SUPPORT FOR DATA MANAGEMENT IN OBJECT-ORIENTED DATA-PARALLEL SCIENTIFIC AND ENGINEERING SIMULATIONS

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Dedicated to My Family
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Abstract

Many data-parallel scientific and engineering simulations require the coordinated movement of large sets of data between multiple storage devices and multiple processors on distributed memory parallel machines. These data management requirements are not met by conventional file systems, so parallel file systems have been developed to support I/O in these programs. Current parallel file systems support I/O on simple data structures such as distributed rectangular arrays of primitive data types; however they do not provide adequate support for I/O on the complex distributed data structures often found in object-oriented parallel programs. This dissertation describes how class libraries can be used to extend the functionality of both parallel file systems and object-oriented database management systems to support the complex I/O requirements of object-oriented data-parallel simulations. The contributions of this dissertation are summarized below:

Several abstract models of parallel file systems are developed; the experimentally measured performance of two commercial parallel file systems is explained in the context of those models. This analysis should help application and library developers obtain high I/O performance from parallel file systems.

Three new abstractions for data management within the file I/O model are proposed. These abstractions can be used to extend parallel file systems to meet the special I/O requirements of object-oriented data-parallel simulations. Implementations of each abstraction are described.
A new abstraction for data management within the persistent data model is proposed. This abstraction can be implemented on object oriented database systems using language-specific class libraries; an implementation of this abstraction is described as well. This work shows how persistence can be used to support several common data management tasks in object-oriented simulations.

Finally, the file I/O model and the persistent data model of data management are compared, in the context of implementations of systems supporting these models for I/O in object-oriented parallel simulation programs. A number of metrics are applied to these systems to perform the comparison: performance, space requirements, productivity effects, ease of implementation, maintainability, and portability.
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Chapter 1: Introduction

1.1 Overview

1.1.1 Scientific and Engineering Simulations

For the purposes of this dissertation, let a *scientific/engineering simulation* be defined as a program that simulates the behavior of a physical system by iteratively solving a discretized system of equations that *models* the physical system by describing the ways in which relevant physical variables affect each other over time and through space\(^1\). Let the equations in such a system be called *model equations*. Analytic solution of model equations may be difficult, time consuming, or impossible, especially when complex (i.e. realistic) boundary and/or initial conditions are involved. For this reason, since the advent of computers, numerical solution of equations modeling physical systems (i.e. simulation) has played an increasingly important role in science, allowing scientists to develop and test complex models of natural processes.

Once a given physical system is well understood, simulation becomes a useful tool for *engineers*. An engineer can combine a mathematical model of a structure being designed with a mathematical model of the physical environment in which that structure is intended to operate. A simulation based on the resulting (combined) model allows detailed measurements of the structure’s predicted performance to be obtained, often at a

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\(^1\) Throughout this dissertation, *italics* are used to denote definitions; *underlining* is used to denote emphasis.
much lower cost than the alternative (building an actual small-scale or full-scale version of the structure, along with a system to measure its performance).

Parallelism is an important consideration in many scientific and engineering simulations (see Appendix A for an introduction to parallelism), for two reasons:

1) There tends to be a large potential for the exploitation of parallelism in these programs, since they generally simulate physical processes that act in parallel at many or all points in space simultaneously.

2) Scientists and engineers benefit from the exploitation of parallelism in simulations, since applying more computational power can allow:
   - a given simulation to run in a shorter amount of time, or
   - a given set of model equations to be discretized more finely and solved in the same amount of time, resulting in a more accurate solution, or
   - a more complex model to be simulated in a given amount of time.

1.1.2 Data Management in Simulations

The terms input/output (I/O) and data management are used interchangeably in this dissertation to refer to the controlled movement of data between transient storage (i.e. physical memory) and stable storage (i.e. disk). Many parallel scientific and engineering simulations have complex I/O requirements; they require the coordinated movement of large sets of data between multiple storage devices and multiple compute nodes on distributed memory parallel machines. These requirements are not met by conventional file systems, so parallel file systems have been developed to support I/O in these programs. Current parallel file systems support I/O on simple data structures such as distributed rectangular arrays of primitive data types; however they do not provide adequate support for I/O on the complex distributed data structures often found in object-oriented parallel simulations. But object-orientation is becoming an increasingly
important issue in the development of simulations, as scientists and engineers seek to obtain the benefits of object-orientation [10] in the development process. This dissertation describes how class libraries can be used to extend the functionality of both parallel file systems and object-oriented database management systems to support the complex I/O requirements of object-oriented parallel simulations.

The remainder of this chapter describes data management tasks common in parallel simulations, identifies two models for data management systems supporting those tasks, summarizes the contributions of this dissertation, and reviews related research.

1.2 Data Management Tasks in Parallel Scientific and Engineering Simulations

This section identifies several types of I/O tasks commonly required in scientific and engineering simulations.

1.2.1 Entire Database Transfer Tasks

In most simulations that predict the dynamic behavior of physical systems (i.e. the manner in which physical systems change over time given a set of initial conditions), simulation proceeds iteratively in discrete time steps; during the computation for each time step, a set of data that defines the predicted state of the simulated physical system is updated to reflect changes in the physical system that are predicted to occur in a certain amount of actual time (the time step size). Let a simulation’s data set or database be defined as the set of data representing the current state of a simulated physical system. Let an entire database transfer task be defined as a task that requires the transfer of an entire data set between memory and disk.
Several types of entire database transfer tasks commonly required in simulations are described below:

• **Initial and Final I/O:** *Initial input* is the transfer into memory of a data set representing initial conditions at the start of a simulation. *Final output* is the transfer of results from memory to permanent storage as a simulation progresses. Final output is required to obtain a permanent copy of a simulation’s predictions. In addition, final output is often used during debugging as follows. Many parallel simulations are manually parallelized versions of sequential programs. During the parallelization process developers often need to compare the predictions of the parallel and sequential versions (the predictions should be the same), to confirm that parallelization has not introduced bugs. The comparison of these results is often performed by outputting corresponding data sets from each program to permanent storage, and using a program such as UNIX’s `diff` to compare them.

• **Checkpointing and Restart:** Many simulations are exceptionally long-running. In order to avoid losing hours or days of results due to program and system crashes, these programs may periodically checkpoint, i.e. save their state, so that the state can be reloaded and computation restarted if a program or system crash occurs.

### 1.2.2 Concurrent Data Access Tasks

Let a *concurrent data access task* be defined as an I/O task in which a data set is accessed by both a simulation and a second program concurrently. Several types of concurrent data access tasks that are commonly performed in simulations are described below:

• **Concurrent Visualization and Computation:** Since parallel simulation applications tend to be long-running, users frequently wish to visualize partial results as a simulation progresses. Visualization facilities may be provided by a
separate program, which then requires a mechanism for accessing the data set concurrently with the simulation.

• **Computational Steering**: Users sometimes wish to change a simulation’s parameters as it executes. For example, if a simulation is executing too slowly, a scientist may wish to reduce the precision of an approximation. Changing this parameter without stopping and restarting the simulation can be accomplished if a second steering program can concurrently access (and modify) the simulation’s parameters. This is one form of computational steering.

### 1.2.3 Tasks Requiring Access to Large Databases

Some simulations operate on data sets that are too large to fit in core, i.e. in physical memory. Some data sets are too large to fit even within address spaces supported by virtual memory systems. Support for access to large databases is an important issue currently being addressed by researchers in both the computational science and database communities. This dissertation focuses on support for entire database transfer tasks and concurrent data access tasks; however, the parallel stream abstractions introduced in Chapters 3 through 5 could be applied to large database access tasks in which a simple sequential data access pattern is required.

### 1.3 Two Models for Systems Supporting Data Management Tasks

Several systems and abstractions for data management are developed in subsequent chapters; in this section, terminology is defined that will be used to describe these systems. Data management systems can be classified into several broad groups by data management model. A data management model specifies two things: the relationships
between the abstractions involved in data management (i.e. the relationships between programs, the data management system, program variables, transient storage, and stable storage), and the programmer interface to the data management system (i.e. the ways in which the programmer interacts with the data management system in order to accomplish data management tasks such as those identified in the previous section: entire database transfers, concurrent data access, and access to large data sets).

Two common data management models, the file I/O model and the persistent data model, are described in the following subsections.

1.3.1 File I/O Model

The data management systems discussed in chapters 3 through 5 operate within a file I/O model of data management; let such systems be called file I/O systems. In the file I/O model, program variables are transient, i.e. their values are always destroyed when the program terminates. Files (strings of bytes) are the units of stable storage and communication between programs. The file I/O system performs the work of moving data between variables and files, but the programmer must manage the variable-to-file mapping. The file I/O system gives the programmer primitives for initiating movement of data between program variables and files, and for flattening and unflattening variables (i.e. converting them to and from strings of bytes).

Data management abstractions in the file I/O model tend to be passive, performing no action until the programmer explicitly identifies an action to be performed. In the file I/O model, I/O tasks are generally coded as follows:

- **Entire database transfers** are coded with explicit commands that move data between files and memory.
• **Concurrent visualization and computation** can be performed by having the simulation program periodically write its data set to a file, and having the visualization program read this file when visualization is to be performed.

• **Computational steering** of a simulation can be coded similarly, by instructing the program to periodically read a file containing any parameters that are to be dynamically modifiable. If a steering program changes a parameter in this file as the simulation runs, the copy of the parameter in the simulation’s transient memory is eventually changed when the file is read.

• **Access to large databases** can be coded by simply storing the entire database in a single file. File I/O systems generally provide primitives that allow programs to sequentially or randomly read and write to files. In the file I/O model it is the programmer’s responsibility to manage the mapping between regions of the file and regions of a data set; in some cases this is a complex task, especially in simulations where the size or structure of the data set changes dynamically at run time.

### 1.3.2 Persistent Data Model

Chapter 6 introduces a data management system that operates within an alternate data management model: the persistent data model. In this alternate model, a program variable may be designated as *persistent*, meaning that it is to be stored in memory like an ordinary variable, but in addition it is to be associated with a second copy that is kept on stable storage. The data management system is responsible for the movement of data between memory and stable storage, and manages the mapping between the two. Persistent data on stable storage may be accessed by more than one program.
concurrently. So units of typed persistent data are the units of both stable storage and communication between programs in this model.

In the persistent data model, the data management system gives the programmer a means of identifying which program variables are to be persistent, a way of associating those variables with persistent data on stable storage, and a way of initiating, committing, and aborting transactions on those variables. Persistent variables can only be modified within transactions. When the programmer commits a transaction, this tells the data management system that the program’s persistent variables are in a consistent state, so at that point their values may be shared with other programs that may have requested access to those variables. When a transaction is aborted, the data management system must ensure that persistent variables are returned to the values they had when the transaction was initiated. Transactions may be aborted explicitly by the programmer or implicitly when a program or system failure occurs. So the data management system is responsible for returning all persistent data to a consistent state when a program is restarted after a program crash or a system failure.

Persistent data systems tend to be more active than file I/O systems in that they may spontaneously (without programmer intervention) initiate the transfer of data between stable storage and memory when this is needed; this is known as transparent I/O.

In the persistent data model, the I/O tasks identified in Section 1.2 can be coded as follows:

- **Initial and final I/O** are automatically performed by the persistent data system when persistent objects are declared and when the program terminates, respectively.

- **Checkpointing**\(^1\) (i.e. saving a simulation’s state) is performed transparently when the programmer commits a transaction; in the persistent data model, committing a

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\(^1\) Do not confuse this computational science term with its homonym, the database systems term "checkpointing" meaning writing database buffers to disk.
transaction within which a data set has been modified causes the data management system to ensure that the modified data is written to stable storage.

- **Concurrent visualization and computation** can be performed by using the persistent data system from within the visualization program to access the persistent data within the simulation. The persistent data system’s transaction mechanism manages the concurrent data access using a locking mechanism such as the one described in Figure 6.3.

- **Computational steering** of a simulation can be coded similarly; the persistent data system can be used within the steering program to concurrently access and modify the simulation’s persistent data as the simulation runs.

- **Access to large databases**: Many persistent data systems are built on object-oriented database management systems (*OODBMS*), some of which have special *query languages* that support optimized access to subsets of large databases. A detailed discussion of OODBMS query languages is beyond the scope of this dissertation; see [39] for more information.

### 1.4 Contributions

This dissertation makes several contributions to the understanding of how data management can be supported in object-oriented data-parallel simulations; these contributions are summarized in this section. An expanded description of these contributions is presented in Section 8.1 of the Conclusion chapter.

Several abstract models of parallel file systems are developed in Chapter 2, and the experimentally measured performance of two commercial parallel file systems is explained in the context of these models. This analysis is intended to help application and library developers obtain high I/O performance from parallel file systems.
Three new abstractions for data management within the file I/O model are proposed in Chapters 3 through 5. These abstractions can be implemented on parallel file systems using language-specific class libraries; implementations of each abstraction are described. These abstractions provide functionality lacking in current parallel file systems, functionality that is required if the special I/O requirements of object-oriented parallel simulations are to be met. Chapter 3 introduces \textit{parallel filestreams}, an extension of ordinary C++ filestreams that supports node-buffered parallel I/O with flattening and unflattening in SPMD parallel C++ programs. \textit{Object-parallel filestreams}, introduced in Chapter 4, extend parallel filestreams to support the element buffering required for file-format-independent I/O on distributed arrays of objects in languages such as pC++. \textit{Distributed array streams}, described in Chapter 5, extend object-parallel filestreams to support a simplified programmer interface as well as automatic bookkeeping of array distribution meta-data in special-format files.

A new abstraction for data management within the persistent data model is proposed in Chapter 6. This abstraction can be implemented on object oriented database systems (\textit{OODBMS}) using language-specific class libraries; an implementation of this abstraction is described. This work shows how persistence can be used to support several common data management tasks in object-oriented parallel simulations.

Chapter 7 carries out a comparison of how well file I/O systems implemented on parallel file systems and persistent data systems implemented on OODBMS support I/O in object-oriented parallel simulations. The comparison is based on experience gained in the implementation of the abstractions described in Chapters 3 through 6, and involves a number of metrics: performance, space requirements, productivity effects, ease of implementation, maintainability, and portability.
1.5 Related Work

Figure 1.1 presents an overview of several research efforts involving data management in parallel programs, and shows where the systems described in this dissertation fit in. Parallel filestreams (Chapter 3), object-parallel filestreams (Chapter 4), and distributed array streams (Chapter 5) differ from ordinary C++ filestreams in that they support I/O on distributed data structures on distributed memory parallel machines; as discussed in Section 3.1, the single-buffered design of ordinary C++ filestreams make them incompatible with existing parallel file systems. These three new parallel stream abstractions also differ from the many research efforts that have been directed towards support for parallel I/O on distributed data structures, in that (like C++ filestreams) the
parallel stream abstractions introduced in this dissertation support I/O on arrays having arbitrarily complex, variable sized elements. The persistent collection abstraction introduced in Chapter 6 differs from all of the abstractions in the left two thirds of Figure 1.1 in that it operates within a persistent data model of I/O, rather than a file I/O model.

The following three subsections describe in more detail several of the research efforts related to the work described in this dissertation.

1.5.1 Parallel File System Performance

A number of studies of parallel file system performance have been undertaken. Kwan and Reed [40] describe the performance of the Scalable File System of the CM-5 at NCSA. Their motivation was to compare the performance of the CM-5 parallel I/O interfaces of CM-Fortran and C/CMMD. Bordawekar, Rosario and Choudhary present an experimental performance evaluation of the Intel Touchstone Delta Concurrent File System in [5]. Chapter 2 differs from this work in that in addition to describing and analyzing the performance of commercial systems, a set of abstract parallel file system models are developed and the performance of two commercial systems is explained in relation to those models. Unlike most of the related work, the emphasis in Chapter 2 is not on measuring the performance of specific I/O systems, but in identifying more general relationships between parallel file system architecture, application I/O behavior, and I/O performance.

1.5.2 Support for File I/O in Parallel Programs

A large number of experimental systems have been developed to support file I/O in parallel programs. The libraries PetSc/Chameleon [29] from Argonne and Panda [46] [45] from UIUC, and the software system PASSION [15] from Syracuse, support I/O on distributed arrays of fixed-sized elements in the context of the distributed-memory
data-parallel programming model. PetSc/Chameleon supports I/O on block-distributed arrays. Panda supports more general HPF-style array distributions and interleaving, as does d/streams. PASSION is a large effort at Syracuse that provides support at the language, compiler, runtime, and file system level for I/O on distributed arrays as well as for computation over out-of-core distributed arrays. (It should be noted that distributed array streams can only be used for out-of-core computation in the simple case where the simulation’s data access pattern is sequential.) The two-phase access strategy described in PASSION for parallel access to files, where data is first read in a manner conforming to the distribution on disk and then redistributed among the processors, is similar to the implementation of the distributed array streams load method. Language extensions for parallel I/O on distributed arrays have been discussed in the HPF Forum [26].

The distributed array streams implementation described in Chapter 5 differs from these systems in that it supports buffered I/O on distributed arrays of objects whose size may vary over the array itself, for example distributed arrays of variable-density grids or distributed arrays of lists of particles, in the context of a portable object-parallel language.

1.5.3 Support for Persistent Data in Parallel Programs

Several systems other than persistent collections have been developed to support persistence in parallel programs, but not nearly as much work has been done in this area as in the area of support for file I/O. ParSets [21], a component of the Shore [7] OODBMS developed at the University of Wisconsin at Madison, is similar to the pC++ implementation of persistent collections described in Chapter 6, in that it provides support for data-parallelism in the form of parallel application of methods of persistent objects. The pC++ programming system, when combined with persistent collections, differs from Shore ParSets in that it supports programmer-controlled alignment and distribution of the elements of aggregate data-structures, and the parallel application of
operators acting simultaneously on multiple aggregate persistent data-structures. It may be possible to provide these facilities in a ParSet abstraction; however, the initial ParSets paper does not describe how such support could be provided.

The Raven [1] operating system and programming language supports task-parallelism and persistent objects, but unlike pC++ persistent collections, no facilities are provided for data-parallelism. Shared Data Abstractions [12], developed at the University of Vienna and ICASE at NASA Langley, extend the support for data-parallelism in FORTRAN 90 with support for task-parallelism and for simplified I/O on data modules, but no support is provided for concurrent multi-program access to data (unlike the persistent collection implementation described in Chapter 6, which does support concurrent access).

A large amount of work has been done on object-oriented database programming systems providing concurrent distributed access to persistent data; however, the emphasis has been on distribution supporting inherently distributed data or inherently distributed users, rather than distribution supporting parallelism for speedup (as is supported in pC++ persistent collections). In the domain of relational DBMS, a large body of work exists on parallel implementation of internal DBMS operations such as the processing of joins. However, this work does not address explicit parallelism in application code, which is useful for exploiting parallelism available in the problem domain (explicit parallelism is supported in pC++ persistent collections).

1.6 Summary and Dissertation Outline

This introductory chapter has identified the data management tasks common to parallel scientific and engineering simulations, described two possible models for systems supporting the data management requirements of those tasks, described the contributions
of this dissertation in terms of those models, and referenced other research work having similar goals.

The remainder of this dissertation is organized as follows. Chapter 2 introduces parallel file systems and relates their performance to their internal architecture and to application I/O patterns. Chapters 3 through 6 describe designs and implementations of four new abstractions for data management within the file I/O and persistent data models in SPMD C++ and object-parallel programs. Chapter 7 evaluates how well file I/O systems implemented on parallel file systems and persistent data systems implemented on OODBMS support I/O in object-oriented parallel simulations. Chapter 8 summarizes the results of this dissertation and describes directions for future research, and Appendices A and B present brief introductions to parallelism and database management respectively.
Chapter 2: I/O Performance in Parallel File Systems

Restricting a parallel machine to the use of a single I/O device would cause I/O to become a bottleneck, since I/O power would be unable to scale with computational power. Therefore, many current distributed memory parallel machines provide a scalable parallel file system (PFS) consisting of several I/O processors each connected to a high speed I/O device. This enables high bandwidth cooperative I/O i.e. parallel I/O to be performed from the compute nodes [47] [36] [17]. Many large-scale parallel applications require the use of parallel I/O to achieve high performance [44].

This chapter introduces parallel file systems, defines several measures of PFS performance, and relates PFS performance to application I/O patterns and to various elements of PFS internal architecture, such as the use of buffering and virtual memory on the I/O nodes. In addition, experimental results are presented that relate the performance of PFS on the Intel Paragon and the Thinking Machines CM-5 to the PFS models. The models and results presented in this chapter should be useful for optimizing I/O performance in applications and I/O libraries, configuring existing PFS and designing new ones. This work was first described in [32].
2.1 Introduction to Parallel File Systems

Figure 2.1 shows the architectural elements of the PFS of several current distributed memory parallel machines [43] such as the Intel Paragon and Thinking Machines CM-5. These PFS support an interface similar to that of a normal UNIX file system. In addition, they are accessible through special system calls provided for parallel I/O. The interface for parallel I/O is similar to the interface for normal UNIX I/O, but an I/O mode associated with each file descriptor coordinates the way in which the compute nodes access and modify files.

2.1.1 An Example Program

This section describes an example program that shows how parallel file systems are used to support the I/O requirements of "polygon overlay", a pC++ [30] program that reads in two lists (or "vectors") of polygons (the "left" and "right" vectors), computes a new polygon vector by overlaying the left and right vectors, then outputs the overlay vector. This program is discussed in detail in [49]. (For a brief description of pC++ see "The
The parallel file system interface described here is supported by the parallel filestream abstraction, which is described in more detail in Chapter 3. Parallel filestreams support additional functionality not found in current parallel file systems; however, the polygon overlay program does not require the use of this additional functionality, so in this case parallel filestreams function purely as I/O middleware, supporting application portability by providing a single I/O interface over multiple parallel file systems. The more advanced features of parallel filestreams are described in Chapter 3.

```c++
void Polygon::readPolygon() {
    // Read the local part of a DISTRIBUTED vector of polygons
    // into leftVec on each node
    leftVec = (polyVec_t *)malloc(sizeof(polyVec_t));

    // Open a read-only input pfstream (s) on each node
    ipfstream s(leftFilename, pfilebuf::rdonly);

    // Read the total size of the polygon vector
    s.setparmode(pfilebuf::unixpar);
    s.unbufread((char *)&totalLeftLength, sizeof(int));

    // Calculate the local size of of the vector, store in leftVec->len
    ...

    // Read the local part of the vector into leftVec->vec
    leftVec->vec = (poly_t *) malloc(sizeof(poly_t)*leftVec->len);
    s.setparmode(pfilebuf::syncpar);
    s.unbufread((char *)leftVec->vec, leftVec->len * sizeof(poly_t));
}

[...]

// Read a COMPLETE COPY of a second vector of polygons
// into rightVec on EACH NODE
rightVec = (polyVec_t *)malloc(sizeof(polyVec_t));

ipfstream s(rightFilename, pfilebuf::rdonly);

s.setparmode(pfilebuf::unixpar);
s.unbufread((char *)&(rightVec->len), sizeof(int));

rightVec->vec = (poly_t *) malloc(sizeof(poly_t)*rightVec->len);
s.unbufread((char *)rightVec->vec, rightVec->len * sizeof(poly_t));
}
```

Figure 2.2: Using a Parallel File System for Input

pC++ Language" on Page 133.) The parallel file system interface described here is supported by the *parallel filestream* abstraction, which is described in more detail in Chapter 3. Parallel filestreams support additional functionality not found in current parallel file systems; however, the polygon overlay program does not require the use of this additional functionality, so in this case parallel filestreams function purely as *I/O middleware*, supporting application portability by providing a single I/O interface over multiple parallel file systems. The more advanced features of parallel filestreams are described in Chapter 3.
An early version of this program that used ordinary UNIX I/O system calls and no parallel I/O had 200 lines of code devoted to I/O, approximately 10% of the total code in the program. Rewriting the original UNIX I/O using a parallel file system reduced the I/O code from 200 to 70 lines, a 65% reduction. Most of this reduction was due to the fact that in the original program, a significant amount of code was required to copy pieces of the (distributed) output polygon vector from their home nodes to node zero, which performed all of the (sequential) file output. When using a parallel file system, all of the nodes of a parallel machine can participate in I/O, so this communication is unnecessary.

**Input**

Figure 2.2 shows how parallel filestreams (pfstreams) can be used to input the left and right polygon vectors in the function readPolygon() of polygon overlay. After allocating memory for the left vector, before reading it in from the file, an input pfstream $s$ is declared as follows:

```cpp
ipfstream s(leftFilename, pfilebuf::rdonly);
```

This opens the file `leftFilename` and attaches the stream $s$ to it.\(^1\)

The left vector is a distributed data structure, i.e. it doesn’t appear in its entirety on any one node, but is distributed across the nodes of the parallel machine. On the other hand the right vector is copied in its entirety into each node. The parallel file systems of the Paragon and CM-5 support I/O modes that allow the programmer to use parallel I/O for each of these types of input. (The names used for these modes in the parallel filestreams abstraction correspond to the I/O mode names used in the Paragon’s parallel file system.) In `unixpar` mode, each processor has its own file pointer and can

---

1. The programmer interface to parallel filestreams is similar to that of ordinary C++ filestreams. A complete description of the parallel filestream interface is presented in Chapter 3. For the purposes of this discussion, the important issue is not parallel filestream syntax, but the way in which the underlying parallel file system’s I/O modes can be used to coordinate parallel I/O from within a parallel application.
perform I/O independently of the other processors. This mode is useful for reading a
copy of a single block of data in the file into each processor in parallel, and for reading
pC++ global variables. The total number of polygons in the left polygon vector is to be
read into the pC++ global variable totalLeftLength, so at this point the parallel I/O
mode of the stream s is set to unixpar mode:

\[ \text{s.setparmode(pfilebuf::unixpar);} \]

Next an unbuffered read of sizeof(int) bytes is performed from the stream s
directly into the global variable totalLeftLength:

\[ \text{s.unbufread((char *)&totalLeftLength, sizeof(int));} \]

The parallel filestream abstraction also supports buffered I/O, which can yield higher I/O
performance when I/O is to be performed on large data sets stored non-contiguously in
memory. However, when data are mostly contiguous, as in this program, unbuffered I/O
is more efficient.

The left polygon vector is to be distributed across the elements of the Polygon
collection using a block mapping from the file to the collection elements. Now that the
total size of this vector is known, the number of polygons to be read into each element is
computed. Then after allocating memory to store the local segment of the vector, the
parallel I/O mode of the stream is set to syncpar mode, in which all the processors use
a single file pointer, reads and writes are synchronizing operations, and data is
distributed from the file to the nodes using a block distribution strategy.

\[ \text{s.setparmode(pfilebuf::syncpar);} \]

In this I/O mode, each node must specify the amount of data it requires from the file
(different nodes may request different amounts). The data blocks for the nodes are read
from the file and are sent to each processor in parallel. Next the local block of data of
size

\[ \text{leftVec->len * sizeof(poly_t)} \]
void Polygon::writePolygon()
{
    int totalOutVecLength;

    // Compute total size of the distributed vector outVec
    totalOutVecLength = outVec->len;
    pcxx_ReduceIntAdd(&totalOutVecLength, 1);

    // Open a write-only output pfstream
    ofstream s(outFn, pfilebuf::wronly | pfilebuf::creat | pfilebuf::trunc, 0644);

    // Use pfilebuf::unixpar mode since we're writing a global variable.
    s.setparmode(pfilebuf::unixpar);
    s.unbufwrite((char *)&totalOutVecLength, sizeof(int));

    // (See the description of pfilebuf::syncpar mode in readPolygon())
    s.setparmode(pfilebuf::syncpar);
    s.unbufwrite((char *)outVec->vec, outVec->len * sizeof(poly_t));
}

Figure 2.3: Using a Parallel File System for Output

is read from the stream $s$ directly into the memory pointed to by $\text{leftVec->vec}$:

\[
\text{s.unbufread((char *)leftVec->vec, leftVec->len * sizeof(poly_t))};
\]

At this point input is completed; the file is closed automatically when the program block containing the declaration of the stream $s$ is exited.

Next the "right" vector is read from $\text{rightFilename}$ in a similar manner, except that this vector is not a distributed data structure; a copy of the entire vector is to be stored on each processor. For this reason, $\text{unixpar}$ mode is used to read this vector, rather than $\text{syncpar}$ mode (which was used for $\text{leftVec}$).

Output

Later in the program, the overlay (output) vector $\text{outVec}$ is computed and then output by the $\text{writePolygon()}$ method (see Figure 2.3). The overlay vector is a distributed data structure having a block distribution. Before it is output, its total size is first computed using a reduction operation. Next an output parallel filestream is declared, and the total size is written to the file using $\text{unixpar}$ mode (since only one copy of the
total size should appear in the file, and each node’s file pointer should be advanced past the just-written size prior to the next write).

Next the local part of the output vector is written to the file using syncpar mode. In syncpar mode, writes are synchronizing operations, and the parallel file system assumes the data structures being written have a block distribution across the nodes; the blocks of data (one per node) are written to the file in order of node number. The file is closed automatically by the stream’s destructor at the end of the block in which the stream was declared.

### 2.1.2 Performance Measures

This section defines several parallel file system performance measures. In later sections, these measures are used to describe the performance both of models of PFS and of commercial PFS.

Assume a parallel machine has $p$ compute nodes and one I/O node. Also assume that:

- An application, when run on the compute nodes, issues a parallel input or a parallel output request to the I/O system.

- The parallel I/O request consists of a system call executed by each compute node, which causes a total of $n$ bytes to be concurrently read or written ($n/p$ bytes from each compute node).

- The time for I/O is the same on each compute node.

For such a system let the following I/O performance measures be defined:

- The *application I/O time* denoted by $Time_{app}(n)$ is the time the system call takes to move data between the application's buffers and the I/O system.
• The actual I/O time denoted by $Time_{actual}(n)$ is the actual time for the I/O operation to complete, i.e. the time taken for the I/O system to move data all the way between the application and the disk.

These times are significant because while data is being transferred between the application’s buffers and the operating system, the application must not disturb the application memory holding the data, and until the I/O operation completes, the data cannot be used by the application (in the case of input) or by other applications (in the case of output).

A rate can be associated with each of these times:

• The application I/O rate denoted by $Rate_{app}(n)$ is:

  \[ n / Time_{app}(n) \]

• The actual I/O rate denoted by $Rate_{actual}(n)$ is:

  \[ n / Time_{actual}(n) \]
In this section a simple model of a PFS is described and successively refined in three steps. The initial model is a system with unlimited memory on the compute nodes and no I/O system memory. In the second model, an I/O buffer is added to the I/O node, and in the third model, virtual memory is added on both the compute and I/O nodes. Figure 2.4 will be referred to throughout the following three subsections; it depicts the internal architecture of these three models, along with the expected I/O performance at each stage. The models described in this section are meant to explain how the interaction of buffering, virtual memory, and application I/O patterns can affect I/O performance.
2.2.1 Unlimited Compute Node Memory and No I/O System Memory

The first parallel file system model that will be analyzed is a system with unlimited memory on the compute nodes and no memory on the (single) I/O node (see the left side of Figure 2.4). In this case, the I/O node moves data directly between the application’s memory (on the compute nodes) and the disk. So by the definition of $\text{Time}_{\text{app}}(n)$ and $\text{Time}_{\text{actual}}(n)$,

$$\text{Time}_{\text{app}}(n) = \text{Time}_{\text{actual}}(n)$$

and this implies that

$$\text{Rate}_{\text{app}}(n) = \text{Rate}_{\text{actual}}(n).$$

Let this rate be called $\text{Rate}(n)$.

As $n$ increases, one would expect to see $\text{Rate}(n)$ start low at first due to latency, and then quickly increase to some limiting rate corresponding to the raw I/O rate of the physical disk device (see the first graph in Figure 2.4). After this point there should be no change in I/O performance when the the size of the data set is scaled upwards. So if the size of the program’s data set is increased by some factor $f$, the time for I/O on that data set should increase by the same factor $f$.

2.2.2 Finite Physical Memory Buffer in the I/O System

Now assume the I/O node has an I/O buffer of finite size $b$ bytes of physical (not virtual) memory (see the central part of Figure 2.4). In this model and the next the discussion is limited to output for clarity.

Parallel output in this model consists of two concurrent processes. Data is transferred 1) from the application through the operating system to the disk buffer, and 2) from the disk buffer to the disk. Assume the transfer of data from the application's
buffer to the disk buffer happens more quickly than the transfer from the disk buffer to
the disk.

As the center graph in Figure 2.4 shows, when the total amount of output \((n)\) that the
application performs is less than the size of the disk buffer \((b)\) then the application output
rate will be much higher than the actual output rate, since the operating system can
quickly copy the contents of the application's buffers into the I/O node's buffer.
Therefore the system calls can complete before the data is actually written to the disk.
When \(n\) is very small, latency will slow the application output rate, so a graph of the
application output rate as a function of \(n\) will start low, rise quickly, then asymptotically
approach a value corresponding to the peak rate at which the operating system can
transfer data from the compute nodes into the I/O node's buffer.

However when the amount of output performed is greater than the I/O node’s buffer
size, the system calls will start to take longer to return, for the following reason. After
the disk buffer fills, the transfer rate from the compute nodes to the I/O buffer will be the
same as the rate at which data is transferred from the I/O buffer to the disk. Therefore as
the output size is increased past \(b\) bytes, the graph of the average application output rate
will start dropping, asymptotically approaching the actual output rate.
2.2.3 Virtual Memory on Compute and I/O Nodes

Actual parallel systems have finite physical memory on both the compute and I/O nodes but provide a larger virtual memory through paging mechanisms. In this section the model from the previous section is extended to include virtual memory on both the compute and I/O nodes (see the right portion of Figure 2.4). Assume the size of the I/O node’s buffer is \( b \) bytes and the physical memory size on both the compute nodes and the I/O node is \( m \) bytes, where \( b > m \) (so the I/O node’s buffer is too large to fit into the I/O node's physical memory).

Figure 2.5: Comparing Output Performance With and Without Virtual Memory on the I/O System

(This figure compares the performance of the models in Sections 2.2.2 and 2.2.3). This data was obtained from a simple virtual memory simulator, in which it was assumed that the raw output disk bandwidth is one fifth the interconnection network bandwidth, and that the raw paging disk bandwidth is one half the raw output disk bandwidth. This graph suggests that the use of virtual memory for the I/O system can improve application output rates at the expense of actual output rates.

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For \( n < m \) the behavior of this model is the same as that of the model in the previous section. When \( n > m \) paging starts on the I/O node. At this point the system’s behavior will be complex, and will be determined by the paging algorithm and the raw I/O rates of the paging disk and the actual disk. Also when the total output size is greater than the total size of the compute nodes’ physical memory (\( n > p^*m \)), paging will occur on both the I/O node and the compute nodes. Analytic modeling of this behavior is difficult, so a simple virtual memory system simulator was developed to examine this behavior.

**Figure 2.6: Comparing Paging Algorithms for the Model with Virtual Memory on the I/O Nodes**

(Simulator data). Both actual and application performance start to drop (due to paging) when the total amount of data written equals the total physical I/O system memory. The severity of the drop depends on the paging algorithm used. With a simple demand paging algorithm, the actual rate drops to 60% of the raw disk rate when 5 times the I/O system memory size is written. With an improved paging algorithm that incorporates early page-in of paged blocks to unused I/O system memory, the actual rate drops to 85% of the raw disk rate, a smaller drop than for the simple algorithm.

For \( n < m \) the behavior of this model is the same as that of the model in the previous section. When \( n > m \) paging starts on the I/O node. At this point the system’s behavior will be complex, and will be determined by the paging algorithm and the raw I/O rates of the paging disk and the actual disk. Also when the total output size is greater than the total size of the compute nodes’ physical memory (\( n > p^*m \)), paging will occur on both the I/O node and the compute nodes. Analytic modeling of this behavior is difficult, so a simple virtual memory system simulator was developed to examine this behavior.
Figures 2.5 and 2.6 depict the results of experiments run on this simulator. Figure 2.5 shows that the use of virtual memory for the I/O system can improve application output rates at the expense of actual output rates. Figure 2.6 shows that if I/O system virtual memory is used then specialized paging algorithms, such as early page-in to unused I/O system memory, can improve actual output performance compared to the performance that can achieved with simpler paging algorithms.

So far it has been assumed that the paging disk device and the destination disk device have the same raw I/O rate. Consider the case where the paging disk is slower than the actual disk. For a write of \( n \) bytes where \( n \) is large, by the time all the data has been transferred out of the compute nodes, a large amount of data will have been transferred onto the paging disk. After this, the amount of data in the I/O node’s buffer will begin to drop, since that data will be transferred out of the buffer by the actual disk faster than it is transferred in to the buffer from the paging disk. At the point where the amount of data in the I/O node’s buffer drops to zero, the rate at which data is written to the actual disk will drop to the paging disk’s raw I/O rate. Therefore, when the paging disk is slower than the actual disk, for large \( n \), it becomes important to avoid causing paging on the I/O node during output. Paging on the I/O node can be avoided by writing only enough data to fill the I/O node’s memory, and then waiting for the memory to be written to the actual disk before writing more data.

### 2.3 Experimentally Measured Parallel I/O Performance

This section describes the performance of parallel file systems on two commercial parallel machines, and relates their measured performance to the performance patterns predicted by the PFS models described in the previous section.
Parallel file systems on some commercial parallel machines (such as the Intel Paragon) perform better when a large parallel write is broken into a series of smaller ones. Let a parallel write operation be defined as a single parallel write system call performed on each compute node of a parallel machine, and a parallel output operation as a series of one or more parallel write operations with no intervening computation. As mentioned in the previous section, there are two significant amounts of time associated with a given parallel output operation: the application output time and the actual output time. Also there are two data transfers rates, the application output rate and the actual output rate, associated with these times. Let parallel read operation, parallel input operation, application input rate, and actual input rate be defined similarly.

The following experimental results for PFS input performance are all in terms of the actual input rate, for the following reason: after initiating a read operation, there will be some period of time during which no data is being transferred into the user's program, while the data is being read from the disk and sent through the network. But neither the Paragon nor the CM-5 provide a way to measure that time. So there is no way to measure the remaining time when data is being transferred into the application's buffers. Therefore, even though it makes sense to talk about an application input rate for a parallel read operation, there is no way to measure that rate on the Paragon or the CM-5.

### 2.3.1 Intel Paragon

This section describes the results of PFS performance experiments that were run on the Intel Paragon at Indiana University in the Fall of 1994. This machine had 92 compute nodes and 2 I/O nodes. Each I/O node had 2 RAID-3 arrays; each array had 5 disks for a total of 10 disks. The programs described here were the only ones running on the machine during most of these experiments. The I/O nodes on this machine were in a service partition. This is not the most optimal configuration since the service partition
nodes also run all the UNIX commands issued by users; these commands use up potential I/O buffer memory.

The test program that was used performed repeated I/O operations, linearly increasing the total amount of I/O. The following optimizations, suggested by the Paragon User's Guide [36], were incorporated into the test program:

1) Read/write buffers were aligned to page boundaries. If properly aligned buffers are not used, the system must copy the data to new, aligned buffers, worsening performance.

2) The Paragon's record (M_RECORD) parallel I/O mode was used. This mode requires that the same amount of data be read or written by each processor in any given parallel I/O operation.

3) Large write operations were broken into multiple smaller operations, where the amount of data written by each processor is either equal to the stripe unit size, or is an integer multiple of the full stripe size. This makes optimum use of file striping, keeping all the I/O nodes busy at once.

4) Read and write requests were initiated with the file pointer located on file system block boundaries.
Figure 2.7: Paragon Input Performance
(in terms of the actual input rate). Input performance drops when the input size exceeds the physical memory of the compute nodes, causing paging.

Parallel Input Performance

Figure 2.7 is a graph of the IU Paragon’s actual input rate for varying sizes of parallel input operations, each comprised of a single parallel read operation, for 2, 4, and 8 processors. The input performance suddenly drops at approximately 8 MBytes of data input per node; at this point the input size exceeds the physical memory of the compute nodes, causing paging.

The Paragon User's Guide indicates that parallel write performance can be improved by breaking large parallel write operations into a series of smaller operations, where the amount of data written by each operation is equal to the stripe unit size. During the course of this experimentation it was determined that the corresponding optimization for input (breaking up large parallel read operations into smaller read operations) does not improve input performance on the Paragon.

Parallel Output Performance

Figures 2.8 through 2.12 describe the empirically measured parallel output performance of Indiana’s Paragon. Parallel output operations in these experiments were comprised of
Figure 2.8: Comparing the Paragon’s Application and Actual Output Performance measured using two compute nodes on IU’s Paragon. The graph shows that the actual output rate remains nearly constant, no matter how much data is written. The graph of the application output rate, on the other hand, rises to a peak around 5MB then suddenly begins to drop. A possible cause of this behavior becomes apparent in Figure 2.9.

a series of smaller parallel write operations of size equal to the I/O system’s stripe unit size.

Figure 2.8 compares the Paragon’s application and actual output rates for output operations of varying size. Note the sudden drop in the application output rate for output operations larger than 5 MBytes. Figure 2.12 shows that this same peaked shape appears after about the same amount of data has been written, regardless of the number of compute nodes involved in the output operation.

The model described in the previous section predicts that if small parallel write operations are performed repeatedly, then the application output rates of those operations should be high until the memory of the I/O nodes is filled, at which point the application
output rates should suddenly drop due to the extra time required for paging on the I/O nodes. This appears to be the cause of the drop in application output rate observed in Figure 2.8. See Figure 2.9, which relates the empirically measured application output rate of parallel output operations of various sizes to the application output rates of the individual parallel write operations that make up those parallel outputs. After 6 Mbytes have been written, the application output rate of the individual parallel write operations suddenly drops. This point roughly corresponds to the total size of the memory on the Paragon’s I/O nodes, so it is likely that this drop is caused by the I/O system’s memory becoming filled. The sudden drop in the rate for the individual writes is what causes the observed peak in the rate for the output operations that those writes comprise.

**Figure 2.9: Paragon Application Output Rates for Output Operations and Their Constituent Writes**

The application output rate for an output operation of a given size is shown in black directly above that size; the rates for the individual write system calls making up that output operation appear in red to the left of that size.
It should be possible to avoid a drop in the application output rate on the Paragon by making sure not to cause paging on the I/O nodes. This could be accomplished by repeatedly writing just enough data to fill the I/O nodes’ memory, then pausing until that data is completely transferred to disk. To test this theory, the application output rates of each of a series of many small writes were measured in two runs (see Figure 2.10). In one run output was paused using the UNIX \texttt{fsync()} system call after roughly 3MBytes of data were written. (\texttt{fsync()} forces a pause until all data is completely written to disk.) In the second run, output was not paused. In both runs there is a drop in the application output rate after a total of about 2.5MBytes of data has been written. In

**Figure 2.10: Using fsync() to Avoid Paging on the I/O Nodes**

Results for two runs are shown, one in which an \texttt{fsync()} call was performed after a total of 3MBytes of data were output, and one without the \texttt{fsync()}. (The individual writes were all of equal size; the rate for the write that brings the total amount of data output in a given run to b bytes appears at b on the x-axis.) The rate starts high in both graphs, then suddenly drops when the I/O system’s memory is filled, due to the onset of paging. But in the run with \texttt{fsync()}, the rate jumps up after the \texttt{fsync()}.
the run with the \texttt{fsync()} , the rate jumps up again at 3MBytes after writing is resumed, while the rate remains low in the run without \texttt{fsync()}. So it would appear that one can avoid a drop in the application output rate by using \texttt{fsync()} to wait for the I/O nodes’ memory to clear after writing enough data to fill it.

Figure 2.11 shows the results of an experiment similar to the one described in Figure 2.10, but where three times as much data was written. In one of the runs in Figure 2.11, \texttt{fsync()} is periodically called before the I/O nodes’ memory fills (instead of just being called once as in Figure 2.10). The model described earlier predicts that as data continues to be written to the I/O nodes, in the run without \texttt{fsync()}, the application output rate should remain low indefinitely after the point where the I/O nodes' memory fills up. The behavior of the non-\texttt{fsync()} run in Figure 2.11 is more complex than the model predicts; the rate jumps up periodically, as if the I/O nodes’
buffers had spontaneously and periodically been cleared. Perhaps this behavior is due to perturbations caused by I/O from other applications or by I/O from the operating system itself.

### 2.3.2 Thinking Machines CM-5

This section describes results of experiments run on the CM-5 at UMIACS, Maryland in the Fall of 1994. This machine had 32 compute nodes and 2 I/O nodes. A wall clock timer was used to obtain these measurements, even though there were other users on the system, since the other timers on the CM-5 cannot be used for I/O measurements. Figure 2.13 shows the UMIACS CM-5’s input and application output performance. The drop in performance for I/O operations greater than 6 MBytes per node is probably due to paging caused by virtual memory on the compute nodes. Figure 2.14 compares the

![Figure 2.12: Paragon Application Output Rates for Varying Numbers of Processors](image)

Notice that the performance drop happens at around the same point (close to 5MB) regardless of the number of processors from which the output takes place. It appears that this drop occurs when the I/O system’s memory becomes filled, causing the onset of paging on the I/O nodes.
application output performance of Maryland’s CM-5 to that of Indiana’s Paragon. Following the publication of [32] (in which this work was first described), Intel reconfigured the I/O system of the Paragon at Indiana, improving its I/O performance.

Figure 2.13: Input and Application Output Performance of the Maryland CM-5. Notice the performance drop for both input and output rates beyond 6MB.
The I/O optimizations that were identified during the course of these experiments are summarized below. These optimizations should be applicable on the Paragon and the CM-5, as well as other machines having similar I/O architectures.

- Blocks of data to be output from the application should be properly aligned in memory so that the operating system doesn't have to make an aligned copy of the data during output.

- Striping mechanisms provided in parallel I/O systems should be used to balance the I/O load among the components of the parallel I/O system. This may require

Figure 2.14: Paragon And CM-5 Application Output Performance for IU's Paragon and Maryland's CM-5. Both measurements are for 32 processors. Each Paragon output operation was comprised of a series of parallel write operations, each of a size equal to the stripe unit size; each CM-5 output operation was comprised of a single parallel read/write operation.

2.4 I/O Optimizations

The I/O optimizations that were identified during the course of these experiments are summarized below. These optimizations should be applicable on the Paragon and the CM-5, as well as other machines having similar I/O architectures.

- Blocks of data to be output from the application should be properly aligned in memory so that the operating system doesn't have to make an aligned copy of the data during output.

- Striping mechanisms provided in parallel I/O systems should be used to balance the I/O load among the components of the parallel I/O system. This may require
carefully setting the size and number of the individual I/O requests making up I/O operations.

- If virtual memory is used for the I/O system then the programmer can improve the actual output rate if I/O system paging is avoided by writing only enough data to fill the I/O system memory, then waiting until that data is completely written to the disk before writing more data. This technique will improve the application output rate as well.

- The simulation results that were described suggest that the use of virtual memory for the I/O system can improve application output rates at the expense of actual output rates. If I/O system virtual memory is to be used, then specialized paging algorithms, such as early page-in to unused I/O system memory, can improve actual output performance.

2.5 Summary

In this chapter, two important measures of parallel file system performance were identified: the application I/O rate and the actual I/O rate. The application I/O rate, the rate at which data is transferred between the application's memory and the operating system, determines the time after an I/O system call during which the application must leave its I/O memory undisturbed. The actual I/O rate, the rate at which the data is transferred between the application's memory and the disk, determines the time that must elapse after an I/O system call before the application receives the data (during input) or before another application can access the data from the file (during output).

Three successively more detailed models of parallel file systems were developed, and the I/O performance of a simple I/O intensive application was described for each model. These models were used to explain the effects of application I/O patterns,
buffering, and I/O system virtual memory on I/O performance. Then the performance patterns predicted by these models were related to the results of experiments performed on two commercial parallel I/O systems (on the Intel Paragon and the Thinking Machines CM-5). Several I/O optimizations identified during the course of this experimentation were described.

This chapter described techniques that parallel application and library developers can use to obtain high I/O performance from parallel file systems. The following three chapters describe three abstractions that support the special I/O requirements of object-oriented parallel programs; these abstractions can be implemented on parallel file systems. Such implementations should incorporate the I/O optimizations identified in this chapter to increase I/O performance.
Chapter 3: Parallel Filestreams

Parallel file systems, described in the previous chapter, effectively support parallel I/O in parallel programs that have relatively simple distributed data structures, such as distributed arrays of floating point numbers. However, *object-oriented* parallel programs have special data management requirements that are not met by parallel file systems.

Object-parallel languages such as pC++ [30] support distributed arrays of arbitrary objects. Complex objects (such as tree-structured objects) often have very fragmented memory usage patterns; parallel file systems only support I/O on blocks of contiguous bytes. So before these programs can perform I/O on these distributed arrays using parallel file systems, some mechanism needs to be provided for aggregating or *flattening* this data into contiguous blocks of memory on each node before output, and for *unflattening* the data during input. So, buffering is required at the application level, along with a class-specific method for programmers to specify how object data is to be flattened and unflattened.

A further complication that arises in object-parallel programs is the fact that distributed arrays of objects may be redistributed between output and input. For many I/O tasks it is important that data written from a given object be read back into the same object, even if the array distribution or number of processors changes between output and input. So some mechanism needs to be provided to keep track of the meta-data describing the source of each data item in a file.
In the absence of programming system support for the above functionality, object-parallel application developers must hard-code this support into each program they write. A better approach is to factor this code out of applications and into language-specific class libraries within object-parallel programming systems. Such libraries can implement new I/O abstractions supporting the I/O functionality object-oriented parallel programs require. In addition, such libraries can function as I/O middleware, improving application portability by insulating applications from the diverse programmer interfaces of underlying parallel I/O systems on different parallel machines, and improving I/O performance by incorporating machine-specific I/O optimizations.

This chapter and the two chapters following describe designs and implementations of three new abstractions supporting I/O in object-oriented parallel programs. This chapter introduces parallel filestreams, an extension of ordinary C++ filestreams supporting node-buffered parallel I/O with flattening and unflattening in SPMD parallel C++ programs. Object-parallel filestreams, introduced in Chapter 4, extend parallel filestreams to support the element buffering required for file-format-independent I/O on distributed arrays of objects in languages such as pC++. Distributed array streams, described in Chapter 5, extend object-parallel filestreams to support a simplified programmer interface as well as automatic bookkeeping of array distribution meta-data within special-format files.

3.1 Goals

As mentioned in the previous section, object-oriented parallel programs require mechanisms for flattening, unflattening, and buffering object data at a level above parallel file systems. In C++, these facilities are provided by the filestream (fstream) abstraction, which is part of the standard C++ iostream library [6]. However, the
single-buffered design of C++ filestreams make them incompatible with existing parallel
file systems; the fact that parallel I/O systems support multiple sources and sinks of data
(i.e. multiple I/O buffers) spread across multiple nodes is precisely what allows them to
support I/O parallelism. The goal of the work described in this chapter was to design
and implement an extension of the filestream abstraction that would support parallel I/O
in SPMD parallel C++ programs, and that could be implemented on existing parallel file
systems.

3.2 Design

The parallel filestream (pfstream) is the abstraction that was designed to meet the above
goals. Figure 3.1 depicts the architecture of this abstraction: a parallel filestream
provides a single buffer on each compute node of a distributed memory parallel machine.

![Figure 3.1: Parallel Filestream Architecture](image.png)

This diagram shows the architecture of the parallel filestream abstraction, along
with a set of objects on four nodes of a distributed memory parallel machine.
### Figure 3.2: Comparing Ordinary and Parallel Filestreams

<table>
<thead>
<tr>
<th></th>
<th>Ordinary Filestreams</th>
<th>Parallel Filestreams</th>
</tr>
</thead>
<tbody>
<tr>
<td>Associated</td>
<td>Associated with a single thread</td>
<td>Associated with one thread per node</td>
</tr>
<tr>
<td>Thread</td>
<td>Single buffer</td>
<td>One buffer per node</td>
</tr>
<tr>
<td></td>
<td>Attached to a single file</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Ordinary file system</td>
<td>Parallel file system</td>
</tr>
<tr>
<td></td>
<td>Flush/load controlled by the stream</td>
<td>Flush/load controlled by programmer</td>
</tr>
<tr>
<td></td>
<td>Static buffer size</td>
<td>Dynamic buffer size</td>
</tr>
</tbody>
</table>

There are several differences between ordinary C++ filestreams and parallel filestreams; these are summarized in Figure 3.2, and described in more detail in the following paragraphs. Both abstractions support one buffer per associated thread — for filestreams, this means a single buffer, whereas parallel filestreams have a buffer per node. A parallel filestream’s buffers can be flushed and loaded concurrently — this is what allows a parallel filestream to take advantage of the underlying parallel file system.

Since filestreams have one buffer associated with a single file, for filestreams, the timing of buffer flushes and loads has no effect on the contents of the file. During writing, data appears in the file in the order it was written to the buffer, no matter when flushes take place; during reading, data is loaded into the buffer in the order it was written in the file. Thus for filestreams, buffer flushing and loading can be controlled automatically by the filestream implementation itself.

In parallel filestreams as well, during writing, considering any particular node, placement of data in the file from that node depends on the order in which data is written to that node’s buffer; however, since a parallel filestream has multiple buffers associated with a single file (one buffer per node), the placement of data within the file also
depends on the amount of data written to the buffers on the other nodes, as well as on the relative timing of buffer flushes on the nodes. So in order to have control over data placement within the file, users of parallel filestreams require control over the timing of buffer flushes (flushes cannot be performed automatically by the parallel filestream implementation, as they are in filestreams). Furthermore, during reading programmers need control over both the relative timing of buffer loads and over the amount of data to be read into each buffer during each load, since in parallel input both factors have an impact on the manner in which data in the file is distributed to the nodes.

Ordinary C++ filestreams generally have fixed sized buffers. But since parallel filestream buffers are flushed manually by the programmer during output, and since the programmer may not know the size of objects that are to be inserted into the buffers...
before flushing, parallel filestreams should have variably-sized buffers that grow (dynamically and automatically) as data is inserted into them.

Figure 3.3 lists per-node methods that implementors of parallel filestreams may wish to support, using a C++ class interface description style. (Of course the parallel filestream abstraction could be implemented in any language.) The state diagrams in Figures 3.4 and 3.5 show the order in which these methods are intended to be used. It may be useful to refer to these figures as the methods are described in more detail in the following paragraphs. `Flush()` flushes the buffer of the node(s) on which it is called, and `load()` causes the node’s buffer to be loaded with data from the disk. (These will be synchronizing operations when they initiate parallel input or output). The `blocksize` methods set and get the current blocksize, which specifies how much data the calling node should read when `load()` is called. This needs to be specified by the user since on machines like the Intel Paragon, in some parallel I/O modes, different data
is is sent to a given node’s buffer during input depending on the amount of data read by other nodes participating in the parallel read. In addition, methods are listed for:

- Opening and closing the file associated with a stream
- Attaching a stream to the file descriptor of a previously opened file
- Setting the parallel I/O mode, which controls the way in which data is mapped between the buffers and the file, and
- Performing unbuffered parallel I/O directly to the underlying parallel file system

The programmer uses insertion functions and extraction functions to copy data from an object into a buffer and from a buffer into an object (respectively). Insertion and extraction functions should be defined by parallel filestream implementors for all of the

Figure 3.5: Parallel Filestream State Diagram for Input.
fundamental (predefined) types in the language, as well as for untyped blocks of memory. In addition, programmers need a means of specifying their own insertion and extraction functions for new types they create themselves. A programmer-defined insertion function decomposes an insertion of a new type into a series of insertions of simpler types for which insertion functions have already been defined, and likewise for programmer-defined extraction functions.

3.3 Implementation

This section describes an implementation of parallel filestreams that I constructed in the summer of 1994 to support parallel I/O in SPMD C++ programs on the Intel Paragon, the Thinking Machines CM-5 and the SGI Power Challenge. The implementation is in the form of a C++ class library; details of the library’s interface and internal architecture are explained. This information should be useful to those planning to implement I/O abstractions similar to parallel filestreams.

3.3.1 Implementation of Insertion and Extraction Functions

This section describes in detail one way of allowing programmers to define their own insertion and extraction functions for flattening and unflattening programmer-defined types. Insertion and extraction functions are implemented by overloading the operators \(<\texttt{<<}\text{and}\texttt{>>}\), allowing a style of I/O similar to formatted text I/O in iostreams [6]:

\[
\begin{align*}
\text{buffer} \texttt{<< object;} & \quad \text{// copy the object into this node’s buffer} \\
\text{buffer} \texttt{>> object;} & \quad \text{// retrieve the object}
\end{align*}
\]

The implementation defines \(<\texttt{<<}\text{ and }\texttt{>>}\) for all the fundamental C++ types. Programmers can define their own \texttt{>>} and \texttt{<<} operators for new types, as in iostreams. For example, for the following type:
class ParticleList {
    int       numberOfParticles;
    double*   mass;              // dynamically allocated
    Position* position;          // arrays
};

the programmer could define insertion and extraction operators as follows:

pfstream& operator<<(pfstream& s, ParticleList& p) {
    // Insert the numberOfParticles field
    // of p (an integer):
    s << p.numberOfParticles;

    // Insert the mass and position fields,
    // variable-sized arrays
    // of size numberOfParticles:
    s << array(p.mass, p.numberOfParticles);
    s << array(p.position, p.numberOfParticles);

    return s;  // allows chaining insertions (s << a << b << c;)
}

pfstream& operator>>(pfstream& s, ParticleList& p) {
    s >> p.numberOfParticles;
    s >> array(p.mass, p.numberOfParticles);
    s >> array(p.position, p.numberOfParticles);
    return s;
}

Pointers (such as mass and position above) require special treatment. Pointers are
generally used either to reference dynamically allocated arrays or to reference
dynamically allocated objects. For arrays, a macro called array() is defined, that
takes a pointer and an integer specifying the size of the array, and returns an object of
type MemBlock containing that information. Insertion and extraction operators for
MemBlocks are defined by the implementation. Pointers to objects are handled by
(possibly recursively) inserting/extracting the object pointed to (see Figure 3.6).
3.3.2 Internal Architecture

Figure 3.7 shows the implementation’s class inheritance structure. There are actually two families of classes:

- **DynamicStreamBuf** plus the four other "DynamicStream" classes implement insertion operators, extraction operators, and a dynamically allocated buffer for a general purpose stream that can be attached to any source and sink of bytes

- **ParallelFileBuf** plus the four "PfStream" classes add the methods and logic necessary to attach that stream to an underlying parallel file system.
Separating the dynamic stream implementation from the parallel file system "glue" code allows the dynamically buffered stream implementation to be easily reused. The only classes in Figure 3.7 that the programmer needs to be aware of are PfStream, InputPfStream, and OutputPfStream; these classes make up the programmer’s interface to the library. InputPfStream and OutputPfStream are similar to

---

Figure 3.7: Class Inheritance Structure in a C++ Parallel Filestream Implementation
PfStream but are intended to be used for either input or output but not both, so their programmer interface can be simplified somewhat.

The DynamicStreamBuf class includes the following member functions:

```cpp
producer(char *buf, int nbytes);  // stores nbytes at buf
c consumer(char *buf, int nbytes);  // copies nbytes from buf
```

These functions act as sources and sinks of bytes, respectively. A DynamicStreamBuf calls these functions when it needs to load or flush its buffer. Of course, since DynamicStreamBufs are meant to be general purpose, these functions cannot actually be implemented within DynamicStreamBuf itself, so they are "pure virtual" functions, implemented within classes derived from DynamicStreamBuf.

DynamicStream also provides the programmer with the following functions for fine-tuning the behavior of the buffers:

```cpp
// Dynamic buffer fine tuning
void  setBufSize(int s);
int   bufSize();
int   bytesInBufferNow();
void  setBufAdd(int a);
int   bufAdd();
void  setBufFactor(float f);
float bufFactor();
```

These allow the programmer to set the size of the buffer and to control how fast the buffer should grow when its capacity is exceeded. BufFactor is a factor by which the size should be multiplied, and BufAdd is a fixed number of bytes that should be added to the size after the multiplication. BufSize is the current maximum size of the buffer, and bytesInBufferNow() returns the number of bytes that are actually in the buffer.

**Middleware Functionality**

In addition to supporting the parallel filestream abstraction, this library functions as parallel I/O middleware, encapsulating machine- and operating system-specific code and supporting a uniform abstraction for parallel I/O across disparate underlying machines and file systems. The parallel I/O modes supported by this implementation are nearly
identical to the those supported by the Paragon’s and CM-5’s parallel file systems, so
implementation was straightforward on these machines, and was slightly more involved
on machines such as the SGI Power Challenge, which lacks a parallel file systems. The
implementation encapsulates machine-specific code within the `ParallelFileBuf`
methods

```c
void parWriteImpl(int fd, char *buf, unsigned nbytes);
void parReadImpl(int fd, char *buf, unsigned nbytes);
```

which are responsible for supporting Paragon-like parallel write and read semantics
based on the current I/O mode, whether or not the underlying file system supports
parallel I/O.

### 3.3.3 Lessons Learned

A lesson learned during the course of this implementation is that the parallel I/O model
is fundamentally incompatible with the stream I/O model employed in the C++ streams
library [6]. A C++ stream is a single source or sink of bytes, but parallel I/O requires
multiple sources and sinks. So while a syntax similar to the C++ iostreams syntax has
been used for the parallel filestreams interface, the semantics of the two interfaces are
somewhat different. An initial intention was to reuse portions of the C++ filestreams
library in this implementation, but the differences in the two I/O models made this
impractical.

### 3.4 Summary

This chapter introduced issues involved in the use of parallel file systems for I/O in
object-oriented parallel programs, and described the design and implementation of
`parallel filestreams`, an abstraction supporting node-buffered parallel I/O in
SPMD-parallel C++ programs. The following chapter extends the parallel filestream
model to support element-buffering, required for I/O in object-parallel (pC++) programs.
Chapter 4: Object-Parallel Filestreams

This chapter describes the *object-parallel filestream* abstraction, which extends the abstraction described in the previous chapter to support parallel I/O in *object-parallel* programs. (*Object-parallelism* is a special type of object-oriented parallelism supported in the language pC++.) First the special I/O requirements of object-parallel programs are identified, then the design and an implementation of object-parallel filestreams is described.

### 4.1 Goals

In SPMD (Single Program Multiple Data) style parallelism, several copies of a program run concurrently on the nodes of a parallel machine, one copy per node, and each copy gets a single thread of control. The programmer may wish to perform I/O operations from any of these threads. To support buffered I/O in these programs, the parallel filestream abstraction (described in the previous section) associates one buffer with each thread — and there is one thread per node, so this means one buffer per node as well. It is important for there to be no more than one thread associated with a given buffer, since this way the threads do not interfere with each other as they read from and write to their respective buffers. And it is useful to associate exactly one buffer with each node, since
this maps well to current parallel file systems, which expect exactly one block of data to be contributed from (sent to) each node, during parallel output (input).

In pC++ [30] style object-parallelism (see "The pC++ Language" on Page 133) there is a single main thread of control from which parallel operations can be performed on collections (distributed arrays of element objects) by invoking a member function of the element objects on the collection itself — initiating one thread per object to execute that function. As in SPMD programs, programmers may wish to perform I/O operations from any of these element threads. However, unlike SPMD programs, if there are more elements than processors, there may be several threads per node in an object-parallel program. This means that parallel filestreams cannot easily be used for I/O in object-parallel programs, since they have only one buffer per node, and as explained above, it is important for there to be no more than one thread associated with a given I/O buffer. The goal of the work described in this chapter was to develop an abstraction that extends the functionality of parallel filestreams to support parallel I/O in object-parallel programs.
This section describes **object-parallel filestreams** (*OpfStreams*), abstractions designed to support parallel I/O on objects in object-parallel programs. Figure 4.1 depicts the architecture of object-parallel filestreams. An object-parallel filestream is similar to a parallel filestream (*pfstream*), but has not only one node buffer on each node (as in parallel filestreams), but also an **element buffer** for each element of the collection on which I/O is to be performed. So object-parallel filestreams support a two-level buffering system: during output, data is first inserted into the element buffers by the element threads, then the element buffers are flushed into the node buffers. Each node buffer accumulates data from its local element buffers; when the node buffers are flushed, parallel output to the file takes place as in parallel filestreams. During input, data is loaded from the file into the node buffers, then again loaded from each node.
buffer into its local element buffers, from which it can be extracted into data structures in each element. (An implementation note: loading a group of element buffers from a local node buffer can be implemented by simply positioning the element buffers within the node buffer’s memory — so during input no data movement between the node and element buffers is actually required.) This two level buffering strategy allows an object-parallel filestream to meet the dual requirements of having both one thread per buffer (so the threads can insert/extract data from their local element buffers in parallel, without synchronizing) and having one buffer per node (this is required by current parallel file systems, which expect exactly one buffer per node for parallel reads and writes).

If programmers are to have control over the order in which data appears in an object-parallel filestream file during output, and if they are to have control over the element into which a given datum in the file is loaded during input, then they must have explicit control over both the timing of buffer flushes and loads and the size of loads, for both the element buffers and the node buffers. The explanation for these requirements

<table>
<thead>
<tr>
<th>Parallel Filestreams</th>
<th>Object-Parallel Filestreams</th>
</tr>
</thead>
<tbody>
<tr>
<td>Controlled by one thread per node</td>
<td>Controlled by one main thread that can invoke one thread per distributed array element</td>
</tr>
<tr>
<td>One buffer per node</td>
<td>One buffer per node, plus one buffer per distributed array element</td>
</tr>
<tr>
<td>Attached to a single file residing on an underlying parallel file system</td>
<td></td>
</tr>
<tr>
<td>Flush/load controlled by the user of the stream</td>
<td></td>
</tr>
<tr>
<td>Dynamic buffer size</td>
<td></td>
</tr>
</tbody>
</table>
follows the same reasoning as the explanation for the similar requirements for parallel filestreams (Page 45). Figure 4.2 compares parallel and object-parallel filestreams.

Figure 4.3 lists methods that implementors of object-parallel filestreams may wish to support, using a pC++ style class and collection interface listing. The state diagrams in Figure 4.4 and 4.5 show the order in which the methods are intended to be used. These methods are described in more detail in the following paragraphs.

At the node level, object-parallel filestreams need to support exactly the same methods that parallel filestreams support for opening and closing files and for loading and flushing the node buffers. Programmers should be able to initiate SPMD-style parallel operations on the node buffers from the main thread of object-parallel programs.
(In Figure 4.3 this is indicated by the inclusion of a data member \texttt{nodeBuffers} of type \texttt{PfStream} in the public (SPMD) section of the \texttt{OpfStream} collection.)

At the element level, object-parallel filestreams need to support methods for loading the element buffers from (and flushing them to) the node buffers and for inserting (extracting) data into (out of) the elements of collections on which I/O is to be performed. Since the element buffers on a given node share a node buffer, in order to insure deterministic behavior during object-parallel flushes and loads, versions of element flush and load need to be provided that serialize the element buffers’ access to the local node buffer during object-parallel flushes and loads. The \texttt{flushAll} and \texttt{loadAll} methods support this functionality, allowing the elements access to their node buffer in order of their element index.

Insertion and extraction should be controllable from both the main thread and from the element threads; see "Implementation of Insertion and Extraction Functions" on Page 49 for an example implementation.
Figure 4.4: Object-Parallel Filestream State Diagram for Output

(Assuming N nodes and E element buffers per node). A state is associated with each buffer (both node and element buffers). Element buffer states are listed below the corresponding node buffer’s states. All buffers start in the empty state.
Figure 4.5: Object-Parallel Filestream State Diagram for Input.
(See the caption of Figure 4.4 on Page 62 for further explanation.)
4.3 Implementation

This section describes an implementation of object-parallel filestreams that I constructed in the summer of 1994 to support parallel I/O in pC++ programs. The implementation is a pC++ class library that builds upon and reuses a large portion of the parallel filestreams implementation described in the previous chapter. In this section details of this library’s interface and internal architecture are explained.

4.3.1 Interface

Parallel Operations on Node and Element Buffers

When working with object-parallel filestreams (OpfStreams), programmers need mechanisms for specifying parallel operations (such as load and flush) on the stream’s node and element buffers. This section describes a straightforward way that control of these operations can be supported from the main thread in pC++ implementations of object-parallel filestreams.

To support parallel operations on the node buffers, the object-parallel filestream can be implemented as a pC++ collection of element buffers. One parallel filestream (PfStream) object per node can be incorporated into the object-parallel filestream by placing it in the "public" part of the OpfStream collection.

```cpp
Collection OpfStream {
    public:
        PfStream n;
    ...
};
```

Then using pC++ syntax, the programmer can initiate SPMD-parallel operations (such as flush) on the node (parallel filestream) buffers using a single statement in the main thread:
OpfStream<ElementStream> s;
...
 s.n.flush();

To support parallel operations on the element buffers using a similar syntax, a 
slightly different implementation technique is required. The element buffers can be 
implemented by a class ElementStream. A virtual reference to an 
ElementStream can be included within the MethodOfElement section of the 
OpfStream collection:

```
Collection OpfStream {
 ...
 MethodOfElement:
  virtual ElementStream &e;
 ...
};
```

indicating to the compiler that a similar (but non-virtual) reference to an 
ElementStream exists in the ElementStream class:

```
class ElementStream {
 public:
  ElementStream &e;
 ...
};
```

The ElementStream constructor should set this reference to refer to *this (the 
constructed ElementStream itself). Then programmers can specify object-parallel 
operations on the element buffers with a single line of code in the main thread:

```
OpfStream<ElementStream> s;
...
 s.e.flushAll();
```

Here the programmer specifies that all element buffers should be flushed to their 
respective node buffers.

**Insertion and Extraction Functions for Collections**

This section describes pC++ implementation techniques for object-parallel filestreams 
that allow programmers to insert (or extract) elements of an entire collection to (or from)
an object-parallel filestream’s element buffers with a single statement. (The following examples are framed in terms of insertion; extraction can be implemented similarly.)

pC++ allows parallel operations on collections to be expressed using a data-parallel style syntax. If s and c are the collections:

```cpp
OpfStream<ElementStream> s;
MyCollection<MyElement> c;
```

and if there is an insertion operator `<<` defined on the `element` types of s and c:

```cpp
ElementStream& operator<<(ElementStream& es, MyElement& me);
```

then the meaning of the following statement in the main thread

```cpp
s << c;
```

is that an element thread should be created for each corresponding pair of elements from s and c, and each of these threads should carry out the application of

```cpp
ElementStream& operator<<(ElementStream& es, MyElement& me);
```

on its pair of elements. But this can be an ordinary programmer-defined insertion operator that inserts the data associated with an object of type `MyElement` into the element buffer of its associated `ElementStream`. These operators are used and defined the same way as insertion operators for parallel filestreams, described in "Implementation of Insertion and Extraction Functions" on Page 49.

The following strategy can be used to save memory during output. As data is inserted into the element buffers, in cases where the amount of data to be inserted is larger than the size of the representation of a pointer, a pointer to the data can be inserted into the element buffer instead of the actual data (since the data doesn’t actually have to be aggregated until it is sent to the node buffers). The size of the data pointed to must be inserted also, and the size information must be interpreted intelligently when data is being copied from the element buffers into the node buffers.
Programmers can insert (or extract) a single field of the elements of an entire collection with a single statement in the main thread as well. Assume the following collections are defined:

```c++
OpfStream<ElementStream> s;
MyCollection<MyElement> c;
```

and assume MyElement has an integer member intMember. Then the intMember field of each of the elements of c can be inserted into the element buffers of s using the following syntax in the main thread:

```
s << c.intMember;
```

This works because in pC++, c.intMember is a collection of type MyCollection<int> having the same distribution and alignment as c, and whose elements are the integer members of the elements of c. So this is just the insertion of an entire (simpler) collection of integers into the OpfStream.

Now assume d is a second collection aligned with c and containing an integer field intMemberOfD. If the programmer invokes

```
s << c.intMember;
s << d.intMemberOfD;
```

this will cause the corresponding intMember and intMemberOfD fields of the elements of c and d to be placed contiguously in the element buffers of the OpfStream (so they will be written contiguously in the file), even if they are scattered in memory. This functionality is called interleaving.
4.3.2 Internal Architecture

Figure 4.6 shows the class inheritance structure of the implementation. Much of the implementation of parallel filestreams (described in section 3.3 on Page 49) is reused; only three new classes were added. The `OpfStream` collection class implements the interface to the node buffers, which can conveniently be implemented by containing a single parallel filestream within the object-parallel filestream. (Note: This works in
pC++ 1.0 because in this version of pC++, non-MethodOfElement data members of collections are always duplicated one per node.) An ElementStreamBuf class (derived from DynamicStreamBuf) implements a single element buffer. Recall that the producer() and consumer() functions of ParallelFileBuf read from and wrote to an underlying parallel file system (see Page 53); in ElementStreamBuf, these functions read from and write to their associated node buffer in the OpfStream collection. To facilitate coding the data movement between the element and node buffers, each ElementStreamBuf maintains a pointer to its associated node buffer. An instance of ElementStreamBuf is contained in the ElementStream class, whose main purpose is to inherit insertion and extraction operators from DynamicStream. ElementStream contains facilities for dynamic buffer fine-tuning similar to those in the parallel filestream implementation (see Page 53).

To instantiate a complete OpfStream, the programmer declares a collection of type OpfStream<ElementStream> (an OpfStream collection having elements of type ElementStream), and gives this collection the same distribution and alignment as the collection(s) on which I/O is to be performed. This ensures there is a one-to-one correspondence between the ElementStreams and the elements of the collection on which I/O is to be performed.

### 4.4 Summary

This chapter discussed the design and implementation of the object-parallel filestream abstraction, an extension to parallel filestreams that can be used to support parallel I/O in object-parallel programming systems such as pC++. The following chapter describes several ways in which object-parallel filestreams can be extended to make their use easier for programmers.
Chapter 5: Distributed Array Streams

This chapter describes the design and an implementation of the distributed array stream, an abstraction that extends the object-parallel filesteam to support a simplified programmer interface as well as automatic redistribution of element data (which is necessary when the element distribution or number of processors changes between output and input).

5.1 Goals

The use of parallel filestreams (Chapter 3) and object-parallel filestreams (Chapter 4) is somewhat more complex than the use of ordinary C++ filestreams. Ordinary filestreams behave like simple first-in first-out queues that reside on persistent storage. Parallel and object-parallel filestreams behave somewhat like queues as data is being inserted and extracted from a given element or node buffer; however, in addition to being concerned with the order in which data are inserted and extracted from each buffer, the programmer must also control the flushing and loading of these buffers. Could this complexity be reduced by an improved design? Probably not, given the requirements of parallel I/O:

1) **Buffering is required for efficient I/O:** If I/O is to be efficient, I/O hardware requires that data be aggregated into relatively large chunks before output is performed. Likewise, I/O hardware returns data to memory in large chunks, from
which it must be de-aggregated. So I/O abstractions must support buffering for aggregation and de-aggregation.

2) **Multiple buffers are required for parallel I/O:** In order to support simultaneous I/O involving multiple nodes of a parallel machine, there must be at least one buffer for aggregation/de-aggregation on each node participating in the I/O. In the case of object-parallel filestreams, if multiple elements on a single node are to be able to aggregate/de-aggregate data in parallel (without synchronizing), then object-parallel filestreams must support an additional buffer per element.

3) **If the parallel I/O is to store or access data in a (single) file, and if the programmer is to control the file’s format, then the mapping between the buffers and the file cannot be controlled automatically:**; there must be some mechanism by which the programmer specifies this mapping. Giving the programmer control over buffer flushes and loads, as the file is written or read from beginning to end, is one way for the programmer to specify this mapping; this is the approach that has been taken in the design of parallel and object-parallel filestreams.

On the other hand, if the above requirements are relaxed, then it may be possible to simplify the coding of parallel I/O in application programs:

- **If data can be stored in multiple files,** then each file can be associated with a single thread (i.e. a single processor), so each processor can use a simple, separate abstraction (such as an ordinary C++ filestream) for I/O. This is often the first I/O approach supported by developers of new parallel systems, since it is relatively simple to implement. This is a workable approach for I/O supporting checkpoint/restart tasks, but there are several potential drawbacks to using this approach for initial input or final output. One problem is that SPMD programs
running on a large number of nodes will generate and consume a large number of files, and the work involved in managing all of these files may not be trivial. A more serious problem is the fact that parallel programs often need to be able to be run with widely varying degrees of parallelism, which often means using a widely varying number of processors. So if there is one file per processor then the number of input files required and output files generated will vary from run to run, and the user must redistribute the input data into separate input files for each degree of parallelism desired. Furthermore, most visualization tools and other programs used to analyze output data expect the data to reside in a single file, so multiple output files will need to be consolidated before analysis can take place.

• **If the requirement that data be stored in files is dropped**, then alternate persistent storage mechanisms such as object-oriented database management systems could be used to support I/O tasks. This approach is explored further in Chapter 6, "Persistent Collections".

• **If the requirement that the programmer have complete control over the file format is relaxed** then some aspects of buffer flushing and loading can be automated, making things easier for application developers, while maintaining the benefits of storing data in a single file.

The goal of the work described in this section was to simplify the programmer’s interface for parallel I/O in object-parallel languages like pC++, taking the latter approach above, i.e. relaxing the requirement that the programmer have complete control over the file format.
5.2 Design

The distributed array stream (d/stream) is an abstraction for parallel I/O in object-parallel languages such as pC++. Like object-parallel filestreams, d/streams support parallel I/O between distributed arrays of objects and a single file; however, d/streams require a that a fixed (d/stream specific) file format be used. This allows d/stream implementations to automatically record information within the file about the size and distribution of the objects stored in the file, as the file is being written. Then when reading, d/stream implementations can automatically route data to the correct distributed array elements, even if these elements appear on different nodes than when the file was written (this may occur if the distribution of the array or the number of nodes has changed since the array was written). In addition, since information about the size of objects in d/stream files is stored within the files themselves, d/streams do not require programmers to specify the amount of data to load into each buffer during input (as is required in parallel and object-parallel filestreams; see "Goals" on Page 43 for further explanation). Programmers can simply invoke "load" and the d/stream implementation determines the amount of data that should be loaded into each buffer.

Figure 5.1 illustrates the architecture of the d/stream abstraction, which is described further below. Like object-parallel filestreams, d/streams have one buffer per distributed
array element. Unlike object-parallel filestreams, loads and stores on these element buffers are synchronizing operations, and the mapping between the element buffers and the file is fixed (the programmer cannot vary the mapping by varying the timing of loads and stores of the element buffers). Though d/stream implementations may make use of node buffers, node buffering is transparent to programmers using d/streams. So node buffering is not part of the architecture of the d/streams abstraction, though it may be a part of a given d/streams implementation architecture. Figure 5.2 summarizes these differences between object-parallel filestreams and d/streams.

Figure 4.3 summarizes the methods that implementors of d/streams may wish to support, using a pC++ style interface listing; these methods are described in more detail below. Insertion and extraction of distributed array data are controllable both from element threads and from the main thread (for insertion of entire distributed arrays), as in object-parallel filestreams (see "Insertion and Extraction Functions for Collections" on

<table>
<thead>
<tr>
<th>Object-Parallel Filestreams</th>
<th>D/streams</th>
</tr>
</thead>
<tbody>
<tr>
<td>Can read and write any file format</td>
<td>Must read and write a special file format</td>
</tr>
<tr>
<td>One buffer per node, plus one buffer per distributed array element</td>
<td>One buffer per distributed array element</td>
</tr>
<tr>
<td>Buffer flushing/loading can be individually controlled</td>
<td>Buffer flushing/loading methods act on all buffers as a group</td>
</tr>
<tr>
<td>User must manage mapping from writing elements to file to reading elements using fine control of buffer flushing and loading</td>
<td>Stream implementations manage the mapping automatically; users have only coarse control over buffer flushing/loading</td>
</tr>
<tr>
<td>Controlled by one main thread that can invoke one thread per distributed array element</td>
<td></td>
</tr>
<tr>
<td>Attached to a single file residing on an underlying parallel file system</td>
<td></td>
</tr>
<tr>
<td>Dynamic buffer size</td>
<td></td>
</tr>
</tbody>
</table>

**Figure 5.2: Comparing Object-Parallel Filestreams and D/streams**
The class \texttt{ElementDStream} is intended to be the element type of collections of type \texttt{DStream}. Both \texttt{load} and \texttt{unsortedLoad} transfer a block of data (written by a corresponding \texttt{flush}) from a file into an input d/stream’s buffer. Implementations of \texttt{unsortedLoad} may have better input performance when reading un-ordered distributed arrays of objects.

Page 65 for an example of how this can be implemented). The d/streams buffer management methods (\texttt{flush()}, \texttt{load()}, and \texttt{unsortedLoad()}) are callable from the main thread only. \texttt{Flush()} flushes all of the data in all of the element buffers to the file using a file format that should be clearly defined by implementations. \texttt{Load()} and \texttt{unsortedLoad()} read the data written by a corresponding \texttt{flush()} back into
the element buffers. So the programmer needs to be aware of element buffering, but not node buffering.

Note that unlike parallel and object-parallel filestreams, d-streams do not require the programmer to specify the amount of data that should be loaded, since this is recorded in the file by flush(). Furthermore, the d/stream load and flush methods are much more intelligent than the corresponding methods in parallel and object-parallel filestreams, in the following respect. In parallel and object-parallel filestreams, the mapping of data from threads performing output to threads performing input is affected by the pattern of insertions and extractions, the pattern of flushes and loads, the distributions of the data structures from which output and input are performed, and the number of nodes that participate in output and input — so the programmer must take all these factors into account when using these abstractions for I/O. When using d-streams, the only variable the programmer needs to be concerned with is the pattern of insertions and extractions; this pattern must be the same for output and input, i.e. the same types must be inserted/extracted from the element threads in the same order during output and input.

The flush and load methods exist only to allow programmers to control interleaving (see Page 67) and buffer memory usage. Implementations in which these are not important issues may choose to fully automate loading and flushing. For a given flush/load pair, d/stream implementations should guarantee that load returns blocks of bytes to exactly the same element buffers from which they were flushed, even if the array distribution or number of nodes (and thus the number of elements per node) changes between output and input. For unsortedLoad, implementations must guarantee only that each of these blocks was written from some (single) element, though not necessarily the one with the same element index. Thus unsortedLoad can be faster than load in some implementations. UnsortedLoad is intended to be used to
read array data in which the element indices perform no important role in the computation.

The state diagrams in Figure 5.4 show the order in which the d/stream methods are intended to be called; this is similar to the usage of ordinary C++ iostreams. Several constraints on the use of the methods cannot be indicated in the state diagrams so they are discussed here. Data must be read in the same order it was written. Specifically:

- When a d/stream file is read by an input d/stream, every load or unsortedLoad must correspond to a flush that occurred when the file was written, and every extract must have a corresponding insert.
- Each extracted object or array must have the same size and type as the corresponding array that was inserted.

5.3 Implementation

This section describes a pC++ implementation of distributed array streams that I constructed in the summer of 1994. Like implementations of the abstractions described
earlier, this implementation is a pC++ class library built on a more primitive library (the object-parallel filestreams library described in the previous chapter).

5.3.1 Interface

This section illustrates how the d/streams interface was implemented in pC++ by showing how the library described in this section would be used to code checkpoint/restart tasks in an example application. Assume the application simulates the behavior of particles in 3D space. Information about the particles can be stored in a distributed array of particle lists, where each list keeps track of the particles in a 3D slab of space. Figure 5.5 shows the declarations that would be used to declare this data structure (called particleArray) in pC++. Using d/streams to write a procedure to checkpoint the entire particleArray from the main thread is fairly straightforward:

```cpp
#include "particleArray.h"
#include "pc++streams.h"

void saveParticleArray() {
    DStream stream(&d, &a, "myCheckpointFileOne");
    stream << particleArray;
    stream.flush();
}
```

It is also possible to checkpoint a single field of the elements of the particleArray:

```cpp
#include "particleArray.h"
#include "pc++streams.h"

void saveNumberOfParticles() {
    DStream stream(&d, &a, "myCheckpointFileTwo");
    stream << particleArray.numberOfParticles;
    stream.flush();
}
```
class Position {
    double x, y, z;
};

class ParticleList {
    int numberOfParticles;
    double * mass;  // variable sized
    Position* position;  // arrays
};

Processors P;
Align a(12,"[ALIGN(collection[i], template[i])]"); -
Distribution d(12, &P, CYCLIC); - - - - -

Collection DistributedArray {
    updateParticles();
} -

DistributedArray<ParticleList> particleArray(&d, &a);

<table>
<thead>
<tr>
<th>Node 1</th>
<th>Node 2</th>
<th>Node 3</th>
<th>Node 4</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>5</td>
<td>6</td>
<td>7</td>
<td>8</td>
</tr>
<tr>
<td>9</td>
<td>10</td>
<td>11</td>
<td>12</td>
</tr>
</tbody>
</table>

Figure 5.5: Declaring a Distributed Array of Lists of Particles in pC++
(These declarations are referred to as particleArray.h subsequently).
Functions to restore the checkpointed data can be written similarly:

```cpp
#include "particleArray.h"
#include "pc++streams.h"

void loadParticleArray() {
    DStream stream(&d, &a, "myFileOne");
    stream.load();
    stream >> particleArray;
}

void loadNumberOfParticles() {
    DStream stream(&d, &a, "myFileTwo");
    stream.load();
    stream >> particleArray.numberOfParticles;
}
```

In the procedures `saveParticleArray()` and `loadParticleArray()` above, the statements

```cpp
stream << particleArray; // and
stream >> particleArray;
```

apply the operators `<<` and `>>` to corresponding pairs of elements of the collections `stream` and `particleArray`, i.e. they invoke the operators

```cpp
DElementStream& operator<<(DElementStream& eltStr,
                         ParticleList& p);
DElementStream& operator>>(DElementStream& eltStr,
                         ParticleList& p);
```

on each pair of elements (see Page 65 for more details). The programmer needs to define these operators to tell the d/streams implementation how `ParticleLists` should be inserted into and extracted from element streams.

```cpp
class ParticleList {
    public:
        int  numberOfParticles;
        double* mass        // dynamically allocated,
                             Position* position; // variable sized arrays
};
```
The programmer can accomplish this by making the following declarations in particleList.h:

```cpp
declareStreamInserter(ParticleList &p) {
    eltStr << p.numberOfParticles;
    eltStr << array(p.mass, p.numberOfParticles);
    eltStr << array(p.position, p.numberOfParticles);
}

declareStreamExtractor(ParticleList &p) {
    eltStr >> p.numberOfParticles;
    eltStr >> array(p.mass, p.numberOfParticles);
    eltStr >> array(p.position, p.numberOfParticles);
}
```

In this d/streams implementation, declareStreamInserter() and declareStreamExtractor() are macros defined in pC++streams.h; the compiler transforms these into the appropriate declarations:

```cpp
DElementStream& operator<<(DElementStream& eltStr,
    ParticleList& p)
```

and

```cpp
DElementStream& operator>>(DElementStream& eltStr,
    ParticleList& p)
```

simplifying the required syntax for the programmer. The array() macro is used to insert/extract dynamically allocated array data; see Page 50 for further details.

Figure 5.6 summarizes the actions that occur when saveParticleArray() is called. The statement

```cpp
stream << particleArray;
```

in the main thread invokes the parallel application of

```cpp
DElementStream& operator<<(DElementStream& eltStr,
    ParticleList& p)
```

in the element threads, on corresponding pairs of elements from stream and particleArray. This operator is defined by the programmer in particleArray.h, using the macro declareStreamInserter(ParticleList &p). The insertion operators (and the array() macro) used there are defined by the d/streams implementation (in pC++streams.h). If the ParticleList class had contained data
members of other programmer-defined types, then programmer-defined insertion functions for those types (defined by other macros similar to `declareStreamInserter(ParticleList &p)`) would be called within `declareStreamInserter(ParticleList &p)`.
Figure 5.7: Class Inheritance Structure in a pC++ D/streams Implementation

(Classes with gray borders are described in more detail in figures on pages 52 and 68.)

5.3.2 Internal Architecture

Figure 5.7 shows the class inheritance structure of my pC++ implementation of d/streams. Note that since this implementation is built directly on the implementation of object-parallel filestreams described in the previous chapter, it was possible to reuse both the object-parallel filestreams implementation (Section 4.3, Page 64) and the parallel
filestreams implementation (Section 3.3, Page 49). There are two new classes. The
DStream collection class is derived from the object-parallel filestream collection, and
contains the main-thread part of the implementations of the flush and load methods.
DElementStream, derived from ElementStream, contains the element-thread part
of the implementation. So to instantiate a complete d/stream, the programmer declares a
collection of type DStream<DElementStream> (a DStream collection having
elements of type DElementStream), and as with the object-parallel filestream
implementation, the programmer must give this collection the same distribution and
alignment as the collection(s) on which I/O is to be performed, so there is a one-to-one
correspondence between element buffers of the stream and elements of the
source/destination collection.

Figure 5.8: A Possible File Format for D/streams Implementations
Figure 5.8 depicts a file format appropriate for d/streams; this is described in greater
detail below. Assume the programmer declares

    DStream<DElementStream> s(...);

in the main thread. The flush method is invoked from the main by simply calling

    s.flush();

which causes the stream implementation to perform the following steps:

1) **Writing distribution and size information**: Information about the distribution of the collection \( s \) (and thus the distribution of all collections that could have inserted data into \( s \)), and about the size of the data to be output from each element, needs to be written to the file prior to the actual data, so that the implementation of load will know how much data is needs to be read and where the data belongs. The implementation writes this data from all the nodes concurrently, using a parallel write operation. For collections having a large number of elements (and thus a large amount of distribution and size information to write), this is a good strategy. For smaller collections, due to the latency concerns involved in writing small chunks of data, it may be more efficient to communicate this information to a single node (node zero on the Paragon and CM-5 would be an appropriate choice). Then instead of writing this information in a separate parallel output step, it can be placed at the head of that node’s buffer so that it can be written with the actual data in step 2 (this data needs to appear before the actual data in the file; thus node zero should be the choice on the Paragon and CM-5).

2) **Writing the actual data**: Next the data in the element buffers (or referenced by pointers in the element buffers) is aggregated into the node buffers, then all the node buffers are written to the file in a single parallel write operation, using the parallel I/O primitives of the underlying parallel file system. On the Intel Paragon
and Thinking Machines CM-5, the implementation uses parallel I/O primitives that transfer a block of data from each compute node to the file system simultaneously, and that write those blocks to the file in order of node number.

The programmer invokes the d/stream load method calling s.load(), and the unsortedLoad method by calling s.unsortedLoad(). When either method is invoked, the implementation first reads the distribution and size information preceding the actual data, using one parallel read operation, then reads the actual data into the node buffers using a second parallel read. (Note that no information about the distribution or size of the data to be read needs to be passed to the library by the programmer when reading, since that information is stored directly in the file, preceding the data itself.) If unsortedLoad was invoked, then at this point the implementation sets up each element buffer so that it falls within the appropriate region in its local node buffer, and the data is ready to be extracted. If load was invoked, then if the number of nodes or the distribution of the array is different than when the array was written, the blocks of data that were read into the node buffers may need to be sent to the actual owner nodes before the element buffers can be set up.

Implementing flush and unsortedLoad using the facilities provided by the underlying object-parallel filestreams implementation is a fairly straightforward task. Flush is implemented by the following main-thread procedure:

```c++
void DStream::flush() {
    // set the parallel I/O mode to syncpar
    // meaning all nodes share a single file pointer,
    // and in a parallel flush, a block of data is sent from
    // each node to the file in node number order
    n.setparmode(pfilebuf::syncpar);

    // write the size of each element buffer
    // into its local node buffer
    n_referencedFromElement() << e.bytesInBufferNow();
    // flush the node buffers to the file
    n.flush();
}```
The object-parallel filestream implementation makes it simple to specify parallel stream operations from the main thread; SPMD-parallel operations on all of the node buffers can be specified using \( n\).operation(), and object-parallel operations on all of the element buffers can be specified using \( e\).operation(). Within an object-parallel operation on the element buffers, the function \( n\).referencedFromElement() can be used to refer to the local node buffer. (See Page 64 for more details). The call \( e\).flushAll(): flushes the element buffers to their respective node buffers in the order of their element indices on each node.

**UnsortedLoad** is implemented as follows:

```cpp
void DStream::unsortedLoad() {
    // set the parallel I/O mode to syncpar
    // meaning all nodes share a single file pointer,
    // and in a parallel load, a block of data is sent from
    // the file to each node in node number order
    n.setParmode(pfilebuf::syncpar);

    // on each node, load a single integer from the file for
    // each local element (this is the size of the block of
    // data to be loaded for each local element)
    n.setBlockSize(elementsOnThisNode * sizeof(int));
    n.load();

    // on each node, for each local element, read an integer
    // from the local node buffer and set the element buffer’s
    // block size accordingly
    e.setBlockSizeFromNodeBuffer();

    // on each node, sum the block sizes of the local element
    // buffers so the node’s blocksize can be set to the
    // total amount of data to load into the node
    int nodeBufferSize = 0;
    ResetLocal(this);
    for (int i=FirstLocal(); i>=0; i=Local(i))
        nodeBufferSize += (*this)(i)->e.blocksize();
    n.setBlockSize(nodeBufferSize);
}
```
// for each node buffer, load that node buffer’s (blocksize)
// bytes from the file (in order of node number)
    n.load();
// for each node, for each local element buffer,
// load that element buffer’s blocksize bytes from the
// node buffer, in order of element index
    e.loadAll();
}

(Note: in both flush and unsortedLoad, the movement of the elements’
buffer-size/blocksize data between the elements and node buffers needs to be serialized
in order of element index on each node. No explicit code is needed to perform this
serialization on the current implementation of pC++, since pC++ itself automatically
serializes object-parallel operations on each node in order of element index.)

UnsortedLoad requires one object-parallel helper function in DElementStream
for reading each element’s blocksize from its local node buffer:

    void DElementStream::setBlockSizeFromNodeBuffer() {
        int size;

        // extract this element’s blocksize from the local node buffer
        n_referencedFromElement >> size;
        setblocksize(size);
    }

Note that the above code does not write the element indices (distribution
information) to the file with the sizes of each element buffer, nor does it read the
distribution information in unsortedLoad. UnsortedLoad does not require
distribution information, since it reads element data in whatever order it appears in the
file. However, load does require distribution information. Load can be implemented
as follows. In flush, each element should send two integers to the node buffer before
the actual data is flushed: the element buffer size (as above) and the element’s index. In
load, the element index data should be read along with the element size data; then after
the actual data has been read, each node knows the actual element index corresponding
to each of the data blocks in the node buffer. In addition, each node knows the current
distribution of the receiving array (which has the same distribution as the stream itself).
So each node should communicate the sizes of its element data blocks to the correct
owner nodes. Then each node should allocate a new node buffer of the appropriate size to hold the element data it is about to receive, after which each node can send its element data blocks to the correct owner nodes, where they can be stored in the newly allocated node buffers.

5.4 Summary

For the programmer, coding parallel I/O on distributed arrays is much simpler with distributed array streams than with object-parallel filestreams. The price for this simplicity is the programmer’s loss of control over the file format. Object-parallel filestreams is the more general I/O abstraction, since it supports parallel I/O on files of any format. But just as object-parallel filestreams can simplify the implementation of dstreams, it can also simplify the implementation of application-specific parallel I/O mechanisms having application-specific file formats. So dstreams should be viewed as just one possible simplified special-format parallel I/O abstraction that can be built on object-parallel filestreams.

The next chapter looks at a totally different approach to I/O in object-oriented parallel simulations: persistence.
Chapter 6: Persistent Collections

Normally, elements of pC++ collections are transitory, i.e., their data disappears when the program terminates. In order to preserve transitory data, the programmer must write code to output that data to a file before the program terminates, using either an operating system supported file I/O mechanism or a higher-level abstraction such as distributed-array streams (described in the previous chapter).

This chapter explores an alternative to the file I/O model for supporting data management tasks in object-parallel simulations: the persistent data model. A new object-parallel I/O abstraction is proposed, the *persistent collection*. Elements of a persistent collection can contain persistent data in addition to ordinary transitory data.

![Persistent Collection Architecture](image)

**Figure 6.1: Persistent Collection Architecture**

A persistent collection is a distributed array of objects, where each object has a persistent part and a transient part.
The persistent section of each element is automatically preserved across program executions; no application I/O code is required to save or load this data. A transaction mechanism is supported, allowing programmers to checkpoint persistent data with a single line of code that commits a transaction. In addition, the persistent part of a collection is concurrently accessible by multiple pC++ programs, with no explicit code for communication required. The following sections describe the design and an implementation of the persistent collection abstraction, and discuss an example program using persistent collections for I/O.

### 6.1 Goals

The goal of the work described in this chapter was to investigate how persistent data systems supporting data management in object-oriented parallel languages can be...
Figure 6.3: Persistent Collection State Diagram

Specifies the dynamic behavior of persistent collections. Several constraints not shown in the diagram are described here. There are (at least) two types of locks on persistent data: shared and exclusive. In state 2, the first attempt to read persistent data causes a shared lock to be acquired on that data before it can be read. Acquisition of the shared lock will succeed immediately as long as no other program has an exclusive lock on the same data (in which case the thread will block until the other program releases the exclusive lock). Similarly, in state 2, an attempt to modify persistent data causes an exclusive lock to be acquired on that data before it can be modified (and before the transition to state 3). Acquisition of the exclusive lock will succeed immediately as long as no other program(s) have exclusive or shared locks on the same data (in which case the thread will block until the other program(s) release the locks). Implementations may include mechanisms for dealing with deadlocks, or may leave deadlock avoidance up to the programmer. A persistent collection must be associated with a collection on stable storage before its persistent data can be accessed or modified.

designed, and to investigate how object-oriented database management systems (OODBMS) can be applied in the implementation of these systems.
6.2 Design

Figure 6.1 sketches the architecture of the persistent collection abstraction, which is described further below. A persistent collection is a distributed array of objects, where the data within each object is divided into a transient part and a persistent part, and access to the persistent data is regulated by a transaction mechanism. So to use a persistent collection, the programmer needs mechanisms for identifying the transient and persistent parts of data within the element class and mechanisms for beginning, committing, and aborting transactions on the persistent data. Figure 6.2 (which lists methods that implementors of persistent collections may wish to support, using pC++ notation) shows these mechanisms being inherited by the programmer’s collection from generic PersistentCollection and PersistentElement classes, but other implementation techniques may be possible. Figure 6.3 describes the transaction and locking mechanisms underlying the dynamic behavior of persistent collections.

6.3 Implementation

I implemented persistent collections for the pC++ language in the summer of 1995 as a pC++ library built on a beta release (version 0.9.3) of the Shore [7] object-oriented database management system. The next section describes the pC++ interface to this library, and the section following describes details of the implementation’s internal architecture.

6.3.1 Interface

This section first describes how programmers can define persistent collection elements, then describes how persistent collections themselves can be instantiated and used.
Elements of persistent collections are defined as follows. The programmer defines the data that is to reside in the persistent part of the elements using the interface definition language SDL (Shore Data Language), in an "interface" called PersistentElementData. This definition takes place in a special file PersistentData.sdl, which is transformed (by Shore) into a C++ header file PersistentData.h that the user includes in pC++ source files that access the persistent data. The following is the SDL definition the programmer would use to obtain a single persistent long integer per element (myPersistentLong):

```yaml
module MyElement {

    interface PersistentElementData {
        public:
            attribute long myPersistentLong;
    }

}
```

All SDL interfaces must be defined within an SDL module. Implementors of persistent collections on Shore may wish to use the module name (MyElement) to distinguish between PersistentElementData definitions for different element classes. (So in the above example PersistentElementData would define the persistent part of pC++ collection elements of type MyElement.)

The programmer defines both the transient part of a persistent collection’s elements and the element’s methods in an ordinary pC++ element class (MyElement in this example), deriving this class from the class PersistentElement (which is defined in PersistentElement.h, part of the implementation):

```cpp
#include "PersistentElement.h"

class MyElement : public PersistentElement {
    public:
        long myTransientLong;
        void P_initialize();
        void hello();
};
```
The class PersistentElement contains a data member \( P \) through which the element’s persistent part can be accessed; data in the transient part is accessed normally:

```cpp
void MyElement::hello() {
    printf("Hello world!\n");
    printf("My persistent data is %ld\n", P->myPersistentLong);
    P.update()->myPersistentLong += 1;
    printf("My transient data is %ld\n", myTransientLong);
}
```

\( P->myPersistentLong \) is used for read-only access to \( myPersistentLong \); \( P.update()\rightarrow myPersistentLong \) is used when \( myPersistentLong \) is to be updated.

Persistent collection "classes" are defined exactly like ordinary collections, except that they must be derived from PersistentCollection (which is defined in PersistentCollection.h, part of the implementation):

```cpp
#include "PersistentCollection.h"

Collection MyCollection: public PersistentCollection {
    public:
        MyCollection(Distribution *D, Align *A,
                     char *persistentCollectionName);
        MethodOfElement:
            virtual void hello();
    };

MyCollection::MyCollection(Distribution *D, Align *A,
                            char *persistentCollectionName)
    :PersistentCollection(D, A, persistentCollectionName) {};
```

Note that the constructor \( MyCollection::MyCollection() \) takes a string argument \( persistentCollectionName \), which it must pass on to the constructor of \( PersistentCollection \).

The programmer can instantiate a persistent collection of type \( MyCollection<\text{MyElement}> \) having \( SIZE \) elements distributed in \( BLOCK \) fashion over the processors in the usual way:
void Processor_Main(int argc, char **argv) {
    Processors P;
    Distribution D(SIZE, &P, BLOCK);
    Align A(SIZE, "[ALIGN(V[i], T[i])]" );

    MyCollection<MyElement> X(&D, &A,
        "/myPersistentCollectionName");

    beginTransaction();
    X.hello();
    commitTransaction();
}

In this example, the programmer passes the string 
"/myPersistentCollectionName" into the collection’s constructor; this string is passed on to PersistentCollection, where it identifies a persistent collection on stable storage (in the Shore persistent object namespace) that is to be associated with the instantiated collection. If no collection named "/myPersistentCollectionName" exists when the program is run, then one is created. PersistentCollection contains a function

    int isPersistentNew();

that returns true if a persistent collection on stable storage is created when the collection is instantiated; the programmer can use this function to perform initialization on the persistent parts of a collection’s elements when the persistent parts are first created.

Access to the persistent part of a collection must be made within a transaction (see Figure 6.3 on Page 92 for a detailed explanation). beginTransaction() initiates a transaction, commitTransaction() ends a transaction normally, and abortTransaction() (or a system or program crash) ends a transaction and rolls back all persistent data modified during the transaction to their pre-transaction values. So to checkpoint the persistent part of a collection, the programmer can simply call commitTransaction(). In addition, if other programs will access the persistent data concurrently, the points at which commitTransaction() are called define the points at which the other programs may gain access to the collection’s persistent data.
Example Programs

To test the persistent collection implementation, I used it to implement checkpointing and inter-program communication for concurrent visualization and computation, in a version of the astrophysics simulation "Self Consistent Field" [35], henceforth referred to as SCF. Figure 6.4 sketches how initial and final I/O and checkpointing/restarting is

```
SelfConsistentField.h:
#include "PersistentCollection.h"
Collection SelfConsistentField : public PersistentCollection {
  ...
};

scf.h:
#include "SelfConsistentField.h"

void Processor_Main(int argc, char **argv)
{
  Processors P;
  Distribution D(num_of_elements, &P, BLOCK);
  Align A(num_of_elements, "{ALIGN(T[i], D[i])}\}"");

  // Declare the persistent collection X and perform initial input
  // (Persistent data is initialized in the constructor for X)
  // (If this is a restart, X will automatically contain its old values)
  // (If this is not a restart, the (programmer-provided) constructor for X will
  //   initialize the collection)
  beginTransaction();
  SelfConsistentField<Segment> X(&D, &A, "/scf");

  // Initialize non-persistent data
  // ...

  // Main computation loop
  while (X.n < nsteps) {
    X.n++;
    // Computation for time step n
    // ...
    if (X.n < nsteps) {
      // Checkpoint all persistent data
      commitTransaction();
      // At this point the data in X is read/write accessible by other programs
      // (e.g. visualization or computational steering programs)
      beginTransaction();
    }
  }

  // Final output of all persistent data
  commitTransaction();
}
```

Figure 6.4: Using Persistent Collections for I/O in an Astrophysics Simulation