application, being a BSP-like algorithm, is a series of computation supersteps separated by calls to PEMS communication functions. These functions serve as both communication supersteps and superstep barriers. When one is called, PEMS performs the necessary communication, then swaps the calling thread out of memory. At this point there is a superstep barrier, so the thread yields and another thread is swapped in, which will eventually call the same communication function and reach the same barrier\(^1\). Thus, all threads will eventually synchronise at this barrier and the next superstep can begin. Several barriers may actually be used to implement a collective communication function in PEMS, but there is always at least one as required by the BSP model and its derivatives.

This process of swapping and synchronisation is relatively straightforward to implement. PEMS1 implements allocation using a basic "bump pointer" allocator, which simply allocates memory in a contiguous range by appending new allocations and "bumping" (increasing) the end pointer. Fig. 2.1 shows an example of such an allocation. To swap, this area of memory is read from / written to disk in a single read / write operation, respectively. Whenever a virtual processor is executing, its context is swapped in to the same area of RAM. This ensures the address to a given memory location remains constant so pointers in the application remain valid.

\(^1\)Recall that all virtual processors run identical programs
Implementing communication is more complex. Following the literature associated with PEMS1 [15][16], the communication strategy used is described here using the ALLTOALLV call as an example. ALLTOALLV is the most powerful collective communication method: all others implemented in PEMS1 can be considered simple cases of ALLTOALLV.

2.2 Alltoallv

PEMS1 performs message delivery using a special disk area separate from the virtual processor contexts, called the "indirect area". The indirect area is statically partitioned such that each virtual processor has a dedicated region of some fixed size for message delivery. Alltoallv is performed in two internal supersteps: messages are first written by the sender to the indirect area in a block-aligned and parallel fashion, then read from the indirect area by the receiver and delivered to the receiver's context on disk.

Alg. 2.2.1 shows a straightforward implementation of this approach for a single processor.

Note the algorithm style used here differs slightly from that used in previous work on PEMS [15] and EM-BSP [13][7][6]. The style used here is more implementation directed, omitting lines such as "for i in 0..v - 1 do in parallel" which do not actually occur in this type of multi-threaded code. When reading or analysing algorithms in this style it is best to think from the perspective of a single thread executing the code. For example, all algorithms in this thesis are written from a perspective such as "first,
deliver *my* messages”; not “first, each virtual processor delivers their messages”. The reader must keep in mind that several threads perform these actions simultaneously.

The notation $m_{i\rightarrow j}$ denotes the message sent from virtual processor $i$ to virtual processor $j$.

---

**Algorithm 2.2.1: SIMPLE-ALLTOALLV-SEQ**

**Data:** $S$ : Array of pointers to $v$ messages to send  
**Data:** $\mathcal{R}$ : Array of pointers to $v$ messages to receive

---

1. **Send Messages**
   - foreach message $m_{p\rightarrow i}$ in $S$ do
   1. Write $m_{p\rightarrow i}$ to $i$’s indirect area on disk
   2. Swap out

2. **Finished Internal Superstep 1**
3. **Begin Internal Superstep 2**

4. **Receive messages**
   - foreach message $m_{i\rightarrow p}$ in $\mathcal{R}$ do
   1. Read $m_{i\rightarrow p}$ from indirect area on disk to the $i^{th}$ location in $\mathcal{R}$
   2. Swap out

---

**Finished Virtual Superstep**

---

In the analysis of Alg. 2.2.1 the following variables are used:

- $\omega$ An arbitrary bound on the simulated algorithm’s message size (i.e. the size of a message sent from one virtual processor to another).

- $\mu$ The (maximum) size of a single virtual processor’s context (i.e. the maximum amount of memory allocated by any virtual processor)

- $B$ The size of a disk block

These variables remain free in the stated run times for various communication methods presented in this thesis. Their actual value depends on the characteristics of a particular application or system configuration. All three are assumed to have the same unit, such as bytes.

To convert I/O volume to run time, the following coefficients are used:
$G$ The time required to read / write a single block from / to disk for message delivery

$S$ The time required to read / write a single block from / to disk for swapping

$G$ and $S$ are identical\(^2\), but different coefficients are used to keep terms related to swapping and terms related to message delivery separate. Chapter 4 describes the reasons for this in further detail.

The notation $\lceil \omega \rceil$ means “$\omega$ rounded up to the next multiple of $B$”.

The terms “I/O volume” or “amount of I/O” are used to refer to an amount of I/O in the same unit as $\mu$ and $\omega$, e.g. bytes. Specifically, it does not refer to number of I/O operations in blocks, often referred to as “I/Os” (note plurality) in EM literature. I/O volume is stated separately because later sections in this thesis investigate non-blocked I/O, and comparison of these approaches is best described in terms of volume. This is done only to facilitate discussion and simplify analysis, the total run time of algorithms is given in terms of I/O operations (“I/Os”), as is typical in EM literature.

The notation “$I_a$” is used to refer to the amount of I/O performed by line $a$ of the algorithm. The amount of I/O performed by a range of lines (inclusive) is denoted “$I_{a..b}$”. The same notation used with $T$ rather than $I$ refers to the time taken, rather than the amount of I/O. For example, line 4 in Alg. 2.2.1 refers to “Swap in”, therefore:

$$I_4 = \mu$$
$$T_4 = S \frac{\mu}{B}$$

Recall that all $v$ virtual processors execute the same code. Thus, if a given line in the algorithm performs $x$ I/O, in total the line is responsible for $vx$ I/O, unless the line is conditionally executed by only some virtual processors. To avoid excessive repetition, phrases such as “for each virtual processor” are omitted from proof explanations where it is clear that all virtual processors perform the same actions (as is the case here).

\(^2\)Except with memory-mapped I/O, see §5.2
The variables and notation used here are used consistently throughout this thesis, and documented (with others) in Appendix B.

Lemma 2.2.1. Alg. 2.2.1 (PEMS1 single processor ALLTOALLvd performs $4v\mu + 2v^2\omega$ total I/O.

Proof. For each of the $v$ virtual processors:
The loop at line 1 first writes all $v$ outgoing messages, each of size $\omega$:

$$I_{1..2} = v^2\omega$$

Line 3 swaps out the partition of size $\mu$:

$$I_3 = v\mu$$

Line 4 swaps in the partition of size $\mu$:

$$I_4 = v\mu$$

The loop at line 5 reads all $v$ incoming messages, each of size $\omega$:

$$I_{5..6} = v^2\omega$$

Line 7 swaps out the partition of size $\mu$:

$$I_7 = v\mu$$

Finally, line 8 swaps in the partition of size $\mu$:

$$I_8 = v\mu$$
The total I/O performed by the algorithm, in the unit of $\mu$ and $\omega$ (e.g. bytes), is therefore:

\[
I_{\text{simple-alltoall-seq}} = I_{12} + I_3 + I_4 + I_{56} + I_7 + I_8
= (v^2\omega) + (v\mu) + (v\mu) + (v^2\omega) + (v\mu) + (v\mu)
= 4v\mu + 2v^2\omega
\]

\[\square\]

**Theorem 2.2.2.** Alg. 2.2.1 (PEMS1 single processor ALLTOALLV) takes $S^4\mu/B + G2v^2\lceil\omega\rceil/B + 2L$ time.

**Proof.** Follows directly from Lem. 2.2.1, since Alg. 2.2.1 performs no network communication and no significant computation.

Since messages are delivered one at a time (i.e. each message delivery is a separate I/O operation), message deliveries are each of size $\lceil\omega\rceil/B$ ($\lceil\omega\rceil$ meaning “$\omega$ rounded up to the next multiple of $B$”). Thus, if messages are smaller than a single block then overhead is accumulated for every message. However, this is not a performance problem since a single block of I/O is the minimal amount of time possible for an I/O operation.

There are two internal superstep barriers, contributing $2L$ to the total run time.

\[\square\]

**Theorem 2.2.3.** Alg. 2.2.1 (PEMS1 single processor ALLTOALLV) requires $v\mu + v^2\omega$ disk space.

**Proof.** Each virtual processor requires $\mu$ disk space for its context regardless of message delivery. $v^2$ messages are delivered in total, each of size $\omega$, therefore an additional $v^2\omega$ space is required for the indirect area.

\[\square\]

### 2.3 Potential for Improvement

While experiments with PEMSl have shown desirable scalability characteristics, the system has significant overhead which requires the use of considerably more computational resources to match the performance of comparable EM algorithms. Though
the ability to take advantage of several machines with parallel disks is a considerable
advantage, reducing this overhead will make the system more competitive on a wider
range of systems and problem sizes.

Additionally, the use of a separate area for message delivery introduces scalability
problems with large contexts (see §6.3) and makes tuning difficult in practice since
the user must know in advance the bounds on a given algorithm’s communication
volume in order to allocate disk.

This thesis introduces several new strategies and capabilities for PEMS intended
to address these issues:

As is usually the case with EM algorithms, the most significant source of overhead
in PEMS1 is unnecessary I/O. There are two cases where PEMS must perform I/O:
swapping and message delivery.

2.3.1 Swapping

Each internal superstep barrier in Alg. 2.2.1 implies a swap out and a subsequent
swap in of each virtual processor. However, many such swaps can be avoided by
making more extensive use of the “direct delivery to context” technique described
in the PEMS1 literature [15]. This technique is based on the observation that a
subsequent swap-in in the second internal superstep is not required, since messages
can be written directly to the context on disk. That is, instead of swapping in the
context, modifying it in memory, then swapping the context back out; the message
can simply be written directly to the appropriate location on disk.

Thus, if the second loop delivers messages directly this way, it is not necessary to
swap in at the first barrier. The final swap-out is also avoided because the context
on disk is already known to be consistent\(^3\), hence this avoids 2\(\mu\) I/O per virtual
processor.

Swapping with finer granularity can avoid slightly more unnecessary I/O: the
swap out at the first internal superstep barrier swaps out the entire context, however
this is not necessary. Virtual processors receive messages to some area within their
context, so when the \texttt{Alltoallv} call is completed and control is returned to user code

\(^3\)In fact a swap out can’t occur here because the context is not swapped in, so a swap out would
write garbage data to disk
this region will have been overwritten with the received messages. Therefore, it is not necessary to swap out this region (the "receive buffer") at the initial superstep barrier.

Some swapping can also be avoided at superstep barriers: in a straightforward implementation, all threads swap at superstep barriers. However, for the last thread to execute in the superstep this is not necessary. The order of execution within a superstep is undefined, so it is wasteful to swap out this thread’s context and allow a different thread to swap in and execute first in the next superstep. Instead, the last thread can simply remain swapped in through the barrier and be the first thread to run in the following superstep, thus avoiding one swap per superstep. More generally, in the case of multi-core, threads execute in parallel rounds of \( k \) threads at a time therefore this technique avoids \( k \) swaps per superstep.

### 2.3.2 Message Delivery

Alg. 2.2.1 writes all messages to be delivered to the indirect area on disk. There is potential for improvement here based on two observations:

1. Each message that must be written in the first loop is a part of the sending virtual processor’s context, and therefore will be written to disk regardless at the first barrier (when the sender’s context is swapped out). Thus, the previous algorithm results in each message being written to disk twice.

2. If the receiving virtual processor of a message is local and has already executed this superstep, then the final destination of the message is known and the message can be delivered directly to the destination context on disk. This avoids reading the message from disk again in order to deliver it to the receiver.

There is an additional downside to delivering messages via a separate disk area: because the indirect area is large and separate from the area on disk where contexts are stored, delivery of messages (and subsequently swapping out) involves seeking across a very large area of disk. In the worst case this results in constantly seeking back and forth between the contexts area and the indirect area. This is potentially a serious performance issue, particularly for large \( \mu \) or \( v \). Since disk seeking is extremely expensive, the performance impact of this behaviour could be as significant as the
actual amount of I/O performed – or even more so. The addition of multi-core support compounds the problem due to several threads seeking simultaneously. Reducing or eliminating this effect is therefore a promising path to improving the performance of PEMS in practice.

Of course, indirect message delivery is not done without reason: the messages are aligned and distributed among disks in a way designed to achieve fully parallel disk I/O, and support “direct” I/O which requires all operations to be block aligned. §6.2 describes new methods of retaining these desirable characteristics without writing messages to a separate area on disk.

2.3.3 Communication Balancing

The original EM-BSP simulation algorithms (and PEMS1) require an upper bound on communication volume so disk space can be allocated accordingly for the indirect area. In the multi-processor case, this is achieved by using a deterministic routing technique [2] which first evenly distributes messages across the network before completing the communication. Messages are first sent to an arbitrary intermediary processor in a round-robin fashion, then sent to their final destination by that intermediate processor. Because messages are evenly distributed to an arbitrary intermediate processor, this technique ensures balanced communication.

This technique is straightforward and works well to ensure balanced communication, but in the context of PEMS incurs a large amount of overhead. In order to be delivered, each message must be (in the worst case):

1. Sent over the network (by the sender)
2. Written to disk (by the intermediary)
3. Read from disk (by the intermediary)
4. Sent over the network (by the intermediary)
5. Written to disk (by the receiver, to the indirect area)
6. Read from disk (by the destination, from the indirect area)
7. Written to disk (by the destination, to its context)
The multiple reads and writes of each message to disk, in particular, is a significant amount of overhead due to the large cost of disk I/O.

2.3.4 Allocation

The simple memory allocation scheme used by PEMSl has a serious limitation for many programs: freeing memory is not possible. Since only a pointer to the end of all allocated memory is stored, there is no way to free a particular chunk of allocated memory.

While some BSP-like algorithms allocate a large amount of memory initially then use it throughout execution (such as the PSRS algorithm presented in §8.3), many have more dynamic memory allocation requirements. PEMSl’s basic allocator is not appropriate for algorithms that continuously allocate and free chunks of memory, since memory consumption will continue to increase until available space is exhausted.

2.3.5 Improvements

Chapter 6 presents solutions to the shortcomings described in this section, all of which are implemented in PEMSl2. These improvements depend on two new fundamental aspects of the design: multi-core support and several I/O drivers, presented in Chapter 4 and Chapter 5, respectively.
Chapter 3

Overview of PEMS2

3.1 Software Design

Fig. 3.1 shows an overview of the PEMS2 design.

The most significant change from the more static architecture of PEMS1 is the addition of abstract interfaces for I/O and threading. All use of these subsystems occurs through these relatively simple interfaces, which makes the addition of new I/O and threading drivers to PEMS2 a straightforward process with little impact on other components.

The original I/O (synchronous) and threading (user-space) implementations from PEMS1 have been modified to fit within this framework. Both remain available for use as user options in PEMS2.

Two new I/O drivers have been implemented: Asynchronous I/O, which allows PEMS to submit many I/O requests to the disk at once and resume computation or communication while they are performed, is described in §5.1. Memory-mapped I/O, which allows PEMS to only swap in the required portions of a virtual processor context at each superstep, is described in §5.2.

A new threading driver based on POSIX threads has been added which supports true concurrency, the implications of which are discussed in Chapter 4.
The PEMS2 API provides a public interface to PEMS for applications. It includes a subset of MPI (communication/synchronization) and standard C (allocation).

### MPI API Functions
- MPI_Init()
- MPI_Finalize()
- MPI_Abort()
- MPI_Comm_size()
- MPI_Comm_rank()
- MPI_Barrier()
- MPI_Scatter()
- MPI_Gather()
- MPI_Bcast()
- MPI_Allgather()
- MPI_Allgatherv()
- MPI_Alltoall()
- MPI_Alltoallv()
- MPI_Reduce()
- MPI_Scan()
- MPI_Barrier()
- malloc()
- realloc()
- free()

### IO Interface
- allocPartition()
- freePartition()
- read()
- write()
- sync()

### STXXL Library
- STXXL (Asynchronous I/O)
- STXXL Library

### Mmap Library
- Mmap (Memory Mapped I/O)

### UNIX Library
- UNIX (Synchronous I/O)

### Pthreads Library
- Pthreads (System Threads)

### GNU Pth Library
- GNU Pth (Userspace Threads)

Figure 3.1: PEMS2 Design
3.2 Computational Model

PEMS2 extends the EM-BSP [13][7][6] models shown in Fig. 1.2 with one or more "cores" per real processor. Each set of cores on a real processor access a single shared main memory, and one or more disks. The cluster of real processors is assumed to be homogeneous, i.e. each real processor has $k$ cores and $D$ disks. This extended model is shown in Fig. 3.2.

Adding the ability for threads to execute concurrently is a relatively straightforward modification to PEMSl (replace the use of GNU Pth functions with POSIX threads equivalents). The difficulty in adding multi-core support lies in the implications, e.g. more sophisticated synchronisation and inter-thread communication methods must be used, and the relevant portions of the system must be made thread-safe. The details of how this has been accomplished are discussed in Chapter 4.

![Figure 3.2: PEMS2 Computational Model](image)

Example: $k = 4$ Cores/Processor, $D = 2$ Disks

Figure 3.2: PEMS2 Computational Model
Chapter 4

Multi-Core Support

PEMS1 supported only user-space threads via the GNU Pth library. On single-core machines this can be advantageous because user-space threads avoid the overhead of context switching. However, achieving true concurrency on a multi-core machine requires the use of “real” system threads.

While it is possible to run several MPI processes concurrently on a single multi-core machine, running a single process with a thread for each virtual processor avoids the overhead of inter-process communication, synchronisation, and context switching. Using threads also allows for more effective parallel disk I/O strategies since PEMs can control parallel access to disk(s) in more flexible ways.

To achieve this, the threading system has been redesigned around a small set of simple synchronisation primitives which are safe for both user-space and kernel threads. PEMs2 can use either user-space threads via GNU Pth, or system threads via the POSIX Threads (“pthreads”) API. In either case there is a 1 : 1 relationship between threads and virtual processors regardless of the number of cores available.

The number of virtual processors that execute concurrently on a local real processor is denoted $k$. The user may choose any value for $k$ provided $1 \leq k \leq \frac{p}{k}$.

4.1 Memory Partitions

Thread concurrency in PEMs2 is achieved by allocating $k$ separate memory partitions (rather than the single partition used by PEMs1). Thus, $k$ separate threads may be swapped in at a given time and perform work concurrently. The user must ensure that $k \mu$ real memory is available for these partitions.

A simple static mapping is used to assign threads to memory partitions: thread $t$ uses partition $t \mod k$. A dynamic mapping would be beneficial in many respects, but this would have the effect of changing the address of a given piece of virtual processor memory, thus invalidating pointers. For example, if a virtual processor is
swapped in at memory address 10, a pointer to the first allocated piece of memory would have the value 10. If that virtual processor was subsequently swapped in at memory address 20, the pointer should have value 20, but still has value 10, thus memory is corrupted. Because of this, a dynamic mapping of contexts on disk to memory partitions is not feasible within PEMS.

4.2 Controlling Concurrency

With the addition of system thread support, several virtual processors may execute in parallel on a single real processor, taking advantage of multiple cores. This raises an issue when $\frac{n}{p} > k$ (which is generally the case for any reasonable configuration): threads can run concurrently, thus more than $k$ threads may attempt to run simultaneously. However, only $k$ memory partitions are available. PEMS itself can not explicitly schedule threads $k$ at a time since the operating system scheduler is used. Instead, an exclusive lock (mutex) is associated with each of the $k$ partitions in main memory. A thread must obtain a lock on its memory partition before executing any part of the simulated virtual processor's algorithm. Therefore, the number of virtual processors which can run concurrently on a single real processor is at most $k$.

4.3 Thread Synchronisation

Superstep synchronisation, a simple barrier, is sufficient for collective communication methods in which all processors participate as equals. However, there are many methods which have more complex synchronisation requirements. A simple example of such a method is the broadcast, or Bcast. In a Bcast, a single virtual processor called the “root” sends a message to every other virtual processor (see §7.2). Thus, other virtual processors have to wait specifically for the root to perform some action. While full superstep barriers could be used for this purpose, synchronisation methods specifically designed for such cases can achieve better performance. With a full barrier, each virtual processor waits for every other virtual processor. However, in a “rooted” case such as Bcast, any virtual processor that reaches the barrier after the root need not wait at all. Since I/O is triggered by virtual processor execution (e.g.
I/O will occur when a virtual processor calls MPI_Bcast), this can be a significant performance factor – the sooner a thread passes a barrier, the sooner it can submit further I/O requests, resulting in higher throughput and more communication/computation overlap.

The collective communication algorithms presented here require three styles of synchronisation (in addition to superstep barriers):

1. **Initial Synchronisation**: Wait for the first thread
2. **Rooted Synchronisation**: Wait for a specific “root” thread
3. **Final Synchronisation**: Wait for all other threads

These operations are implemented to work with any number of threads running at a time, swapping virtual processors in or out as required.

Since each thread holds its memory partition lock while executing, and other inactive threads require the same partition, simply using a primitive signal (e.g. that provided by pthreads) would result in a deadlock and/or missed signals. This is because primitive signals are not persistent, i.e. only those threads waiting on a signal at the moment it fires are notified. In PEMS2, a primitive signal with an associated counter and flag make up a composite synchronisation structure. This allows for synchronisation both between threads which are currently swapped in (via the primitive signal) and threads which are not (via the counter or flag).

The primitive signal is only used to synchronise the \( k \) currently swapped in threads, eliminating the possibility of deadlock. The counter keeps track of how many threads have reached the synchronisation barrier, and the flag is used to signal an arbitrary condition (e.g. “the root has finished”).

This composite signal structure is simply referred to as a “signal”; which is the main threading abstraction used to implement our synchronisation primitives.

All functions described in this section are called while the thread holds the lock on its memory partition. Because swapping is generally the most expensive operation performed by PEMS during a simulation, the goal of these primitives is to swap only when necessary. Run times stated for these methods only consider time spent performing I/O, since no significant computation takes place.
4.3.1 Rooted Synchronisation

Alg. 4.3.1 EM-WAIT-FOR-ROOT waits for the root thread to signal. This is generally necessary for any rooted collective communication method (e.g. Bcast, Gather). Swapping to disk occurs only when a thread is blocking the memory partition required by the root. The return value indicates whether the partition has been swapped out, which allows the caller to only swap in/out again if necessary. Only the non-root threads call this function; the root thread must perform whatever work is required, then signal (using Alg. 4.3.5, EM-SIGNAL-THREADS) to unblock the other threads.

Algorithm 4.3.1: EM-WAIT-FOR-ROOT

<table>
<thead>
<tr>
<th>Data:</th>
<th>s (signal), t (this thread ID), r (root thread ID)</th>
<th>t ≠ r</th>
</tr>
</thead>
<tbody>
<tr>
<td>Result:</td>
<td>True iff thread was swapped out</td>
<td></td>
</tr>
</tbody>
</table>

1. result ← false
2. s.lock()
3. if s.flag = false then — If the root has not already signalled
   4. pt ← t mod k — Current thread’s partition
   5. pr ← r mod k — Thread r’s partition
   6. if pt = pr then — If t and r share a partition
      — Yield to root —
      7. result ← true
      8. Swap out
      9. Unlock partition
   10. s.wait() — Wait for root to signal
   11. if pt = pr then — If t and r share a partition
      — Yielded above, so re-lock partition —
      12. s.unlock() — Release signal lock to prevent deadlock
   13. Lock partition
   14. s.lock()
15. s.count ← s.count + 1
16. if s.count = \(\frac{n}{p}\) then — If all non-root threads are finished waiting
   — Reset signal —
   17. s.count ← 0
18. s.flag ← false
19. s.unlock()
20. return result

Lemma 4.3.1. Alg. 4.3.1 takes \(S\frac{\nu}{P\cdot k\cdot B}\) time in the worst case.
Proof. The only possible I/O occurs at line 8, which is only executed by virtual processors which share a memory partition with the root processor. There are \( k \) partitions per real processor, shared by \( \frac{n}{p} \) virtual processors, thus \( \frac{n}{pk} \) virtual processors may perform I/O. If a virtual processor performs I/O, it swaps out once at line 8, resulting in \( \mu \) I/O per virtual processor that shares a partition with the root. Since all virtual processors that perform I/O share a memory partition, only one may be swapped in at a given time, therefore no disk parallelism occurs in the worst case when striping is not in use.

Note that Lemma 4.3.1 does not take disk striping into consideration, i.e. it is assumed that each virtual processor is mapped to a single disk. If PEMS is being used on a configuration where all data is striped across all disks, then all I/O is inherently fully parallel, and therefore Alg. 4.3.1 would take \( S \frac{n}{FkBd} \) time.

### 4.3.2 Initial Synchronisation

Implementations of several collective communication functions require an arbitrary single thread to do some work (e.g. perform MPI communication) before any other threads continue. Alg. 4.3.2 (EM-FIRST-THREAD), when called by all threads, will return true immediately if the caller is the first thread, or otherwise block until the first thread has signalled and return false. Note that when true is returned the signal is still locked; this allows the first thread to perform the necessary work while other threads wait. The first thread must signal (using Alg. 4.3.5 with false as the “lock” parameter) when it has completed the work in order to wake any waiting threads.

**Lemma 4.3.2.** Alg. 4.3.2 performs no I/O.

### 4.3.3 Final Synchronisation

Collective communication calls which collect data at a single root processor (e.g. Gather) must wait for other threads to finish their work before the results can be gathered and delivered to their final destination. Alg. 4.3.3 (EM-ALL-THREADS-FINISHED) along with Alg. 4.3.4 (EM-WAIT-THREADS) provides the required mechanism. If true is returned, all threads have reached the call and the work may be safely performed. Whether or not a swap has occurred is passed as an input/output
Algorithm 4.3.2: EM-FIRST-THREAD

Data: $t$ (this thread ID), $s$ (signal)
Result: True iff caller is the first thread

1. s.lock()
2. if $s$.count = 0 then \hspace{1em} \textit{If this is the first thread}
   \hspace{1em} \textit{Keep lock and return true}
   \hspace{1em} s.flag \leftarrow false
   \hspace{1em} \textbf{return} true
3. \hspace{1em} s.count \leftarrow (s.count +1) \mod \frac{p}{v}
4. if s.flag = false then \hspace{1em} \textit{If first thread has not finished}
   \hspace{1em} s.wait()
5. if s.count = 0 then \hspace{1em} \textit{If this is the last thread}
   \hspace{1em} \textit{Reset signal}
   \hspace{1em} s.flag \leftarrow false
6. s.unlock()
7. \textbf{return} false

Parameter (e.g. a pointer) to allow cascading several calls without performing unnecessary swaps: if true is passed for this parameter, no swap will be performed. Otherwise, if a swap is performed, the parameter will be set to true to notify the caller.

Like Alg. 4.3.2 (EM-FIRST-THREAD), if false is returned the lock is not released. When this happens the caller must call Alg. 4.3.4 (EM-WAIT-THREADS) which will block until all threads have completed.

Lemma 4.3.3. Alg. 4.3.4 performs at most $v\mu$ I/O.

Proof. The only I/O performed is a swap out of size $\mu$, which is called $v$ times in the worst case (once by each virtual processor).

\section*{4.3.4 Signalling}

Both Initial and Rooted synchronisation require a thread to signal the others once some work has been performed. Alg. 4.3.5 (EM-SIGNAL-THREADS) is used for this purpose in both cases. Since these cases have different locking semantics, whether the signal lock should be taken is passed as a parameter (specifically: false must be passed in the Initial case, and true in the Rooted case).
Algorithm 4.3.3: EM-ALL-THREADS-FINISHED

Data: $t$ (this thread ID), $s$ (signal), $w$ (whether swap has occurred)

Result: True iff the caller is last

```plaintext
1. result ← true
2. last ← false
3. s.lock()
4. if s.count = $\frac{n}{p} - 1$ then — If this is the last thread
   - Signal others, reset signal and return true —
   5. s.count ← 0
   6. s.broadcast()
   7. s.unlock()
   8. return true
9. else — This is not the last thread
10. s.count ← (s.count +1) mod $\frac{n}{p}$
11. if s.flag = true then — If the last thread has not already finished
12.   if w = false then — If this thread hasn't already swapped out
13.      — Swap out and notify caller —
14.      Swap out
15.      w ← true
16.   — Wait for last thread to finish —
17.   Unlock partition
18.   s.wait()
19.   Lock partition
20. if s.flag = true then — Last thread has finished
21.   s.unlock()
22. else — This thread is blocking the last thread
23.   — Keep lock and return false —
24.   return false
25. return result
```
Algorithm 4.3.4: EM-Wait-Threads

**Data:** \( s \) (signal), \( w \) (whether swap has occurred)

1. **if** \( w = \text{false} \) **then** — *If this thread hasn’t been swapped out yet*
   2. Swap out
   3. \( w \leftarrow \text{true} \) — Yield partition and wait for signal —
4. Unlock partition
5. \( s\text{-wait}() \)
6. Lock partition
   — *Reset signal* —
7. \( s\text{-flag} = \text{false} \)
8. \( s\text{-count} = 0 \)
9. \( s\text{-unlock}() \)

Algorithm 4.3.5: EM-Signal-Threads

**Data:** \( t \) (this thread ID), \( s \) (signal), \( l \) (whether to lock)

1. **if** \( l = \text{true} \) **then**
2. \( s\text{-lock}() \)
3. \( s\text{-count} \leftarrow (s\text{-count} + 1) \mod \frac{p}{k} \)
4. \( s\text{-flag} \leftarrow \text{true} \) — *Set flag for threads yet to run*
5. \( s\text{-broadcast}() \) — *Signal the \( k-1 \) other currently running threads*
6. \( s\text{-unlock}() \)
Chapter 5

New I/O Drivers

5.1 Asynchronous I/O

5.1.1 Background

The UNIX system I/O used by PEMSl is synchronous, i.e. a call to read or write blocks until the I/O operation has finished. In some cases this is necessary because execution can not continue until I/O is finished, typically because the buffers used are required for the next operation. In other cases, however, there is useful work that can be safely performed in parallel with the I/O operation. In these cases, asynchronous I/O is advantageous. Asynchronous I/O allows an I/O request to be submitted with a non-blocking call, and provides a separate mechanism to wait for completion. This allows I/O to proceed in parallel with computation, improving overall performance.

An additional benefit of asynchronous I/O is the ability to send many I/O requests to the operating system (OS) at once. With synchronous I/O, this is not possible because all I/O requests block. Asynchronous I/O, however, allows submitting many requests in rapid succession, keeping the OS and disk busy with I/O requests. This is beneficial because the OS attempts to schedule disk I/O optimally when several requests are pending. Several algorithms exist for this purpose which yield better performance than a trivial First Come First Served (FCFS) algorithm [21]. All modern commonly used operating systems include at least one disk scheduling algorithm; Linux in particular provides several which may be selected at runtime for a specific disk volume (see §9.1).

5.1.2 Design

PEMS2 uses the STXXL [10] file layer for asynchronous I/O. This is the lowest level abstraction in STXXL, essentially a portability layer for asynchronous I/O with a more elegant interface than the operating system’s API. The scheduling and caching
mechanisms in other layers of STXXL are not used (see §1.3.1 for additional discussion of STXXL).

The non-trivial modifications to PEMS required for asynchronous I/O are concerned with waiting for the necessary I/O requests to finish. All I/O in PEMS is performed by some virtual processor, and written to / read from the context of another virtual processor. It is important that threads only wait when necessary to avoid blocking other threads which could otherwise proceed. Generally, the thread that initiated the I/O request is the only thread that should wait. Accordingly, PEMS2 has \( k \) independent I/O request queues per real processor, one for each local virtual processor that is swapped in. Each virtual processor can make multiple I/O requests (e.g. during message delivery) and explicitly wait for all, or some, of its own requests to finish if necessary. Otherwise, all requests are waited on at the next superstep barrier before the virtual processor is swapped out.

5.2 Memory Mapped I/O

5.2.1 Background

The I/O approaches previously discussed (both synchronous and asynchronous) have a major disadvantage for certain algorithms: at each virtual superstep, the entire context of every virtual processor is swapped regardless of how much data the algorithm actually uses. In cases where the algorithm only accesses a small portion of the data (e.g. sampling) this can result in a very large amount of unnecessary I/O. This can cause the I/O complexity of the simulation to be far from optimal, particularly for algorithms with many supersteps each of which do not access the majority of memory. This problem cannot be solved with explicit I/O (i.e. read/write calls) because PEMS has no way of knowing which areas of memory are actually used by the simulated algorithm.

Special API calls could be added to PEMS to address this problem, but this conflicts with the goal of simulating generic BSP-like algorithms, and would not be compatible with MPI. Fortunately, there is a mechanism available in all modern operating systems which can solve this problem: memory mapped I/O. Memory mapping is a facility which allows a file (or other addressable resource) to be mapped
onto a range of virtual memory and used normally like any other region of memory, without actually reading the entire file into physical memory. Pages are swapped to/from disk by the OS as necessary without any effort on behalf of the programmer.

The critical property of memory mapped I/O is that this page swapping is performed by the OS kernel which, unlike "userland" code such as PEMS, does know which areas of memory are accessed. This allows PEMS to avoid unnecessary swapping, since the kernel will only swap in/out those regions of memory which are actually used by the simulated algorithm. This implies the cache behavior of the algorithm may also affect I/O performance – algorithms with favourable memory access patterns will make use of the kernel-managed cache more effectively, and achieve better performance with memory mapped I/O.

Experiments in §8.4 confirm experimentally that memory mapping avoids a significant amount of I/O in some cases.

5.2.2 Design

When used with memory mapping, PEMS2 simply maps the entire used portion of disk into memory. Rather than allocate in-memory partitions and swap in/out from/to disk, the simulated algorithm works directly with a range of this mapped memory. All other aspects of the simulation remain the same, in particular, only \( k \) virtual processors execute at a given time. If suitable parameters are chosen such that \( k\mu \) fits within physical memory (as it must with explicit I/O), this ensures that the amount of virtual memory used at any given time fits within physical memory, so thrashing is avoided.

Because memory-mapped disk regions are used in the same way as any other region of memory, message delivery in PEMS2 with memory-mapped I/O is simply a direct virtual memory copy (e.g. using `memcpy`). In degenerate cases where the problem size is smaller than the available physical memory, this effectively makes PEMS an in-memory multi-core MPI system. This allows PEMS to scale gracefully over a wide range of problem sizes from very small, to the majority of physical memory, to much larger than physical memory.
Chapter 6

Simulation Enhancements

6.1 Swapping

A straightforward implementation of many communication algorithms could perform many complete swaps in a virtual superstep (i.e. a superstep in the simulated algorithm), since virtual supersteps may be composed of several internal supersteps (i.e. a superstep performed by PEMS). A careful implementation, however, can ensure that each virtual processor is completely swapped out and completely swapped in only once per virtual superstep. Thus, for explicit I/O, \( L \geq S^{2p_B} \).

With the use of memory mapped I/O, supersteps cause no explicit I/O at all. In this case the analysis of a simulated algorithm must take into consideration any swapping I/O it would cause by accessing its own memory mapped partition. Because of this generic bounds for a PEMS simulation using memory mapped I/O can not be given, the analysis is specific to a particular algorithm.

6.2 Message Delivery

This section introduces a new communication strategy for PEMS which addresses the limitations discussed in §2.3. For illustrative purposes, the basic concept is first presented in the form of a simplified algorithm, Alg. 6.2.1, which does not consider details such as block alignment.

Message delivery in PEMS2 is discussed using Alltoallv as an example. Rather than write/read messages to a separate area on disk as in PEMS1, all virtual processors record in a table where in their contexts they expect to receive incoming messages. Then, they deliver directly to other virtual processors' contexts on disk. Thus the additional communication area on disk (and with it a significant amount of I/O and disk seeking) is eliminated.

\(^1\)Note that \( L \) in this thesis differs from previous work on PEMS, see Appendix B.4
**Algorithm 6.2.1: SIMPLE-DIRECT-ALLTOALLV**

Data: $S$: Array of pointers to $v$ messages to send

Data: $R$: Array of pointers to $v$ messages to receive

1. Let $T$ be a shared $v \times v$ table of incoming message offsets

   — *Store message offsets* —

2. Store incoming message offsets from $R$ in $T$

3. Swap out

   — *Finished Internal Superstep 1* —

   — *Begin Internal Superstep 2* —

   — *Deliver messages* —

4. Swap in

5. foreach message $m_{p-i}$ in $S$ do

6. \[ Write\ m_{p-i}\ to\ \bar{T}_{p-i}\]

7. Swap out

   — *Finished Virtual Superstep* —

This strategy avoids out-of-place message delivery, but is not an ideal solution for two reasons: I/O operations are not necessarily block aligned, and messages are written to disk and read again in cases where this can be avoided.

One possible approach to eliminate alignment issues is to simply use buffered I/O. However, the caching and copying inherent to buffered I/O is not suitable for a system like PEMS which consumes as much main memory as possible. PEMS1 resolved this by organizing all message data in an appropriate way in a separate area on disk. PEMS2 instead directly delivers the largest aligned portion of a message possible, and keeps a cache of remaining blocks which require “cleaning up”. The key observation is that for a given message, a maximum of 2 blocks may not be properly aligned (namely the first and last block of the message$^2$). Since each virtual processor receives $v$ messages, a given virtual processor must receive at most $2v$ unaligned blocks – dramatically less than the total message volume for a typical coarse-grained algorithm.

The overhead inherent in buffered I/O is due to the fact that a portion of a block can not physically be written to disk. For example, if the first half of a block must

---

$^2$Note that while messages may be distributed in any way across the context, an individual message is a contiguous range
be written, the contents of the second half must be in memory so the complete block

be assembled and written to disk. Therefore, some sort of cache is required to

avoid corrupting blocks when a partial block is written. We will need to emulate this

behavior to achieve our goal, but are able to do so more efficiently than the generic
cache mechanism in the kernel since we know precisely what the kernel must guess$^3$.

The general solution to this problem is trivial: simply read in the desired block

from the destination context, modify it, and write it back out again. Unfortunately

this is not sufficient for our purposes since two (or more, in cases with very small
messages) messages can overlap a single block, which raises synchronisation issues
when $k > 1$. The overhead associated with a read/write cycle is also undesirable.

Instead, we will cache the blocks containing unaligned messages ends ("boundary
blocks") in memory throughout the course of the Alltoallv call. As virtual processors
deliver the bulk of their messages directly to their destinations, they update this cache
with the remaining fragments of the delivered messages. Since this is done when the
relevant contexts are already swapped in, the read/write cycle is avoided. Finally,
when the bulk of all messages have been delivered each processor flushes the necessary
boundary blocks from the cache in memory to its context on disk and the algorithm
completes.

Another issue arises with the direct delivery of messages in the absence of buffered
I/O: while the source and destination of each message contain aligned regions of equal
size, these regions may not have equivalent alignment (i.e. their start offsets are not
equivalent mod $B$). Fig. 6.1 illustrates such a case (the top and bottom regions rep-
resent the sender’s and receiver’s contexts, respectively). The largest aligned region
within the message source does not correspond directly to the largest block-aligned re-
gion within the message destination because their alignment differs. This is a problem
because non-buffered I/O requires that all offsets be block aligned, both in memory
and on disk. To resolve this we take advantage of the fact that the source context is
both in memory and on disk at write time, so we can destroy the context in memory
and avoid swapping out to prevent corruption. We shift each message in memory left-
ward so the regions in the sender and receiver align properly (Step 1 in Fig. 6.1). Thus
aligned, these regions can be delivered directly, and the remainder of the message is

$^3$"There are only two hard problems in Computer Science: cache invalidation and naming things.”
– Phil Karlton
handled by the boundary block cache in memory.

![Figure 6.1: Direct Message Delivery](image)

### 6.3 Disk Space Reduction

In addition to reducing the volume of I/O performed, the elimination of the indirect area significantly reduces the amount of disk space required to run a given simulation, particularly with large numbers of virtual processors. This allows a given system configuration to handle larger problem sizes.

In PEMSl, each real processor required $\frac{v\mu}{p}$ disk space for its local virtual processor contexts, and $v\mu$ disk space for the indirect area. Note that the size of the indirect area increases with $v$ rather than $\frac{v}{p}$. This has the effect of increasing disk space when real processors are added even if $\frac{v}{p}$ remains constant, which can be a significant scalability problem. The ideal strategy for scaling up a PEMS simulation is to determine the parameters that fully utilize the resources available to a single machine, then be able to easily add real processors as necessary to reach the desired problem size. An area on disk that scales with $v$ rather than $\frac{v}{p}$ conflicts with this concept. In practice this makes scaling more tedious than necessary, since predicting the amount of required disk space is more difficult. Additionally, as experiments in §8.3.3 show, this large
region of disk space can incur a serious performance penalty when $\mu$ is large, due to both disk seek time and file system overhead.

During the course of a simulation the disk is continually reading and writing for swapping and message delivery. As a result the disk head must constantly seek across the entire region of disk space, including, in PEMS1, the huge indirect area. This occasionally had the counter-intuitive effect of making the simulation slower when more RAM was added and $\mu$ correspondingly increased, because the disk seek time dwarfed the time spent actually swapping a given context.

To solve this problem, the improved simulation algorithms introduced in this thesis eliminate the indirect disk area entirely, so the amount of disk space required per real processor is precisely $\frac{\mu}{p}$. As a result, real processors can be added to increase problem size without increasing the disk space requirement for each real processor. In practice, this makes tuning a PEMS simulation much more manageable.

Fig. 6.3 illustrates the difference in disk space consumption between the two strategies. Even for a modest $\frac{\mu}{p}$ the disk space requirements for PEMS1 rapidly increase; in this case with 16 processors the disk space required of a single real processor exceeds the total problem size. PEMS2, in contrast, only uses disk for virtual processor contexts, so the amount of disk space required precisely matches the problem size regardless of how many real processors are added.

\[
\begin{array}{ccccccc}
 p & v & \text{Required} & \text{PEMS1/Proc} & \text{PEMS1} & \text{PEMS2/Proc} & \text{PEMS2} \\
 1 & 8 & 16 \text{ GiB} & 32 \text{ GiB} & 32 \text{ GiB} & 16 \text{ GiB} & 16 \text{ GiB} \\
 2 & 16 & 32 \text{ GiB} & 48 \text{ GiB} & 96 \text{ GiB} & 16 \text{ GiB} & 32 \text{ GiB} \\
 4 & 32 & 64 \text{ GiB} & 80 \text{ GiB} & 320 \text{ GiB} & 16 \text{ GiB} & 64 \text{ GiB} \\
 8 & 64 & 128 \text{ GiB} & 144 \text{ GiB} & 1152 \text{ GiB} & 16 \text{ GiB} & 128 \text{ GiB} \\
 16 & 128 & 256 \text{ GiB} & 272 \text{ GiB} & 4352 \text{ GiB} & 16 \text{ GiB} & 256 \text{ GiB} \\
\end{array}
\]

\[
\left(\frac{\mu}{p} = 8, \mu = 2 \text{ GiB}\right)
\]

Figure 6.2: Disk Space Requirements

6.4 Communication Buffer Size

Due to the removal of the indirect message area, PEMS2 does not require an upper bound on communication volume in order to allocate disk space, but communication
volume must still be bounded in many cases to avoid exceeding the available communication buffer. Note that one bound on communication volume is inherent: each virtual processor can send at most $\mu$ data in total, since each virtual processor has $\mu$ memory and messages must reside in that memory before being sent.

Each virtual processor sends at most $v$ messages in a communication superstep. The user may configure how many of these messages are sent at once using the parameter $\alpha$. By choosing an appropriate value for $\alpha$, the user may ensure there is always sufficient buffer space to handle communication. This strategy removes the need for indirect routing as in PEMSl (see §2.3.3). Performance is therefore improved since each message is sent over the network precisely once, to its destination real processor.

The amount of I/O performed by communication methods depends on the size of messages. To represent this in analytical results, the variable $\omega$ is used to represent an arbitrary bound on virtual message size. Specific values may be substituted for $\omega$ to find the run time for a particular call, or a particular computational model. For example, if a Bcast is performed where the message is simply a single 32-bit integer, $\omega = 4$ bytes; for a CGM algorithm, $\omega = \Theta(\frac{N}{v})$; etc.

### 6.5 Scheduling and Disk Parallelism

If each virtual processor is mapped to a single disk (i.e. striping or similar techniques are not in use), the runtime of a collective communication method depends on the order of execution of virtual processors. This is because virtual processors execute in synchronised rounds $k$ at a time, where each round includes a single virtual processor mapped to each memory partition ($0 \ldots k - 1$). However, this does not automatically imply that each round contains a virtual processor mapped to each disk. In the worst case, only a single disk may be used despite several disks being available. Fig. 6.3 shows such a case: if the virtual processors shown in bold (0, 4, and 8) are executed in a round, only disk 0 is used for that round and thus disk parallelism is not exploited.

This problem must be addressed in order to precisely analyse the communication functions in PEMSl and applications built with them. Restrictions on $k$ and $D$ could solve the problem, but this approach is not realistic since $k$ and $D$ reflect physical system characteristics. Defining the scheduler’s behaviour such that these situations
are avoided is more flexible, and feasible to implement in practice. Conveniently, a trivial scheduling algorithm results in the desired behaviour: if virtual processors are executed in ID order, then message delivery is distributed across all disks. For example, with $k = 3$ as in Fig. 6.3, processors 0, 1, 2 would execute in the first round, 3, 4, 5 in the next round, etc. If $k \geq D$, then clearly each round uses $D$ disks in parallel (since an increasing sequence of $k$ integers mod $D$ contains all integers in $0 \ldots D - 1$ if $k \geq D$). If $k < D$, then this is not the case, and virtual processor contexts should be distributed across disks to exploit disk parallelism.

When each virtual processor context is distributed across disks, all disk I/O of sufficient size is fully parallel, so the scheduler behaviour need not be defined and no restriction is required of $k$ and $D$. In this case, a separate restriction is necessary: individual reads and writes must be large enough that they will be performed across all $D$ disks. With a straightforward round-robin block distribution strategy as used by striped RAID systems and the STXXL block layer, this requires $\omega \geq BD$ for fully parallel message delivery, and $\mu \geq BD$ for fully parallel swapping. For any reasonable configuration, $\mu \gg BD$. $\omega$ may be $< BD$, but if this is the case messages are so small that full disk parallelism is impossible (since disks can not perform transfers smaller than $B$), so we will simply assume $\omega \geq BD$ to simplify analysis.

Def. 6.5.1 summarises these conditions.

**Definition 6.5.1** (Fully Parallel Swapping). If each virtual processor context resides on a single disk, $k \geq D$, and virtual processors are scheduled in increasing order by ID, then PEMS2 performs all swapping I/O across all $D$ disks in parallel.
If each virtual processor context is distributed evenly across all disks in a blockwise fashion, and $\mu \geq BD$, then PEMS2 performs all swapping I/O across all $D$ disks in parallel.

Unfortunately, this behaviour conflicts with the potential swapping optimisation described in §2.3.1 where $k$ swaps can be avoided at each virtual superstep barrier. Accordingly, the run times given in this thesis do not include that optimisation.

6.6 Allocation

When PEMS is initialised it first allocates all memory required by virtual processors. It then intercepts allocation requests from the simulated algorithm and satisfies them by allocating the requested memory from this pool.

To fully support dynamic memory allocation and deallocation, PEMS2 uses a more sophisticated allocation scheme than PEMSl (see §2.3.4). All virtual processor memory is still contained within a single region of size $\mu$. Unlike PEMSl, however, PEMS2 stores the offset and size of each allocation. This enables freeing of allocated memory, which can then be reused by future allocations.

The allocation records are stored using a simple balanced binary search tree in memory. Since the number of allocations is relatively small and the overhead of this data structure is not significant compared to disk I/O, a more sophisticated structure would not likely show any significant improvement.

The allocation algorithm is simple: search from the lowest address until a large enough free chunk is found, then split the start of this chunk into a newly allocated area of appropriate size.

Deallocation is also straightforward: remove the allocated chunk, and merge with any adjacent free chunks. If there are no adjacent free chunks, simply record the area as deallocated.

More sophisticated strategies are of course possible; efficient allocation with minimal fragmentation is a much researched problem. In the context of PEMS, however, the most important benefit of an allocator over the basic design of PEMSl is the ability to re-use deallocated memory, and avoid I/O for currently unallocated memory regions. The relatively simple allocator presented here, though not optimal with respect to fragmentation, does provide these two advantages.
The swapping related I/O function in PEMS2 have been modified to only swap currently allocated regions of memory, rather than swap the entire partition in a single read/write operation as in PEMS1. As a result, programs which free memory as soon as possible see improved performance due to less I/O. For programs with very dynamic allocation behaviour, this can amount to a significant reduction in I/O and total run time compared to the PEMS1 strategy.
Chapter 7

New and Improved Communication Algorithms

7.1 Alltoallv

In an ALLTOALLV, every virtual processor sends a message of arbitrary size to every other virtual processor, thus \( v^2 \) messages are exchanged in total. ALLTOALLV is the most powerful collective communication operation implemented in PEMS that performs only communication (i.e. new values are not computed as part of the operation).

Due to the complexity and size of the EM-ALLTOALLV algorithm, the single-processor and multi-processor versions are presented here separately. These are referred to as EM-ALLTOALLV-SEQ and EM-ALLTOALLV-PAR, respectively. The algorithm in general (i.e. for both single processor and multi-processor cases) is referred to as EM-ALLTOALLV. Note the implementation makes no such distinction and simply provides an implementation of the MPI_Alltoallv function that works in both cases.

![Diagram of Alltoallv Operation](image.png)

Figure 7.1: Alltoallv Operation
7.1.1 Single Processor

Algorithm

Alg. 7.1.1 describes the single processor implementation of Alltoallv in PEMS2. Note that all I/O (including swapping) is explicitly performed, superstep barriers do not imply swapping. This algorithm and the others in this section perform fine-grained swapping, e.g. “Swap message in” means the message (which resides in the virtual processor’s context) should be swapped in from disk to its usual location in the memory partition, just as if the entire partition was swapped in.

The reader is encouraged to review the simpler implementations of Alltoallv (Alg. 2.2.1 and Alg. 6.2.1), since the algorithm given here solves the same problem in a similar but more intricate way.
Algorithm 7.1.1: EM-ALLTOALLV-SEQ

Data: \( S \) : Array of pointers to \( v \) messages to send
Data: \( R \) : Array of pointers to \( v \) messages to receive

1. Let \( T \) be a shared \( v \times v \) table of incoming message offsets
2. Let \( E \) be a shared array of \( v \) execution states, all initially false
3. Let \( M \) be a cache of at most \( 2v^2 \) border blocks (2\( v \) per virtual processor)
   - Store message offsets and synchronise —
4. Swap out everything except regions in \( R \)
5. Store incoming message offsets from \( R \) in \( T \) — \( T_{s \rightarrow p} \) is valid
6. Set \( E_p \) to true — This thread has reached this point
7. Synchronise with the \( k-1 \) other currently running threads
   - Deliver messages if possible —

8. foreach message \( m_{p \rightarrow i} \) in \( S \) do
   9.   Update \( M \) with the start and end of this message
   10.  if \( E_i \) is true then — Thread \( i \) has recorded its offsets in \( T \)
      11.    Align and deliver directly to \( T_{p \rightarrow i} \) on disk

   — Finished Internal Superstep 1 —
   — Begin Internal Superstep 2 —

   — Deliver remaining messages —

12. foreach message \( m_{p \rightarrow i} \) in \( S \) not delivered in superstep 1 do
13.    Swap message in
14.    Align and deliver directly to \( T_{p \rightarrow i} \) on disk

   — Finished Internal Superstep 2 —
   — Begin Internal Superstep 3 —

   — (Blocked I/O only) Flush border block cache —

15. Flush border blocks in \( M \) to our context
    — Finished Virtual Superstep —
Analysis

Similar disk parallelism issues arise in the analysis of EM-ALLTOALLV as those described in §6.5, but with respect to message delivery rather than swapping. Since message delivery, like swapping, happens in the same order as virtual processor execution, the same arguments used for swapping (Def. 6.5.1) apply to message delivery as well. Def. 7.1.1 summarises the necessary conditions for communication methods that, like EM-ALLTOALLV, perform message I/O to/from all virtual processors.

**Definition 7.1.1** (Fully Parallel Message Delivery). If each virtual processor context resides on a single disk, \( k \geq D \), and virtual processors are scheduled in increasing order by ID, then a communication function which performs message I/O to/from all virtual processors does so across all \( D \) disks in parallel.

If each virtual processor context is distributed evenly across all disks in a blockwise fashion, and \( \omega \geq BD \), then a communication function which performs message I/O to/from all virtual processors does so across all \( D \) disks in parallel.

**Definition 7.1.2** (Fully Parallel I/O). A communication function has "fully parallel I/O" if it has both fully parallel swapping (Def. 6.5.1) and fully parallel message delivery (Def. 7.1.1).

**Lemma 7.1.3.** When used with explicit I/O, EM-ALLTOALLV-SEQ performs \( v\mu + \frac{v^2-v_k}{2}\omega + 2v^2B \) I/O.

*Proof.* The fundamental difference between Alg. 7.1.1 and Alg. 2.2.1 is that the amount of I/O performed by a given virtual processor depends on how many virtual processors have finished executing previously.

Let \( \delta \) be the number of messages delivered directly on line 11.

Let \( t \) be the number of messages delivered indirectly on line 14.

In lines 8...10, threads deliver directly to all threads that have completed Internal Superstep 1. Since threads execute in synchronised rounds \( k \) at a time, the first round of \( k \) threads *each* deliver \( k \) messages directly, the next round \( 2k \) (since \( 2k \) threads have
now run), the next round \(3k\), etc. Hence:

\[
\delta = \sum_{i=1}^{\frac{k}{2}} ik^2
\]

\[
= k^2 \left[ \frac{v}{k} \left( \frac{\frac{v}{k} + 1}{2} \right) \right]
\]

\[
= vk \left( \frac{\frac{v}{k} + 1}{2} \right)
\]

\[
= \frac{v^2 + vk}{2}
\]

\[
\omega = v^2 - \delta
\]

The remaining analysis is straightforward:

\[
I_{\text{seq}} = I_4 + I_{11} + I_{13..14} + I_{15}
\]

\[
= (v \mu - v^2 \omega) + (\delta \omega) + (2 \omega \omega) + (2v^2 B)
\]

\[
= v \mu - v^2 \omega + \delta \omega + 2v^2 \omega - 2\delta \omega + 2v^2 B
\]

\[
= v \mu + v^2 \omega - \left( \frac{v^2 + vk}{2} \right) \omega + 2v^2 B
\]

\[
= v \mu + \frac{v^2 - vk}{2} \omega + 2v^2 B
\]

\[
\Box
\]

**Corollary 7.1.4** (Improvement). When used with explicit I/O, EM-ALLTOALLV-SEQ performs \(2v\mu + \frac{3v^2 + vk}{2} \omega - 2v^2 B\) less message delivery I/O per virtual superstep than Alg. 2.2.1 (PEMS1-ALLTOALLV-SEQ).

**Proof.**

\[
\Delta I = I_{\text{orig-seq}} - I_{\text{seq}}
\]

\[
= (3v \mu + 2v^2 \omega) - \left( v \mu + \frac{v^2 - vk}{2} \omega + 2v^2 B \right)
\]

\[
= 2v \mu + \frac{3v^2 + vk}{2} \omega - 2v^2 B
\]
Lemma 7.1.5. EM-ALLTOALLV-SEQ uses at most $\frac{2u^2B}{P}$ shared buffer space.

Proof. The only buffer space used is for the block cache, when direct I/O is in use. Each of the $\frac{v}{P}$ local virtual processors has 2 blocks in the cache for each of its $v$ received messages. □

Theorem 7.1.6. Given fully parallel I/O (Def. 7.1.2), EM-ALLTOALLV-SEQ takes $S_{BD}^{\frac{v}{P}} + G_{2BD}^{\frac{v^2-vk}{D}} + G_{BD}^{2u^2} + L$ time.

Proof. Follows directly from Lem. 7.1.3, since EM-ALLTOALLV-SEQ performs no network communication and no significant computation. □

Benchmarks

Fig. 7.2 shows the run time of a single call to EM-ALLTOALLV-SEQ for various numbers of 32-bit integers. The x-axis represents total problem size as a number of 32-bit integers, and the y-axis represents total run time. Times are shown for both memory-mapped ("mmap") and explicit ("unix") I/O, for $k = 1$ and $k = 4$ cores (e.g. alltoall-mmap-k1 represents memory-mapped I/O with 1 core).

No action is performed by the program other than a single ALLTOALLV on the complete data set. Note in particular the performance improvement seen with UNIX I/O when using 4 cores compared to using a single core. Since the test program performs no significant computation, this shows that the run time of EM-ALLTOALLV-SEQ itself improves when $k$ increases, as Thm. 7.1.6 predicts (since the $vk$ term in the message delivery time is subtracted).

The situation is reversed with memory mapped I/O, due to the overhead of the operating system’s cache mechanism. This is not surprising: since communication primitives in PEMS2 are carefully tuned to minimise I/O, explicit I/O will always result in better performance for a trivial program that simply calls a collective communication function once. The potential benefit of memory-mapped I/O is that the operating system’s cache mechanism can avoid a large amount of I/O for certain programs, but that is not the case here.

Note that the experiment shown in Fig. 7.2 is a trivial program that does not represent a case where multi-core or memory-mapping are expected to show much
benefit. Fig. 7.2 is not intended to illustrate the improved performance of PEMS2, only that an improvement is seen when using multiple cores in spite of the fact that no computation is performed. Experiments to illustrate the improvements in PEMS2 for realistic use cases are shown in Chapter 8.

Figure 7.2: Single Processor EM-ALLTOALLv Performance
7.1.2 Multiple Processor Algorithm

Alg. 7.1.1 describes the multiple processor implementation of Alltoallv in PEMS2. Local message delivery occurs in an identical manner as in the single processor case. Remote messages are handled in the second internal superstep, when all message destinations are known (since they have been recorded in the shared table in the previous internal superstep). Using this information, virtual processors receive on behalf of their local peers and deliver directly to their contexts on disk.

Analysis

Lemma 7.1.7. EM-ALLTOALLV-PAR takes \( g \frac{akw}{b} + l \frac{v^2}{Pka} \) communication time, assuming \( akw \geq b \), with \( g \), \( b \), and \( l \) as in the BSP model (see Appendix B).

Proof. Each virtual processor must send one message to each other virtual processor, and all messages are sent directly to the real processor that hosts the destination virtual processor.

All communication performed by EM-ALLTOALLV-PAR is performed by EM-ALLTOALLV-PAR-COMM. This algorithm sends messages from the “round” of \( Pk \) currently executing virtual processors in “chunks” of size \( \alpha \) (where \( \alpha \) is a user-defined parameter indicating the number of messages to send at once, \( 1 \leq \alpha < v \)).

In each round, \( k \) virtual processors are active on each real processor, and each of these virtual processors sends \( v \) messages over the network\(^1\). Thus, each round consists of \( \frac{v}{P\alpha} \) separate \( akw \)-relations.

\( \frac{v}{P\alpha} \) such rounds occur, therefore there are \( \frac{v^2}{P^2\alpha} \) such relations in total. \( \square \)

Lemma 7.1.8. When used with explicit I/O, EM-ALLTOALLV-PAR performs \( \frac{u}{P} + \left( \frac{v^2}{P} - v^2 + \frac{3v^2}{2P^2} - \frac{kP}{2P} \right) \omega + 2v^2B \) I/O.

Proof. The local message delivery of the parallel version of EM-ALLTOALLV-SEQ is identical to that of EM-ALLTOALLV-SEQ, therefore this portion of the analysis \( (I_4, I_{11}, I_{13..14}, \text{and } I_{15}) \) is identical except with \( \frac{v}{P} \) local virtual processors rather than \( v \).

\(^1\)For simplicity of analysis, messages to local virtual processors are included in this figure though in reality this is optimised away and each virtual processor sends \( v - \frac{v}{P} \) messages over the network
The remaining I/O is \( I_{17..18} = \frac{v^2}{P} \omega \) for the received network messages.

\[
\delta = \sum_{i=1}^{Nk} i k^2 \\
= k^2 \left[ \frac{v}{P_k} \left( \frac{v}{P_k} + 1 \right) \right] \\
= \frac{vk}{P} \left( \frac{v}{2P_k} + 1 \right) \\
= \frac{v}{P} \left( \frac{v}{2P} + \frac{k}{2} \right) \\
= \frac{1}{2} \left( \frac{v}{P} \right)^2 + \frac{k}{2} \left( \frac{v}{P} \right)
\]

\[
\iota = \left( \frac{v}{P} \right)^2 - \delta
\]

\[
I_{seq} = I_4 + I_{11} + I_{13..14} + I_{17..18} + I_{15} \\
= \left( \frac{v \mu}{P} - v^2 \omega \right) + (\delta \omega) + (2 \omega) + \left( \frac{v^2}{P} \omega \right) + (2v^2 B) \\
= \frac{v \mu}{P} - v^2 \omega - \delta \omega + 2 \left( \frac{v}{P} \right)^2 \omega + \frac{v^2}{P} \omega + 2v^2 B \\
= \frac{v \mu}{P} - v^2 \omega - \frac{1}{2} \left( \frac{v}{P} \right)^2 \omega - \frac{k}{2} \left( \frac{v}{P} \right) \omega + 2 \left( \frac{v}{P} \right)^2 \omega + \frac{v^2}{P} \omega + 2v^2 B \\
= \frac{v \mu}{P} + \left( \frac{v^2}{P} + \frac{3v^2}{2P^2} - \frac{kv}{2P} - v^2 \right) \omega + 2v^2 B
\]

\[\Box\]

**Lemma 7.1.9.** EM-ALLTOALLV-PAR uses at most \( \frac{2v^2B}{P} + \alpha k \omega \) shared buffer space.

**Proof.** The size of the block cache is equivalent to the sequential case (Lem. 7.1.5). Additional space is used by EM-ALLTOALLV-PAR-COMM to assemble messages contiguously for communication. The "chunk size", \( \alpha \), is a user parameter which controls the amount of buffer space used for this purpose: at most \( \alpha k \omega \) buffer space is used. \[\Box\]

**Theorem 7.1.10.** Given fully parallel I/O (Def. 7.1.2), EM-ALLTOALLV-PAR takes \( S_{PDB} \frac{v \mu}{P} + G \left( \frac{v^2}{P} + \frac{3v^2}{2P^2} - \frac{kv}{2P} - v^2 \right) \frac{\omega}{PDB} + G2v^2 B + g \frac{\alpha k \omega}{b} + l \frac{v^2}{PDB} + L \) time.

**Proof.** Follows directly from Lem. 7.1.8 and Lem. 7.1.3. \[\Box\]
Algorithm 7.1.2: EM-ALLTOALLV-PAR

**Data:** $S$: Array of pointers to $v$ messages to send
**Data:** $R$: Array of pointers to $v$ messages to receive

1. Let $T$ be a shared $v \times \frac{p}{v}$ table of incoming message offsets
2. Let $E$ be a shared array of $\frac{p}{v}$ execution states, all initially false
3. Let $M$ be a cache of at most $\frac{2v^2}{p}$ border blocks (2v per local thread)
   - Store message offsets and synchronise —
4. Swap out everything except regions in $R$
5. Store incoming message offsets from $R$ in $T$ — $T_{\rightarrow p}$ is now valid
6. Set $E_p$ to true — This thread has reached this point
7. Synchronise with the $k - 1$ other currently executing local threads
   - Deliver messages if possible —
8. **foreach** local message $m_{p\rightarrow i}$ in $S$ **do**
   9. Update $M$ with the start and end of this message
   10. **if** $E_i$ is true **then** — Thread $i$ has recorded its offsets in $T$
   11. Align and deliver directly to $T_{p\rightarrow i}$ on disk

   — Finished Internal Superstep 1 —
   — Begin Internal Superstep 2 —
   — Deliver remaining messages —
12. **foreach** local message $m_{p\rightarrow i}$ in $S$ not delivered in superstep 1 **do**
13. Swap message in
14. Align and deliver directly to $T_{p\rightarrow i}$ on disk
15. Communicate using Alg. 7.1.3
16. **foreach** remote message $m_{i\rightarrow j}$ received **do**
17. Update $M$ with the start and end of this message
18. Align and deliver directly to $T_{i\rightarrow j}$ on disk

   — Finished Internal Superstep 2 —
   — Begin Internal Superstep 3 —
   — (Blocked I/O only) Flush border blocks —
19. Flush border blocks in $M$ to our context
   — Finished Virtual Superstep —
Algorithm 7.1.3: EM-ALLTOALLV-PAR-COMM

Data: $S$ : Array of pointers to $v$ messages to send
Data: $R$ : Array of pointers to $v$ messages to receive
Data: $T$ : Shared $v \times \frac{p}{p}$ table of message offsets

1. Let $\alpha$ be the network "chunk size" parameter
2. Let $B$ be the shared communication buffer
3. foreach $i$ in $0, \alpha, 2\alpha, 3\alpha, \ldots, \frac{p}{p} - 1$ do
4.   Assemble messages to threads $i \ldots i + \alpha - 1$ on each real processor contiguously in $B$
5.   if this is the last of $k$ threads to reach this point then
6.     Send/Receive assembled messages with MPI_Alltoallv
7.     Deliver received messages using $T$
7.2 Bcast

In a BCAST (broadcast), a single root virtual processor sends a single message to every other virtual processor, i.e. every virtual processor receives the same individual message.

![Figure 7.3: Bcast Operation](image)

The PEMS2 implementation of Bcast uses rooted synchronisation on the real processor that contains the root, and initial synchronisation on all other real processors (see §4.3).

All threads on the same real processor as the root wait for the root to write the message to the shared buffer. They then copy the message from the shared buffer to their receive buffer.

The message is delivered remotely using a single MPI_Bcast: the root sends, and the first virtual processor to run on other real processors receives into the shared buffer. This receiving virtual processor then signals, and other virtual processors copy from the shared buffer to their receive buffers.
Algorithm 7.2.1: EM-Bcast

**Data:** $S$: Send buffer of size $\omega$ (valid only at root)

**Data:** $R$: Receive buffer of size $\omega$

1. Let $B$ be the initial portion of shared buffer of size $\omega$

   — Broadcast —

2. if this is the root then
   3. Copy $S$ to $B$
   4. EM-SIGNAL-THREADS() — Signal that data is ready
   5. if $P > 1$ then
   6. MPLBcast from $S$ — Send to other real processors

7. else — This is not the root

8. if the root is on this real processor then
   9. EM-WAIT-FOR-ROOT()
   10. Copy from shared buffer to $R$

11. else root is on another real processor

12. if $P > 1$ and EM-FIRST-THREAD() then
   13. MPLBcast to $B$ (receive from root)
   14. EM-SIGNAL-THREADS()
   15. EM-WAIT-THREADS()
   16. Copy from $B$ to $R$

18

— Finished Virtual Superstep —
7.2.2 Analysis

Lemma 7.2.1. EM-BCAST takes \( S \frac{2\nu}{PKB} + G \frac{\nu \omega}{PDB} \) time to perform I/O (not including virtual superstep overhead).

Proof. EM-WAIT-FOR-ROOT takes \( S \frac{2\nu}{PKB} \) time (Lem. 4.3.1), and each virtual processor delivers the buffer (of size \( \omega \)) to its context. \( \square \)

Lemma 7.2.2. EM-BCAST performs a single network \( \omega \)-relation, where \( \omega \) is the size of the buffer to broadcast.

Proof. MPI_Bcast is called exactly once with a buffer of size \( \omega \). \( \square \)

Theorem 7.2.3. EM-BCAST takes \( S \frac{2\nu}{PKB} + G \frac{\nu \omega}{PDB} + g \frac{\omega}{b} + I + L \) time where \( \omega \) is the size of the buffer to broadcast, assuming \( v \omega > B \) and \( \omega > b \).

Proof. Follows directly from Lemma 7.2.2 and Lemma 7.2.1, since extra swapping only occurs for a single virtual processor and message delivery occurs for all virtual processors in parallel. \( \square \)
7.3 Gather

In a GATHER, each virtual processor sends a message to the root virtual processor.

![Figure 7.4: Gather Operation](image)

The PEMS2 implementation of Gather uses final synchronisation (see §4.3). In both the single and multiple processor cases, the gathered messages are assembled in the shared buffer before finally being collected by the root virtual processor.

In the single processor case, the virtual processors simply copy to the appropriate location in the shared buffer, then signal. When all threads have signalled, the root copies the result from the shared buffer to its receive buffer and the operation is complete.

In the multiple processor case, each virtual processor participates in an MPI_GATHER to send data to the real processor which hosts the root. When these communication rounds are completed, all gathered data resides in the shared buffer at the real processor which hosts the root. The root virtual processor then copies this data to its receive buffer and the operation is complete.

7.3.1 Algorithm
Algorithm 7.3.1: EM-GATHER

Data: \( S \): Send buffer of size \( \omega \)
Data: \( R \): Receive buffer of size \( v\omega \) (valid only at root)

1. Let \( B \) be the initial portion of shared buffer of size \( v\omega \)
2. Let \( y = \) false (yielded, swapped out)
3. if \( P > 1 \) then
   if the root is on this real processor then
      MPI.GATHER \( P \) ranks of current senders (receive)
      MPI.GATHER(\( S, B \)) (receive)
      if this is the root then
         if not EM-ALL-THREADS-FINISHED(\( y \)) then
            EM-WAIT-THREADS(\( y \))
         Copy data from \( B \) to \( R \)
      else
         EM-THREAD-FINISHED()
   else
      EM-THREAD-FINISHED()
   else — Root is not on this real processor
      MPI.GATHER \( P \) ranks of current senders (send)
      MPI.GATHER(\( S \)) (send)

else — Single processor
   if this is the root then
      if not EM-ALL-THREADS-FINISHED(\( y \)) then
         EM-WAIT-THREADS(\( y \))
      if \( y \) then
         Swap in \( R \)
      Copy \( S \) to \( B \)
      Copy \( B \) to \( R \)
   else — This is not the root
      Copy \( S \) to \( B \)
      EM-THREAD-FINISHED()

— Finished Virtual Superstep —
7.3.2 Analysis

Lemma 7.3.1. EM-GATHER takes at most $S_{BD}^{\mu+\omega}$ time to perform I/O (not including virtual superstep overhead), assuming $\omega > B$.

Proof. In the multi-processor case, the root may swap its context out via EM-WAIT-THREADS ($\mu$ I/O since only the root calls this function). In this case, $\mathcal{R}$ is not swapped in at line 10, so this copy is actually a disk write ($\omega$ I/O), for a total of $\mu + \omega$ I/O in the worst case. Other virtual processors perform no additional I/O. The single processor case clearly performs less I/O in the worst case. □

Lemma 7.3.2. EM-GATHER takes $g\frac{\omega}{P_0} + l\frac{\nu}{P}$ communication time, assuming $\omega > b$.

Proof. EM-GATHER performs an MPI_GATHER for $P$ threads at once, where each real processor sends one message of size $\omega$ to the root. Thus, EM-GATHER performs $\frac{\nu}{P}$ network $\omega$-relations. The lemma follows directly from the definitions of $l$, $g$, and $b$ in the BSP model. □

Theorem 7.3.3. EM-GATHER takes $S_{BD}^{\mu+\omega} + g\frac{\omega}{P_0} + l\frac{\nu}{P} + L$ time, assuming $\omega > b$ and $\omega > B$.

Proof. Follows directly from Lem. 7.3.1 and Lem. 7.3.2. □
7.4 Reduce

A Reduce operation applies an associative and commutative\(^2\) operator to \(v\) values (one from each virtual processor), placing the single result in a buffer on some root virtual processor. This operation is vectorized across \(n\) values, i.e. a single Reduce performs \(n\) reductions of size \(v\) resulting in \(n\) values at the root. (This definition corresponds to MPI\_Reduce, which is not implemented by PEMSl).

A Reduce can be performed with less I/O and communication than an Alltoall, since several values may be reduced to a single value before delivery. On each real processor, for each of the \(n\) reductions, EM-REDUCE performs \(k\) operations at a time in parallel into the shared buffer. \(\sigma\) must therefore be large enough to hold \(kn\) values. When all local threads have finished, the final thread reduces these \(k\) values to a single value, and communicates that value to the root. This requires 2 internal supersteps, but only a single swap and a single network communication (if necessary).

\(^2\)MPI allows the user to define non-commutative operators, but PEMSl currently requires operators to be commutative
7.4.1 Algorithm

**Algorithm 7.4.1: EM-REDUCE**

**Data:** $S$: Array of $n$ values to send  
**Data:** $R$ (root only): Array of $n$ values for result  
**Data:** $r$ (root): ID of root virtual processor

1. Let $B$ be one of $k$ shared buffer portions of size $n$
   
   — Partially reduce local data —

2. Swap out

3. Reduce $S$ into $B$
   
   — Finished Internal Superstep 1 —

4. If $p = r$ or $p$ is thread 0 on a different real processor than $r$ then

5. Reduce $kn$ values in shared buffer to $n$ local results

6. If $P > 1$ then

7. $\text{MPI-Reduce } n$ local results to $r$'s real processor

8. If $p = r$ then

9. If $P > 1$ then

10. $\text{MPI-Reduce } Pn$ values from the network into $R$

11. Swap $R$ out to partition on disk
   
   — Finished Virtual Superstep —

7.4.2 Analysis

**Lemma 7.4.1.** EM-REDUCE takes $\frac{nL}{P} + nk$ computation time to reduce all real processor’s local values to a local result.

*Proof.* All operations are performed on vectors of size $n$. For each of these $n$ elements: Each real processor first reduces $\frac{n}{P}$ values on $k$ cores in parallel, resulting in $k$ intermediate values (Step 1 in Fig. 7.5), which takes $\frac{n}{Pk}$ time. Then, these $k$ intermediate local results are combined by application of the reduction operator (Step 2 in Fig. 7.5), which takes $k$ time. Thus, it takes $\frac{n}{Pk} + k$ time to reduce a vector of size 1, or $\frac{nL}{P} + nk$ time to reduce a vector of size $n$. □

**Lemma 7.4.2.** EM-REDUCE takes $G\frac{nw}{B}$ time to perform I/O (not including virtual superstep overhead).
Proof. The root processor delivers the final result of size $n\omega$ to its context on disk. Precisely one swap occurs per virtual processor, which is accounted for by by $L$ if necessary.

When $P > 1$, EM-REDUCE performs a single network MPI.Reduce operation. The precise communication and computation time may vary between MPI implementations, but we can find reasonable bounds by assuming the MPI implementation is at least as good as the “obvious” algorithm, as described in Lem. 7.4.3.

**Lemma 7.4.3.** A reasonable MPI.Reduce implementation on a switched network reduces $nP$ values across $P$ processors (Step 3 in Fig. 7.5) in $n\lg(P) + g \frac{n\omega \lg(P)}{b} + l \lg(P)$ time, assuming $n\omega \lg(P) \geq b$.

Proof. MPI.Reduce can be implemented as a parallel tree reduction to achieve logarithmic time, as shown in Fig. 7.6. The result is computed as $\lg(P)$ parallel partial reductions. Each partial reduction combines two vectors of length $n$ ($n\lg(P)$ total time), and is the result of sending a single vector of $n$ values each of size $\omega$ over the network ($g \lg(P) \frac{n\omega}{b}$ total time).

![Logarithmic MPI.Reduce](image)

Figure 7.6: Logarithmic MPI.Reduce

**Theorem 7.4.4.** EM-REDUCE takes $G \frac{n\omega}{B} + g \frac{n\omega \lg(P)}{b} + l \lg(P) + n \lg(P) + \frac{n\omega}{P_k} + nk + L$ time.

Proof. Follows directly from Lem. 7.4.1, Lem. 7.4.2, and Lem. 7.4.3. 

\[ \square \]
### 7.5 Summary

<table>
<thead>
<tr>
<th>Operation</th>
<th>Buffer Space</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bcast</td>
<td>$\omega$</td>
</tr>
<tr>
<td>Gather</td>
<td>$v\omega$</td>
</tr>
<tr>
<td>Reduce</td>
<td>$kn$</td>
</tr>
<tr>
<td>Alltoallv-Seq</td>
<td>$\frac{2v^2B}{P}$</td>
</tr>
<tr>
<td>Alltoallv-Par</td>
<td>$\frac{2v^2B}{P} + \alpha kW$</td>
</tr>
</tbody>
</table>

Figure 7.7: Communication Algorithm Buffer Space

<table>
<thead>
<tr>
<th>Operation</th>
<th>Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bcast</td>
<td>$S \frac{2uv}{PkB} + G \frac{vw}{PDB} + g \omega + l + L$</td>
</tr>
<tr>
<td>Gather</td>
<td>$S \frac{vw}{BD} + g \frac{vw}{Pb} + l \frac{P}{b} + L$</td>
</tr>
<tr>
<td>Reduce</td>
<td>$G \frac{vw}{B} + g \frac{nw \log(P)}{b} + n \log(P) + \frac{nw}{pk} + nk + L$</td>
</tr>
<tr>
<td>Alltoallv-Seq</td>
<td>$G \frac{v^2}{2BD} \omega + G \frac{2v^2}{D} + L$</td>
</tr>
<tr>
<td>Alltoallv-Par</td>
<td>$S \frac{vw}{PDB} + G \left( \frac{v^2}{P} + \frac{3v^2}{2P} - \frac{kv}{2P} - \omega\right) \frac{v}{PDB} + G2v^2B + g \frac{akw}{b} + l \frac{v^2}{Pka} + L$</td>
</tr>
</tbody>
</table>

Figure 7.8: Communication Algorithm Run Time
8.1 Experimental Setup

All experiments in this chapter were performed on the HPCVL cluster described in detail in Appendix C.1.

8.2 Plot Style

Labels for plot lines show the program name, PEMS version, I/O style, and number of processors. For example, “PSRS PEMS2 (mmap) P=2” refers to the PSRS algorithm running on PEMS2 with mmap I/O on 2 real processors. The three types of I/O style referred to are shown in Fig. 8.2.

<table>
<thead>
<tr>
<th>Label</th>
<th>I/O Style</th>
</tr>
</thead>
<tbody>
<tr>
<td>unix</td>
<td>Synchronous UNIX I/O</td>
</tr>
<tr>
<td>stxxl-file</td>
<td>Asynchronous STXXL File I/O (§5.1)</td>
</tr>
<tr>
<td>mmap</td>
<td>Memory Mapped I/O (§5.2)</td>
</tr>
</tbody>
</table>

Figure 8.1: PEMS2 I/O Styles

The label “stxxl” refers to STXXL’s included sorting algorithm, run on a single processor (the program does not support distributed processors). This data is included on all plots to provide a consistent baseline for comparison of other results.

Variables shown below the plot (e.g. $\mu$ or $\frac{\pi}{\nu}$) apply to all runs shown in that plot, with the exception of the “stxxl” data.
8.3 Sorting

The well-known Parallel Sorting by Regular Sampling [17] (PSRS) algorithm is a good candidate for use with PEMS and explicit I/O due to its coarse granularity and small, constant number of supersteps. Alg. 8.3.1 shows a simple high-level description of this algorithm, with calls to collective communication functions (i.e. calls that result in I/O via PEMS) shown in bold.

8.3.1 Algorithm

Algorithm 8.3.1: PSRS

<table>
<thead>
<tr>
<th>Data: $\mathcal{D}$ (data) : Array of size $\frac{n}{v}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 Sort $\mathcal{D}$</td>
</tr>
<tr>
<td>2 Choose $v$ equally spaced splitters in $\mathcal{D}$</td>
</tr>
<tr>
<td>3 Gather all $v^2$ splitters at root</td>
</tr>
<tr>
<td>4 Sort all $v^2$ splitters at the root</td>
</tr>
<tr>
<td>5 $\text{Bcast}$ splitters evenly to all processors (each receives $v$ splitters)</td>
</tr>
<tr>
<td>6 Locate splitters in (sorted) $\mathcal{D}$</td>
</tr>
<tr>
<td>7 Compute the number of elements in $\mathcal{D}$ in each bucket</td>
</tr>
<tr>
<td>8 $\text{Alltoall}$ bucket sizes (each sends/receives $v$ sizes)</td>
</tr>
<tr>
<td>9 $\text{Alltoallv}$ buckets to final destination processor</td>
</tr>
<tr>
<td>10 Merge received buckets</td>
</tr>
</tbody>
</table>

8.3.2 Analysis

Let $\pi$ be the size of an integer used for counts.

Let $\epsilon$ be the size of an individual data element.

Alg. 8.3.1 consists of four supersteps, the first three of which communicate only counts of a fixed size.

The remaining call, $\text{Alltoallv}$, does all the work of distributing data among processors. The message sizes in this step therefore depends on how balanced the global partitioning is. The partitioning scheme used in PSRS guarantees the balancing is within a factor of 2 of an ideal partitioning [17] We can assume, then, that the worst case virtual message size for this final step is $\omega \leq \frac{2n\epsilon}{v^2}$.
<table>
<thead>
<tr>
<th>Operation</th>
<th>Size</th>
<th>Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gather</td>
<td>$v\pi$</td>
<td>$S^{\mu+2v\pi}<em>{BD} + g^{kv</em>\pi}_b + l^{v}_P + L$</td>
</tr>
<tr>
<td>Bcast</td>
<td>$v\pi$</td>
<td>$S^{2v\mu}<em>{PDB} + G^{v^2\pi}</em>{PDB} + g^{v\pi}_b + l + L$</td>
</tr>
<tr>
<td>Alltoall</td>
<td>$v\pi$</td>
<td>$O\left(S^{\mu}<em>{PDB} + G^{v^2\pi}</em>{PDB} + g^{akv\pi}<em>b + l^{v^2}</em>{Pka} + L\right)$</td>
</tr>
<tr>
<td>Alltoallv</td>
<td>$2\nu e/v^2$</td>
<td>$O\left(S^{\nu e}<em>{PDB} + G^{2new}</em>{PDB} + g^{ak2me}<em>{v^2b} + l^{v^2}</em>{Pka} + L\right)$</td>
</tr>
</tbody>
</table>

### 8.3.3 Performance

**PEMS1 vs. PEMS2 Time (Scaling $v$)**

To allow direct comparison with previous experimental results for PEMS1 [15], these experiments use a small virtual processor context size, $\mu = 64$ MiB, with an additional 64 MiB for the shared buffer. Because direct I/O is used the operating system does not use extra RAM for caching, i.e. performance is as if the system had only this amount of RAM and is unaffected by additional memory which goes unused\(^1\).

The context size, $\mu$, remains constant for all runs, while the problem size is increased via $v$. This is the ideal way to scale PEMS: choose the memory parameters suitably for the available hardware, then scale $v$ up to reach the desired problem size.

The PEMS1 and PEMS2 programs are identical, and experiments were run on the same machines with identical configuration and PEMS parameters.

As the figures in this section show, PEMS2 is significantly faster than PEMS1, particularly with several real processors. When run with 8 processors, PEMS1 is still not competitive with STXXL, taking over twice as long. PEMS2, however, is faster than STXXL with 8 processors, and very close in speed with 4. PEMS2 also scales better than PEMS1, with a slope nearly identical to that of STXXL. In contrast, the performance gap between PEMS1 and STXXL gets larger as the problem size increases.

---

\(^1\)This has been verified by monitoring resource consumption and running identical experiments on machines with reduced RAM
Figure 8.2: PEMS1 vs. PEMS2 (PSRS, P=1)

\[ n = 8000000, P = 1, \mu = 64 \text{ MiB} \]

Figure 8.3: PEMS1 vs. PEMS2 (PSRS, P=2)

\[ \frac{n}{v} = 8000000, P = 2, \mu = 64 \text{ MiB} \]
\[ \frac{n}{v} = 8000000, \ P = 4, \ \mu = 64 \text{ MiB} \]

Figure 8.4: PEMS1 vs. PEMS2 (PSRS, P=4)

\[ \frac{n}{v} = 8000000, \ P = 8, \ \mu = 64 \text{ MiB} \]

Figure 8.5: PEMS1 vs. PEMS2 (PSRS, P=8)
PEMS1 vs. PEMS2 Speedup

Fig. 8.6 shows the relative speedup of the experiments in the previous section where \( n = 4 \) billion. The speedup shown is relative to the sequential execution of the same system, i.e. PEMS1 speedup is relative to PEMS1 with \( P = 1 \) and PEMS2 speedup is relative to PEMS2 with \( P = 1 \).

This figure illustrates that PEMS2 performance improves as real processors are added significantly more than that of PEMS1.

![Graph showing speedup comparison between PEMS1 and PEMS2](image)

**n = 4096000000, v = 512, \( \frac{n}{v} = 8000000 \), \( \mu = 64 \) MiB**

Figure 8.6: PEMS1 vs. PEMS2 PSRS Relative Speedup

PEMS1 vs. PEMS2 Time (Scaling \( \mu \))

The results in §8.3.3 confirm the hypothesis that always delivering directly to virtual processor contexts on disk results in improved performance compared to delivering indirectly via a separate area on disk. However, the parameters used here are not realistic – modern machines have far more RAM than 128 MiB. While these smaller runs can be extrapolated to estimate the performance of runs using more RAM (as
mentioned in previous work [15][16], when comparing PEMS1 and PEMS2, the context size has an impact on performance due to the differing disk layouts and delivery strategies.

As described in §2.3.2, a significant motivation factor for the new direct delivery strategy was the reduction of disk seeking. Because virtual processor contexts and the message area reside on separate areas of disk in PEMS1, during a simulation the disk must seek back and forth between these (possibly very distant) areas in order to perform swapping and delivery. This effect should increase as $\mu$ increases, since increasing $\mu$ increases the distance from a context to the indirect area, as well as the distance between each region of the indirect area itself.

Fig. 8.7 shows the results of an experiment with the PSRS algorithm that confirms this observation. In this experiment, the context size ($\mu$) increases, while $v$ remains constant. Thus there is an equal number of virtual processors for every run, but each virtual processor handles a larger number of elements. The results clearly show that PEMS2 scales significantly better than PEMS1 with respect to increasing $\mu$. This is an important observation, since experiments using small contexts do not illustrate this aspect of performance. Because modern machines have several GiB of memory, the experiments in the previous section (and similar experiments in the PEMS1 literature) do not realistically reflect the performance of PEMS when used for practical purposes, where it is desirable to use as much RAM as possible to maximise performance.
Figure 8.7: Increasing Context Size with Constant $v$
PEMS2 Large Runs

The dramatically different slopes in Fig. 8.7 suggest that PEMS2 should be more suited to exploiting the full capabilities of modern machines. In order to investigate this, the experiments in this section use a more realistic context size, $\mu = 1$ GiB. Each machine has 4 cores ($k = 4$), thus 4 GiB RAM per machine is used for virtual processor contexts. An additional GiB is used for the shared buffer, for a total of 5 GiB$^2$

These runs illustrate performance with much larger problem sizes: up to roughly 32 billion ($32 \cdot 10^9$) 32-bit integers; or about 119 GiB of data. Because the PSRS algorithm requires twice the space in order to sort, as well as additional space for counts, this amounts to well over 200 GiB of data; significantly larger than the total amount of physical memory available on the machines used.

The improved performance of PEMS2 with larger context sizes can be seen by comparing performance to STXXL with the small context results in §8.3.3 (the STXXL data is identical). In those experiments with small contexts, PEMS2 only surpasses STXXL performance at $P = 8$ (Fig. 8.5). In the experiments here with larger contexts, PEMS2 surpasses STXXL performance at $P = 4$ (Fig. 8.10), and is very close when $P = 2$ (Fig. 8.9), a significant improvement.

Direct comparison of individual runs with similar problem sizes illustrates the improvement clearly. For example, with $P = 8$, PEMS2 with small contexts takes 1441 seconds to sort 4 billion elements (Fig. 8.5). PEMS2 with large contexts takes only 704 seconds to sort 4 billion elements (Fig. 8.11), more than twice as fast as the comparable run with small contexts. PEMS1 with small contexts takes 3925 seconds to sort 4 billion elements (Fig. 8.5), making PEMS2 with large contexts more than 5 times as fast. Considering the fact that PEMS2 scales better than PEMS1 with respect to context size (Fig. 8.7), it is clear that PEMS2 is a significant improvement over PEMS1 for practical applications which utilize the resources available on modern hardware.

Performance is best, and most predictable, when using UNIX I/O. Memory-mapped I/O performs significantly worse, though this is not surprising since PSRS is

---

$^2$Of course, a small amount of additional memory is used for internal control structures, thread stacks, the operating system, etc.
not a good choice of algorithm for memory mapping: most memory is used in all steps, so caching provides little benefit but a large amount of overhead. More surprisingly, asynchronous STXXL I/O does not outperform the synchronous UNIX I/O with the exception of a few runs. If the \( n = 32 \) billion data point is ignored, the asynchronous performance for \( P = 8 \) looks promising; it may be the case that further optimisation of the implementation to increase I/O, computation, and communication overlap will allow asynchronous I/O to show a consistent improvement over synchronous I/O.

\[ \frac{n}{v} = 128000000 \, , \, P = 1 \, , \, k = 4 \, , \, \mu = 1024 \text{ MiB} \]

Figure 8.8: PSRS PEMS2 P=1
Figure 8.9: PSRS PEMS2 P=2

\[ \frac{n}{\nu} = 128000000, \ P = 2, \ k = 4, \ \mu = 1024 \ \text{MiB} \]

Figure 8.10: PSRS PEMS2 P=4

\[ \frac{n}{\nu} = 128000000, \ P = 4, \ k = 4, \ \mu = 1024 \ \text{MiB} \]
Figure 8.11: PSRS PEMS2 P=8

\[ \frac{n}{\nu} = 128000000, \ P = 8, \ \mu = 1024 \text{ MiB} \]
**Internal Benchmarks**

PEMS includes a more fine-grained benchmarking system which plots the time of a program's execution at each superstep barrier. This can be used to precisely investigate the run time of a program and see which sections of code spend the most time. This data is shown in Fig. 8.12, Fig. 8.14, and Fig. 8.13 for a PSRS run using unix, mmap, and stxxl-file I/O, respectively. These plots clearly illustrate where time is consumed among the 4 communication calls in the PSRS algorithm.

Each plot shows a single PSRS execution on a single real processor (runs are using 2 real processors and the same parameters as in the previous section, but each plot shows only a single real processor). Each line represents the elapsed time of a single thread. Because $\frac{n}{p}$ is relatively high for these runs, there are many lines on these plots and distinguishing each individually is difficult. However, it is the overall trend and distribution that is interesting in these plots, not the time taken by any particular thread.

These plots illustrate the fundamental performance difference between explicit and memory mapped I/O. The UNIX and STXXL plots have similar structures - time increases in jumps at each superstep, roughly corresponding to the amount of I/O performed in that superstep. MMap, however, differs significantly: elapsed time is nearly flat until the final Alltoallv. This illustrates the effect of caching and benefits of memory mapping - the first 3 steps deal with only splitter data, thus a small amount of data is accessed each step. This allows the cache to work effectively, keeping the splitter data in memory and avoiding I/O. The advantage of memory-mapped I/O in PEMS is clearly visible: these steps take almost no time at all, because no swapping I/O is performed. The final Alltoallv, however, moves all the data to its final location, accessing the majority of the virtual processor's memory. This data is large and not cached, so a large amount of I/O is performed. This last step which accesses and moves the majority of memory on every virtual processor causes PSRS to not see much overall performance advantage with memory-mapped I/O. Other algorithms which communicate in smaller chunks would see a significant improvement in overall runtime, as the first 3 steps in Fig. 8.14 illustrate. Thus, memory mapping expands the scope of PEMS: with explicit I/O as in PEMS1, only algorithms like PSRS with a small number of very coarse supersteps are appropriate. PEMS2 with memory
mapped I/O, however, can see good performance with algorithms that use a large number of supersteps and finer grained communication, since a superstep barrier no longer forces a complete swap.

Figure 8.12: PSRS PEMS2 (unix) Elapsed Time Per Thread

(Each line represents elapsed time for a single virtual processor)
Figure 8.13: PSRS PEMS2 (stxxl-file) Elapsed Time Per Thread

(Each line represents elapsed time for a single virtual processor)

Figure 8.14: PSRS PEMS2 (mmap) Elapsed Time Per Thread

(Each line represents elapsed time for a single virtual processor)
8.4 CGMLib

CGMLib/CGMGraph [4] (here collectively referred to as “CGMLib”) is an implementation of several CGM algorithms and associated utilities. CGMLib is a high-level object based C++ library implemented on top of MPI, which implements several communication methods:

oneToAllBCast(int source, CommObjectList &data) Broadcast the list data from processor number source to all processors.

allToOneGather(int target, CommObjectList &data) Gather the lists data from all processors to processor number target.

hRelation(CommObjectList &data, int *ns) Perform an h-Relation on the lists data using the integer array ns to indicate for each processor which list objects are to be sent to which processor.

allToAllBCast(CommObjectList &data) Every processor broadcasts its list data to all other processors.

arrayBalancing(CommObjectList &data, int expectedN=-1) Shift the list elements between the lists data such that every processor contains the same number of elements.

partitionCGM(int groupId) Partition the CGM into groups indicated by groupId. All subsequent communication operations, such as the ones listed above, operate within the respective processor’s group only.

unPartitionCGM() Undo the previous partition operation.

These communication methods are implemented using MPI collective communication methods. All methods are supported by PEMS excluding partitionCGM and unPartitionCGM, which depend on the MPI_Comm_split call which is not currently implemented by PEMS.

CGMLib also provides additional utilities, such as communication and computation benchmarking, a system for routing data requests between processors, and commonly useful algorithms such as sorting, prefix sum, and list ranking.
8.4.1 Sort

The sorting algorithm included in CGMLib is a simple deterministic parallel sample sort, based on PSRS [17] and techniques described in [5]. The figures in this section show the performance of this sort under PEMS.

Though CGMLib Sort is similar to PSRS, performance under PEMS differs dramatically from the straightforward PSRS MPI implementation presented in §8.3. Unfortunately, characteristics of CGMLib and PEMS interact in ways that limit the problem size achievable for a given system. In particular, the CGMLib sort allocates much more memory. In the context for which CGMLib was originally designed (direct execution on a cluster using MPI) this does not significantly impact performance. However, when explicit I/O is used in PEMS, the amount of memory allocated has a very large impact on performance since this dictates the amount of swapping I/O performed. The problem is amplified by the fact that the CGMLib communication primitives typically use several MPI communication functions, which results in a larger number of swaps each superstep.

Though $n$ is much smaller than the PSRS results from §8.3, because of a larger constant factor of memory consumption, the runs in this section represent a large problem in terms of the amount of data handled by PEMS. The largest runs reach the limit of available disk space on our test configuration. Importantly, though $n$ itself is not very large from an EM perspective, the actual amount of allocated memory used is well in excess of the total amount of available system RAM.

8.4.2 Prefix Sum

The CGMLib Prefix Sum application finds the inclusive prefix sum of an array distributed across all processors.

The inclusive prefix sum of an array $[a_0, a_1, \ldots, a_{n-1}]$ is the array $[a_0, a_0+a_1, \ldots, a_0+a_1+\ldots+a_{n-1}]$, i.e. each element in the result is the sum of that element and all previous elements. For example, the prefix sum of $[1,2,3,4]$ is $[1,3,6,10]$.

This application shows similar performance to CGMLib Sort. This is expected, since both algorithm perform a small constant amount of communication.
\section*{Figure 8.15: CGMLib Sort PEMS2 P=1}

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure8_15}
\caption{CGMLib Sort PEMS2 P=1}
\end{figure}

\begin{align*}
\frac{n}{v} &= 5120000, \; P = 1, \; k = 4, \; \mu = 1400 \text{ MiB}
\end{align*}

\section*{Figure 8.16: CGMLib Sort PEMS2 P=2}

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure8_16}
\caption{CGMLib Sort PEMS2 P=2}
\end{figure}

\begin{align*}
\frac{n}{v} &= 5120000, \; P = 2, \; k = 4, \; \mu = 1400 \text{ MiB}
\end{align*}
Figure 8.17: CGMLib Sort PEMS2 P=4

Figure 8.18: CGMLib Prefix Sum PEMS2 P=1
Figure 8.19: CGMLib Prefix Sum PEMS2 P=2

Figure 8.20: CGMLib Prefix Sum PEMS2 P=4
8.4.3 Euler Tour

The CGMLib Euler Tour application[9] finds the Euler Tour of a forest.

The Euler Tour of a graph is a path which traverses every edge of the graph exactly once and returns to the starting point. In order to apply this problem to a tree, or a forest (a collection of trees), each edge is doubled. Figures 8.21, 8.22, and 8.23 show example input, transformed input, and output for this problem, respectively. The labels on nodes in 8.23 represent the order each node is visited in the Euler Tour.

![Figure 8.21: Euler Tour Input](image1)

![Figure 8.22: Euler Tour Input (Doubled Edges)](image2)

![Figure 8.23: Euler Tour Solution](image3)
This application is significantly more complex than CGMLib Sort and CGMLib Prefix Sum, and uses several other facilities of CGMLib (including sorting and list ranking). Fig. 8.24 shows the performance of the CGMLib Euler Tour application with memory mapped I/O. Here, \( n \) refers to the total number of trees in the forest, each of which contains \( n^2 \) nodes.

\[
\begin{align*}
\text{Seconds} & \quad 900 & 800 & 700 & 600 & 500 & 400 & 300 & 200 & 100 & 0 \\
0 & 5 \times 10^0 & 5 \times 10^1 & 5 \times 10^2 & 5 \times 10^3 & 5 \times 10^4 & 5 \times 10^5 & 5 \times 10^6 & 5 \times 10^7 & 5 \times 10^8 & 5 \times 10^9 \\
\end{align*}
\]

CGMLib Euler Tour (mmap) \( P=4 \)

\( v = 16 \), \( P = 4 \), \( \frac{v}{P} = 4 \), \( k = 4 \), \( io = \text{mmap} \)

Figure 8.24: CGMLib Euler Tour
8.4.4 CGMLib + PEMS Conclusions

Though the high constant factor of memory consumption prevents the CGMLib sort from being competitive with the simpler PSRS implementation, the results do show favourable scalability characteristics. It is likely that improvements to the PEMS memory allocator to reduce fragmentation, reductions in CGMLib’s memory usage, and other improvements would result in a significant reduction in constant overhead and make PEMS+CGMLib a competitive solution.

A positive outcome of these experiments can be seen in the results where \texttt{mmap} I/O is used. The problems described above are directly related to the use of explicit I/O – more allocated memory translates to more I/O. Memory mapped I/O, however, avoids this problem (as described in detail in §5.2). CGMLib, then, provides an ideal example of where the new memory mapped I/O capability of PEMS can be advantageous. As can be seen in the results, the CGMLib applications perform dramatically better with memory mapped I/O compared to Unix and STXXL I/O. This is because the large amount of allocated memory is not entirely used in each superstep, allowing the Operating System’s cache to do a good job of keeping required data in memory across supersteps, avoiding disk I/O. This confirms that memory mapping is an effective strategy in PEMS for improving the performance of algorithms with certain characteristics.
Chapter 9

Conclusions

This thesis presents PEMS2, an improved version of PEMS (Parallel External Memory System). PEMS can be used to execute BSP-like algorithms implemented as MPI programs with massive data sets larger than main memory by utilising disk.

PEMS2 incorporates most of the future work mentioned in the literature associated with PEMS1 [15]. In particular, PEMS2 adds multi-core support, asynchronous I/O, and reduces disk requirements. Beyond this, a new message delivery strategy has been introduced, and the implementation heavily reworked to a more flexible and easy to use form, with MPI compatibility and a run-time configuration system which allows simple experimentation with any algorithm.

Experiments show that PEMS2 performs significantly better than PEMS1, particularly when using the full resources of modern hardware.

9.1 Future Work

Since PEMS2 is simple to use with existing MPI code, much of the interesting work to be done based on this thesis is experimentation with various algorithms and configurations. It is hoped that PEMS2 will prove useful in practice to other researchers and practitioners interested in very large problems.

However, there are several potential avenues of investigation related to PEMS itself:

- Further MPI Compatibility: Unfortunately, many existing MPI programs are not actually BSP or BSP-like algorithms. It may be possible to implement non-collective communication functions (e.g. MPI_Send and MPI_Recv) in PEMS. However, because these functions do not adhere to the superstep model (i.e. they are not collective communication methods), implementation in PEMS may be difficult. If possible, though, this would increase the number of readily available PEMS compatible programs dramatically. While there may be negative
performance implications with algorithms that use these functions, the fact that a great deal of MPI programs use them is undeniable. Such algorithms, if well designed, could be practical EM solutions if PEMS had efficient support for non-collective communication.

- **Asynchronous and Memory-Mapped I/O**: Further investigation is required to evaluate the effectiveness of these modifications. The experiments in this thesis do not fully investigate the full potential of these new styles of I/O. While experiments with CGMLib have shown memory mapping to be a useful strategy, results for asynchronous I/O have (perhaps surprisingly) not shown much advantage.

- **Dynamic α**: The communication “chunk size” parameter used by Alltoallv, α, is currently specified as a user parameter. This could be made dynamic, so α is as large as possible for each communication (e.g. an Alltoallv with very small messages would only perform a single network communication).

- **Fully Asynchronous Design**: Lack of significant performance increases seen when using asynchronous STXXL I/O suggest that synchronisation in PEMS limits performance. A fully asynchronous design where both network communication and disk delivery are handled by a separate “controller” thread could alleviate this problem. In such a design, virtual processors would first record their message destinations much like the current Alltoallv design in PEMS2 (see §6.2). Rather than delivering in synchronised rounds, however, messages would be sent over the network immediately. The controller thread on the receiving real processor would receive the message and immediately write it to the correct location on disk. This way, delivery of messages does not require synchronisation between the sending and receiving virtual processors, thus communication, computation, and I/O overlap would be significantly increased. Such a design would also be more appropriate for implementing non-collective communication methods like MPI_Send and MPI_Recv.

- **Multi-Core and Multi-Disk**: Due to hardware limitations, the multiple-disk capabilities of PEMS2 have not been tested. It is likely that the use of several
cores and several disk will show significant performance improvements since several virtual processors running at once can make full use of up to $k$ disks (or more, depending on the type of I/O used).

- Disk Scheduling: Recent versions of Linux include several disk scheduling algorithms. It would be interesting to investigate the impact these have on the performance of PEMS.

- CGMLib: The scalability shown by CGMLib+PEMS2 experiments in §8.4 is promising, but absolute performance is hindered by excessive copying and memory consumption. Improving these characteristics of CGMLib would make the combination of CGMLib and PEMS2 a more practical solution, and due to the impressive breadth of functionality available in CGMLib, simplify the development of many advanced EM algorithms with PEMS2.

- New Architectures: There are interesting similarities between disk-based models such as those used in this thesis and increasingly popular special purpose multi-core (e.g. the Cell BE) and General Purpose Graphics Processing Unit (GPGPU) architectures. Both have a fast local memory store, and a slower external memory store. Transferring data between these two stores is a key factor in performance. PEMS, with some modifications, may allow suitable BSP algorithms to run on these new architectures with good performance. The implementation currently contains a “mem” I/O driver which simply uses allocated memory and does no I/O at all. This driver shows good performance and multi-core speedup (but of course can not scale beyond the limits of RAM). This illustrates that PEMS2 is not inherently tied to disk I/O, and adapting PEMS2’s strategy to novel architectures is an interesting avenue for future research.
Bibliography


Appendix A

Availability of PEMS2

PEMS is freely available online at http://pems.sourceforge.net/. PEMS is Free / Open Source software licensed under the GNU General Public License (GPL). Essentially this means PEMS is free to use, modify, and distribute, but all derivative works must also be released with source code under the GPL. The PEMS2 implementation is designed to be simple to use and extend (e.g. use with applications is trivial due to MPI compatibility, adding new I/O drivers is straightforward). Use, experimentation, and modification is encouraged; if licensing is an issue please contact the authors. Contact information is available at the PEMS website.

PEMS can be compiled and installed using the typical process for UNIX software: ./configure; make; sudo make install. Run ./configure --help for a summary of the available compile-time options which can be passed to ./configure.

Using PEMS with MPI programs is as simple as using any other MPI implementation. No source code modifications are necessary. There are two ways of doing so:

1. PEMS uses the pkg-config system for compiler and linker flags. The command pkg-config --cflags pems2 will return the compiler flags required for building against PEMS2, and pkg-config --libs pems2 will return the linker flags required for linking against PEMS2.

2. Like many MPI implementation, PEMS ships with compiler wrapper scripts to automatically add the necessary compiler and linker flags. Simply replacing uses of mpicc and mpic++ with pemsc and pemsc++, respectively, will build an MPI program against pems. Most MPI programs ship with a Makefile where this modification can be easily made.
Appendix B

Conventions

B.1 Terminology

real processor A single physical computer which runs a single process (of possibly many threads) and may have several cores which share main memory.

virtual processor A processor in the simulated bulk-synchronous algorithm.

thread The implementation of a virtual processor. While there is a 1 : 1 correspondence between threads and virtual processors, the two are not identical – a thread performs work internal to the PEMS implementation in addition to the work of the simulated virtual processor.

context The memory of a virtual processor, which may exist on disk or in main memory depending on whether or not the virtual processor is currently swapped in.

memory partition A context-sized block of real main memory into which a context is swapped. Unlike contexts, all memory partitions fit into main memory at once.

internal superstep A superstep performed internally by PEMS (e.g. as part of a multi-superstep communication method).

virtual superstep A superstep performed by the simulated algorithm (the simulation of a virtual superstep may require several internal supersteps).

swap The process of writing/reading an entire context to/from disk. The more specific terms swap in and swap out are used where necessary.
B.2 Notation

\( n \) = Size of the problem to be solved
\( t \) = a thread ID which is a local identifier for a thread (\( 0 \leq t < \frac{n}{P} \)).
\( \rho \) = the current virtual processor's global ID (\( 0 \leq \rho < v \)).
\( m_{i\rightarrow j} \) = a message sent from virtual processor \( i \) to virtual processor \( j \).
\( \lfloor x \rfloor \) = \( x \) rounded down to the nearest disk block boundary.
\( \lceil x \rceil \) = \( x \) rounded up to the nearest disk block boundary.
\( \lfloor r \rfloor \) = the smallest block aligned region containing range \( r \)
\( \lceil r \rceil \) = the largest aligned region within range \( r \)

B.3 Simulation Parameters

A PEMS simulation has the following run-time parameters:

\( P \) = Number of real processors
\( \mu \) = Memory size of a single virtual processor
\( D \) = Number of disks per real processor (\( D \geq 1 \))
\( v \) = Total number of virtual processors (\( v > P \))
\( k \) = Number of concurrent threads per real processor (\( k < \frac{v}{P} \))
\( \sigma \) = Size of the “shared buffer” in main memory

B.4 System Parameters

For performance analysis we make use of the following variables, which are essentially those of the BSP* model (in lowercase) with analogous variables (in uppercase) to represent EM performance:

\( b \) = Minimum size of a network message to achieve rated throughput (BSP*)
\( g \) = Time to deliver a network packet of size \( b \), or 0 if \( P = 1 \) (BSP*)
\( l \) = Overhead of a single network superstep (BSP*)
\( B \) = Size of a single disk block (EM)
\( G \) = Time to write/read a single block of size \( B \) to/from disk (EM)
\( L \) = Overhead of a single virtual superstep (EM)
\( S \) = Time to swap a single block of size \( B \) to/from disk (EM)
$L$ represents the constant overhead of a virtual superstep, including any synchronisation time and swapping I/O. $L$ may vary depending on the I/O system in use. Note that, due to the introduction of memory-mapped I/O, this is in contrast to previous work related to PEMS [13][7][6][15][16] where $L$ does not include any I/O.

$S$ is used to separate terms representing swap I/O from terms representing message delivery I/O. $S$ is identical to $G$ when using explicit I/O (i.e. UNIX or STXXL), and 0 by definition when using memory mapped I/O.

$g$ represents parallel network performance given a fully connected network, i.e. each processor can send messages directly to each other processor. When two processors communicate, network bandwidth between other processors is not affected. This is true for switched ethernet networks, but not true for ethernet hubs. Hubs are not suitable for high performance computing/networking, but this is not a concern with modern hardware since switches have reduced in price so much that even low-end consumer hardware is typically switched.
Appendix C

Methodology

C.1 Hardware/Software Configuration

Experiments were run on the HPCVL Beowulf cluster at Carleton University. Each node has 2 dual-core\(^1\) AMD Opteron 2214 processors at 2.2 GHz with 1 MiB cache per core, 8 GiB RAM, and a single 200 GiB disk. Nodes are interconnected with a high-end gigabit ethernet switch.

Linux 2.6.28 was used, with the ext4 filesystem and standard I/O scheduling. All code was compiled with GCC 4.1.1.

C.2 File Systems

EM algorithms, including PEMS, attempt to optimise disk access for performance reasons. Locality of reference and favourable access patterns (e.g. linear sweeps) provide the best performance, since disk seeking is extremely expensive. However, in practice most applications running on a modern operating system are not actually accessing disk directly – disk access is provided via a file system. This has the implication that a linear sweep in code may not actually translate to a linear sweep on disk due to file system fragmentation (files are generally not guaranteed to be contiguous ranges of blocks). This can result in unpredictable performance.

Thankfully, some file systems take this into consideration and provide facilities for allocating large areas of disk. The new default file system for Linux, ext4, includes this ability (support for “extents”). PEMS2 makes use of this functionality on systems modern enough to support this feature. All experiments in this thesis explicitly allocate disk via this mechanism.

Fig. C.1 shows the potential impact a fragmented filesystem can have. In this experiment, all parameters remain constant including \(n\) and only \(\mu\) is increased. That

\(^1\)i.e. 4 cores total
is, more disk space is used, but the actual problem size remains constant. Notice ext4 (with extents) has consistent performance regardless of the space used, but ext3 (without extents) degrades in performance severely as more space is used. Further experiments have shown that, without extents, disk performance can be very unpredictable.

Note that Fig. C.1 illustrates a particular pathological case. In particular, it is possible (with luck) to achieve nearly² equivalent performance without explicitly allocating disk. However, unless the file system is entirely empty, this is extremely unlikely for large files.

Those working with external memory algorithms should be careful to choose an appropriate filesystem, and make use of explicit disk space allocation routines (fallocate or posix_fallocate in Linux) to ensure good, predictable performance.

²Using extents actually reduces filesystem overhead in general, since blocks are much larger and less tree traversal is required. Thus, using extents can yield a performance improvement even on a completely unfragmented filesystem.
Appendix D

MPI Compatibility

Fig. D.1 shows the subset of MPI implemented by PEMS2. Additionally, malloc, realloc, and free are wrapped by PEMS to allocate memory in the virtual processor context rather than system RAM.

- MPI_Allgather
- MPI_Allgatherv
- MPI_Allreduce
- MPI_Alltoall
- MPI_Alltoallv
- MPI_Bcast
- MPI_Gather
- MPI_Gatherv
- MPI_Reduce
- MPI_Scatter
- MPI_Barrier
- MPI_Wtime
- MPI_Init
- MPI_Finalize
- MPI_Abort
- MPI_Comm_rank
- MPI_Comm_size

Figure D.1: Supported MPI Functions