

Sun[™] MPI 6.0 Software Programming and Reference Manual

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Contents

Preface vii

Data Types 17

1.

2.

Introduction to Sun MPI 1 Sun MPI Features 1 Sun MPI I/O 2 Sun MPI Library 3 Types of Libraries 3 Sun MPI Routines 4 Point-to-Point Communication Routines 4 One-Sided Communication Routines 5 Collective Communication Routines 12 Using the In-Place Option 13 Using Persistent Communication Requests 14 Managing Process Topologies 14 Name-Publishing Routines 14 Environmental Inquiry Routines 15

Packing and Unpacking Functions 15

Managing Communicators, Groups, and Contexts 15

```
Resource Reservation for Batch Processing 20
 Programming With Sun MPI 20
    Fortran Support 21
    Recommendations for All-to-All and All-to-One Communication 22
    Signals and MPI 22
 Multithreaded Programming 23
    Guidelines for Thread-Safe Programming 23
        MPI Wait(), MPI Waitall(), MPI Waitany(), MPI Waitsome() 24
        MPI_Cancel() 24
        MPI_Probe(), MPI_Iprobe() 24
        Collective Calls 25
        Communicator Operations 25
    Error Handlers 25
 Profiling Interface 26
 MPE: Extensions to the Library 29
       To Obtain and Build the MPE 29
Getting Started 31
 Header Files 31
    Sample Code 32
 Compiling and Linking 34
    Choosing a Library Path 37
    Stubbing Thread Calls 37
 Profiling With mpprof 38
 Basic Job Execution 38
    Executing With CRE 39
    Executing With LSF Suite 39
 Debugging
    Debugging With the Prism Environment 40
```

Starting Up Prism 40

Debugging With TotalView 41

Limitations 41

Related Documentation 42

Starting a New Job Using TotalView 42

Attaching to an mprun Job 44

Launching Sun MPI Batch Jobs Using TotalView 45

Debugging With MPE 47

4. Programming With Sun MPI I/O 49

Data Partitioning and Data Types 49

Definitions 50

Note for Fortran Users 51

Routines 51

File Manipulation 52

File Hints 52

File Views 53

Data Access 53

Data Access With Explicit Offsets 54

Data Access With Individual File Pointers 55

Pointer Manipulation 55

Data Access With Shared File Pointers 56

File Interoperability 57

File Consistency and Semantics 58

Sample Code 59

Partitioned Writing and Reading in a Parallel Job 59

Data Access Styles 63

Overlapping I/O With Computation and Communication 64

A. Sun MPI and Sun MPI I/O Routines 67

Sun MPI Routines 67

Sun MPI I/O Routines 95

B. Environment Variables 103

Informational Variables 103

General Performance Tuning 104

Tuning Memory for Point-to-Point Performance 106

Numerics 109

Synchronization of One-Sided Communications 109

Tuning Rendezvous 111

MPProf 112

Miscellaneous 113

C. Troubleshooting 117

MPI Messages 117

MPI I/O Error Handling 120

Adjusting System V Shared-Memory Limits 122

Preface

The *Sun MPI Software Programming and Reference Manual* describes the Sun[™] MPI library of message-passing routines and explains how to develop an MPI program on a Sun HPC system.

For the most part, this guide does not repeat information that is available in detail in the *MPI Standard*; it focuses instead on what is specific to the Sun MPI implementation.

You should be familiar with programming in C or Fortran, with parallel programming, and with the message-passing model.

Before You Read This Book

For general information about writing MPI programs, see "Related Documentation" on page x. Sun MPI is part of the Sun HPC ClusterToolsTM suite of software. For more information about running Sun MPI jobs, see the *Sun HPC ClusterTools Software User's Guide*. Product notes for Sun MPI are included in *Sun HPC ClusterTools Software Release Notes*.

Using UNIX Commands

This document does not describe how to use basic $UNIX^{\circledR}$ commands. For that type of information, see:

- Solaris Handbook for Sun Peripherals
- Online documentation for the Solaris[™] operating environment

■ Other software documentation that you received with your system

Typographic Conventions

Typeface*	Meaning	Examples
AaBbCc123	The names of commands, files, and directories; on-screen computer output	Edit your.login file. Use ls -a to list all files. % You have mail.
AaBbCc123	What you type, when contrasted with on-screen computer output	% su Password:
AaBbCc123	Book titles, new words or terms, words to be emphasized. Replace command-line variables with real names or values.	Read Chapter 6 in the <i>User's Guide</i> . These are called <i>class</i> options. You <i>must</i> be superuser to do this. To delete a file, type rm <i>filename</i> .

^{*} The settings on your browser might differ from these settings.

Shell Prompts

Shell	Prompt
C shell	machine-name%
C shell superuser	machine-name#
Bourne shell and Korn shell	\$
Bourne shell and Korn shell superuser	#

Related Documentation

Application	Title	Part Number
Sun HPC ClusterTools Documentation	Read Me First: Guide to Sun HPC ClusterTools 5 Software Documentation	817-0096-10
Sun HPC ClusterTools Software	Sun HPC ClusterTools 5 Software Release Notes	817-0081-10
	Sun HPC ClusterTools 5 Software Installation Guide	817-0082-10
	Sun HPC ClusterTools 5 Software	817-0090-10
	Performance Guide Sun HPC ClusterTools 5 Software User's	817-0084-10
	Guide Sun HPC ClusterTools 5 Software Administrator's Guide	817-0083-10
Sun S3L	Sun S3L 4.0 Software Programming Guide	817-0086-10
	Sun S3L 4.0 Software Reference Manual	817-0087-10
Prism™ graphical	Prism 7.0 Software User's Guide	817-0088-10
programming environment	Prism 7.0 Software Reference Manual	817-0089-10

A wealth of documentation on MPI is available on the World Wide Web. It includes the following:

■ The MPI home page, with links to specifications for the MPI-2 Standard:

```
http://www.mpi-forum.org
```

■ Additional Web sites that provide links to papers, talks, the MPI Standard, implementations, information about MPI-2, tutorials, and pointers to many other sources:

```
http://www.erc.msstate.edu/mpi/
http://www.arc.unm.edu/
```

Man Pages

Man pages are also available online for all the Sun MPI and MPI I/O routines and are accessible by means of the Solaris man command. These man pages are usually installed in /opt/SUNWhpc/man. Ask your system administrator for their location at your site.

Accessing Sun Documentation

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Introduction to Sun MPI

The MPI specification was developed by the MPI Forum, a group of software developers, computer vendors, academics, and computer-science researchers whose goal was to develop a standard for writing message-passing programs that would be efficient, flexible, and portable.

The outcome, known as the *MPI Standard*, was first published in 1993. The most recent version (MPI-2) was published in 1997. It was well received, and several implementations are available publicly.

 Sun^{TM} MPI is a complete library of message-passing routines, including all MPI 1.2–compliant and MPI 2–compliant routines. Chapter 2 provides an overview of the routines, Appendix A summarizes them, and the Sun man pages provide detailed descriptions.

Sun MPI Features

- Optimization for running with Sun HPC ClusterTools 5 software using C, C⁺⁺, Fortran 77, or Fortran 90.
- Integration with the Sun HPC ClusterTools[™] Runtime Environment (CRE) and the Sun GridEngine, LSF, and PBS distributed resource-management systems.
- Support for multithreaded programming.
- Seamless use of various network protocols.
- Multiprotocol support so that MPI selects the fastest available medium for each type of connection (such as shared memory, RSM, and ATM).
- Communication by shared memory for fast performance on clusters of SMPs
- $\blacksquare \ \ \ Finely \ tunable \ shared-memory \ communication.$
- Optimized collectives for symmetric multiprocessors (SMPs) and clusters of SMPs.

- MPI I/O support for parallel file I/O.
- Prism support, so that users can develop, run, and debug programs in the Prism programming environment.
- Implicit co-scheduling, whereby the Sun HPC spind daemon enables certain processes of a given MPI job on a shared-memory system to be scheduled at approximately the same time as other related processes. This co-scheduling reduces the load on the processors, thus reducing the effect that MPI jobs have on each other.
- Complete support of one-sided communication routines and name-publishing routines.
- Dynamic library support.
- MPI-2 dynamic support.

Sun MPI and MPI I/O provide full F77, C, and C++ support, as well as Basic F90 support.

Sun MPI I/O

File I/O in Sun MPI uses MPI 2–compliant routines for parallel file I/O. Chapter 4 describes these routines. Their man pages are provided online, and the routines are summarized in Appendix A.

Sun MPI Library

This chapter describes the Sun MPI library:

- "Types of Libraries" on page 3
- "Sun MPI Routines" on page 4
- "Managing Communicators, Groups, and Contexts" on page 15
- "Data Types" on page 17
- "Resource Reservation for Batch Processing" on page 20
- "Programming With Sun MPI" on page 20
- "Multithreaded Programming" on page 23
- "Profiling Interface" on page 26
- "MPE: Extensions to the Library" on page 29

Note – Sun MPI I/O is described separately, in Chapter 4.

Types of Libraries

Sun MPI contains four types of libraries, which represent two categories.

- 32- and 64-bit libraries —If you want to take advantage of the 64-bit capabilities of Sun MPI, you must explicitly link to the 64-bit libraries. The 32-bit libraries are the default in each category.
- Thread-safe and non-thread-safe libraries —For multithreaded programs, link with the thread-safe library in the appropriate category unless the program has only one thread calling MPI. For programs that are not multithreaded, you can link against either the thread-safe or the default (non-thread-safe) library. However, nonmultithreaded programs have better performance using the default library, as it does not incur the extra overhead of providing thread safety. Therefore, use the default libraries whenever possible for maximum performance.

For full information about linking to libraries, see "Compiling and Linking" on page 34.

Sun MPI Routines

This section gives a brief description of the routines in the Sun MPI library. All the Sun MPI routines are listed in Appendix A with brief descriptions and their C syntax. For detailed descriptions of individual routines, see the man pages or the *MPI Standard*. The routines are divided into these categories:

- "Point-to-Point Communication Routines" on page 4
- "One-Sided Communication Routines" on page 5
- "Collective Communication Routines" on page 12
- "Name-Publishing Routines" on page 14
- "Environmental Inquiry Routines" on page 15
- "Packing and Unpacking Functions" on page 15

Point-to-Point Communication Routines

Point-to-point communication routines include the basic send and receive routines in both blocking and nonblocking forms and in four modes.

A *blocking send* blocks until its message buffer can be written with a new message.

A blocking receive blocks until the received message is in the receive buffer.

Nonblocking sends and receives differ from blocking sends and receives in that they return immediately and their completion must be waited or tested for. It is expected that eventually nonblocking send and receive calls will allow the overlap of communication and computation.

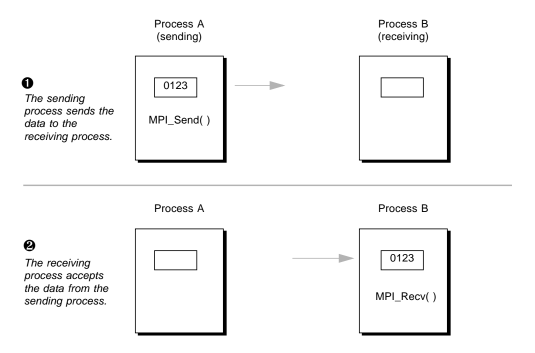
The four modes for MPI point-to-point communication are as follows:

- Standard The completion of a send implies that the message either is buffered internally or has been received. Users are free to overwrite the buffer that they passed in with any of the blocking send or receive routines, after the routine returns.
- *Buffered* The user guarantees a certain amount of buffering space.
- *Synchronous* Rendezvous semantics occur between sender and receiver; that is, a send blocks until the corresponding receive has occurred.

Ready – A send can be started only if the matching receive is already posted. The ready mode for sends is a way for the programmer to notify the system that the receive has been posted, so that the underlying system can use a faster protocol if it is available.

One-Sided Communication Routines

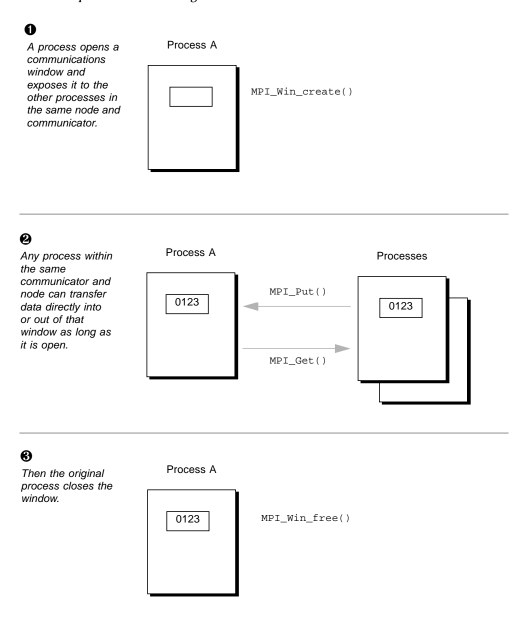
Standard MPI communication is two-sided. To complete a transfer of information, both the sending and receiving processes must call appropriate functions. The operation proceeds in two stages, as shown in the following figure.



This form of communication requires regular synchronization between the sending and receiving processes. That synchronization can become complicated if the receiving process does not know which process is sending it the data it needs. One-sided communication was developed to solve this problem and to reduce the amount of synchronization required even when both sending and receiving processes know each other's identities.

In one-sided communication, a process opens a window in memory and exposes it to all processes that belong to a particular communicator, provided they reside on the

same node. As long as that window is open, any process in the communicator and node can *put* data into it and *get* data out of it.



The *put* requires no complementary operation from the process that opened the window, and is equivalent to the combination of a *send* and *receive* operation in two-

sided MPI communication.

The functions used to implement one-sided MPI communication fall into three categories and are summarized in TABLE 2-1. You can find their definitions in the MPI Standard. Also, Appendix A of this document provides syntax summaries.

One-Sided Communication Routines TABLE 2-1

ndow Creation	
MPI_Win_create()	Creates a window in memory and exposes it to all processes in the communicator and node.
<pre>MPI_Win_free()</pre>	Closes the window created with ${\tt MPI_Win_create()}$. Requires barrier synchronization.
MPI_Win_get_group()	Returns a duplicate of the group of the communicator used to create the window.
ta Transfer	
MPI_Accumulate()	Combines data from a process with data already in the window. Different from MPI_Put() in that new data is appended to existing data instead of replacing it.
MPI_Get()	The calling process takes data directly from the window. The opposite of MPI_Put(). Equivalent to the combination of a send and receive operation originated by the receiving process.
MPI_Put()	The calling process loads its data directly into the buffer of the target process. Equivalent to the combination of a send and receive operation originated by the sending process. The opposite of MPI_Get().
nchronization	
MPI_Win_fence()	Blocks any process from operating on a particular window until all operations relating to that window have completed. Similar to MPI_Barrier(), but applies to a window instead of a communicator.
MPI_Win_lock()	Starts an RMA access epoch. While the lock is in place, only the process whose rank is specified in the function call can be accessed by RMA operations on the window.
MPI_Win_unlock()	Closes an RMA access epoch begun with a call to ${\tt MPI_Win_lock()}.$
MPI_Win_start()	Starts an RMA access epoch for window.
MPI_Win_complete()	Completes an RMA access epoch on <i>window</i> started by a call to MPI_Win_start().

TABLE 2-1 One-Sided Communication Routines (Continued)

MPI_Win_post()	Starts an RMA exposure epoch for the local window associated with <i>window</i> .
MPI_Win_wait()	Completes an RMA exposure epoch started by a call to MPI_Win_post() on <i>window</i> .
MPI_Win_test()	Attempts to complete an RMA exposure epoch; a nonblocking version of MPI_Win_wait().

Some special considerations apply to allocating memory for one-sided communications. For example:

- MPI_Alloc_mem allocates memory by means of SysV shared memory. If you find that you cannot allocate any more memory because of system-imposed limits, you can either increase shmsys:shminfo_shmseg in /etc/system or try to preallocate a large segment and then use parts of it for your needs.
- If the memory used for a communication window has been allocated by means of the MPI_Alloc_mem function, the memory region specified by the base and size parameters passed to MPI_Win_create should not exceed the limits of the memory region allocated using a call either to malloc or to the MPI_Alloc_mem function. If the memory has been allocated using a call to MPI_Alloc_mem, these limits are checked internally and the call to MPI_Win_create() returns MPI_ERR_OTHER.

Several one-sided communications routines support info keys. These keys, and their descriptions, are listed in TABLE 2-2.

 TABLE 2-2
 Info Keys Supported By One-Sided Communications Routines

Routine	Info Key	Description
MPI_Win_create()	no_locks	If MPI_Win_lock is called on a window created with this info key, the call fails. If this info key is present, it is assumed that the local window is never locked, allowing several internal checks to be skipped, permitting a more efficient implementation.
	sun_noexpose	An info key (unique to Sun MPI) that is interpreted only by the remote shared memory (RSM TM) protocol module when passed using MPI_Win_create (and while the RSM protocol is in use). When explicitly passed with a base address that was not allocated using MPI_Alloc_mem, sun_noexpose notifies the library not to expose address spaces not specified by the application.
	sun_rsmrexmit_hw	An info key (unique to Sun MPI) that is interpreted only by the RSM protocol module when passed using MPI_Win_create (and while the RSM protocol is in use). The sun_rsmrexmit_hw info key defines a high-water mark, in bytes, of data that will be sent before any error checking and retransmission of data is done. By default the RSM protocol module assumes the high-water mark to be 16KB.
	sun_shmeuid	The effective user ID (UID) to which the shared-memory segment is set for memory-based protocols, including shared memory (SHM) and RSM. Set this key only when you anticipate connections from programs run by other users. Valid only for server programs run as root.

 TABLE 2-2
 Info Keys Supported By One-Sided Communications Routines (Continued)

Routine	Info Key	Description
	sun_shmegid	The effective GID to which the shared-memory segment is set for memory-based protocols, including SHM and RSM. Set this key only when you anticipate connections from programs run by other users.
	sun_shmperm	The permissions to which the shared-memory segment is set, in octal, for memory-based protocols, including SHM and RSM. Set this key only when you anticipate connections from programs run by other users.
MPI_Alloc_mem	sun_shmeuid	The effective UID to which the shared-memory segment is set for memory-based protocols, including SHM and RSM. Set this key only when you anticipate connections from programs run by other users. Valid only for server programs run as root.
	sun_shmegid	The effective GID to which the shared-memory segment is set for memory-based protocols, including SHM and RSM. Set this key only when you anticipate connections from programs run by other users.
	sun_shmperm	The permissions to which the shared-memory segment is set, in octal, for memory-based protocols, including SHM and RSM. Set this key only when you anticipate connections from programs run by other users.

Several one-sided communications routines support assertions. These assertions, and their descriptions, are listed in TABLE 2-3.

Assertions Supported by One-Sided Communications Routines TABLE 2-3

Routine	Assertion Values	Description
MPI_Win_fence()	MPI_MODE_NOPRECEDE	The RSM protocol module recognizes this value and does not issue any RSM close barriers on the window passed into the MPI_Win_fence() call. The effect of this assertion value is that fence calls execute faster.
	MPI_MODE_NOSTORE MPI_MODE_NOPUT MPI_MODE_NOSUCCEED	All native Sun HPC ClusterTools protocol modules and generic one-sided functions ignore these values. However, they are available for use by third-party protocol modules
MPI_Win_start()	MPI_MODE_NOCHECK	When this value is passed in to this call, the library assumes that the post call on the target has been called and it is not necessary for the library to check to see if such a call has been made.
MPI_Win_post()	MPI_MODE_NOCHECK	When this value is passed in to this call, the library assumes that the post call on the target has been called and it is not necessary for the library to check to see if such a call has been made.

 TABLE 2-3
 Assertions Supported by One-Sided Communications Routines (Continued)

Routine	Assertion Values	Description
MPI_Win_lock()	MPI_MODE_NOCHECK	The Sun MPI library supports MPI_MODE_NOCHECK for the RSM protocol module only. When this lock is set, the lock and unlock calls in the RSM protocol module do not acquire the lock. However, data synchronization does occur.

Collective Communication Routines

Collective communication routines are blocking routines that involve all processes in a communicator and, in most cases, an intercommunicator. Collective communication includes broadcasts and scatters, reductions and gathers, all-gathers and all-to-alls, scans, and a synchronizing barrier call.

TABLE 2-4 Collective Communication Routines

Routine	Description
MPI_Bcast()	Broadcasts from one process to all others in a communicator or intercommunicator.
MPI_Scatter()	Scatters from one process to all others in a communicator or intercommunicator.
MPI_Scatterv()	Scatters from all processes to all others in a communicator or intercommunicator.
MPI_Reduce()	Reduces from all to one in a communicator or intercommunicator.
<pre>MPI_Allreduce()</pre>	Reduces and then broadcasts result to all nodes in a communicator or intercommunicator.
MPI_Allreducev()	Reduces from all processes and then broadcasts result to all nodes in a communicator or intercommunicator.
<pre>MPI_Reduce_scat ter()</pre>	Scatters a vector that contains results across the nodes in a communicator.
<pre>MPI_Gather()</pre>	Gathers from all to one in a communicator or intercommunicator.
MPI_Gatherv()	Gathers information from all processes in a communicator or intercommunicator.

Collective Communication Routines (Continued) TABLE 2-4

Routine	Description
MPI_Allgather()	Gathers and then broadcasts the results of the gather in a communicator or intercommunicator.
MPI_Allgatherv()	Gathers from all processes and then broadcasts the results of the gather in a communicator or intercommunicator.
MPI_Alltoall()	All processes send data to, and receive data from, all other processes in a communicator or intercommunicator.
MPI_Alltoallv()	Like MPI_Alltoall(), but user can use vector style (displacement and element count) to specify what data to send and receive.
MPI_Alltoallw()	$\label{like MPI_Alltoallv} Like ~ \texttt{MPI_Alltoallv}(), ~ but ~ user ~ can ~ specify ~ database ~ of individual ~ datablocks, in addition to ~ displacement ~ and ~ element ~ count.$
MPI_Scan()	Scans (performs a parallel prefix) across processes in a communicator or intercommunicator.
MPI_Exscan()	Performs an exclusive prefix reduction on data distributed across the calling processes.
<pre>MPI_Barrier()</pre>	Synchronizes processes in a communicator or intercommunicator (no data is transmitted).

Many of the collective communication calls have alternative vector forms, with which various amounts of data can be sent to or received from various processes. In addition, MPI_Alltoallw() accepts a database of individual datablocks.

The syntax and semantics of these routines are basically consistent with the point-topoint routines (upon which they are built), but there are restrictions to keep them from becoming too complex:

- The amount of data sent must exactly match the amount of data specified by the
- There is only one mode, a mode analogous to the standard mode of point-to-point routines.

Using the In-Place Option

Several collectives can pass MPI_IN_PLACE as the value of *send-buffer* at the root. When they do, *sendcount* and *sendtype* are ignored, and the contribution of the root to the gathered vector is assumed to be already in the correct location in the receive bugger. The collectives are as follows:

- MPI_Gather()
- MPI_Gatherv()
- MPI_Scatter()
- MPI_Scatterv()

```
■ MPI_Allgather()
```

- MPI Allgatherv()
- MPI_Reduce()
- MPI_AllReduce()
- MPI Reduce scatter()
- MPI_Scan()

Using Persistent Communication Requests

Sometimes within an inner loop of a parallel computation, a communication with the same argument list is executed repeatedly. The communication can be slightly improved by using a *persistent* communication request, which reduces the overhead for communication between the process and the communication controller. A persistent request can be thought of as a communication port or *half-channel*.

Managing Process Topologies

Process topologies are associated with communicators; they are optional attributes that can be given to an intracommunicator (not to an intercommunicator).

Recall that processes in a group are ranked from 0 to *n*–1. This linear ranking often reflects nothing of the logical communication pattern of the processes, which may be, for instance, a two- or three-dimensional grid. The logical communication pattern is referred to as a *virtual topology* (separate and distinct from any hardware topology). In MPI, two types of virtual topologies can be created: Cartesian (grid) topology and graph topology.

You can use virtual topologies in your programs by taking physical processor organization into account to provide a ranking of processors that optimizes communication.

Name-Publishing Routines

Name-publishing routines enable client applications to retrieve system-supplied port names. A server calls the MPI_Publish_name() function to publish the name of the service associated with a particular port name. A client application calls the MPI_Lookup_name(), passing it the published service name, and in return gets its associated port name. The server can also call the MPI_Unpublish_name() function to stop publishing names.

Sun's implementation of the *MPI Standard* does not provide a *scope* for the published names and does not allow a server to publish the same service name twice. The implementation consists of three routines:

```
■ MPI_Publish_name()
```

- MPI Unpublish name()
- MPI_Lookup_name()

Environmental Inquiry Routines

Environmental inquiry routines are used for starting up and shutting down error-handling routines and timers.

Few MPI routines can be called before $\texttt{MPI_Init()}$ or after $\texttt{MPI_Finalize()}$. Examples include $\texttt{MPI_Initialized()}$ and $\texttt{MPI_Version()}$. $\texttt{MPI_Finalize()}$ can be called only if there are no outstanding communications involving that process.

The set of errors handled by MPI depends upon the implementation. See Appendix C for tables listing the Sun MPI error classes.

Packing and Unpacking Functions

Sun's implementation of the *MPI Standard* provides functions for packing and unpacking messages to be exchanged within an MPI implementation, and in the external32 format used to exchange messages between MPI implementations.

```
    MPI_Pack()
    MPI_Unpack()
    MPI_Pack_size()
    MPI_Pack_external()
    MPI_Unpack_external()
    MPI_Pack_external_size()
```

Managing Communicators, Groups, and Contexts

A distinguishing feature of the *MPI Standard* is that it includes a mechanism for creating separate worlds of communication, accomplished through *communicators*, *groups*, and *contexts*.

A *communicator* specifies a group of processes that will conduct communication operations within a specified context without affecting or being affected by operations occurring in other groups or contexts elsewhere in the program. A communicator also guarantees that, within any group and context, point-to-point and collective communication are isolated from each other.

A *group* is an ordered collection of processes. Each process has a rank in the group; the rank runs from 0 to n–1. A process can belong to more than one group; its rank in one group has nothing to do with its rank in any other group.

A *context* is the internal mechanism by which a communicator guarantees safe communication space to the group.

At program startup, two default communicators are defined:

- MPI_COMM_WORLD, which has as a process group all the processes of the job
- MPI_COMM_SELF, which is equivalent to an identity communicator

The process group that corresponds to MPI_COMM_WORLD is not predefined, but can be accessed using MPI_COMM_GROUP. One MPI_COMM_SELF communicator is defined for each process, each of which has rank zero in its own communicator. For many programs, these are the only communicators needed.

Communicators are of two kinds: *intracommunicators*, which conduct operations within a given group of processes; and *intercommunicators*, which conduct operations between two groups of processes.

Communicators provide a *caching* mechanism, which allows an application to attach attributes to communicators. Attributes can be user data or any other kind of information.

New groups and new communicators are constructed from existing ones. Group constructor routines are local, and their execution does not require interprocessor communication. Communicator constructor routines are collective, and their execution can require interprocess communication.

You can also create an intercommunicator from two MPI processes that are connected by a socket. Use the MPI_Comm_join() function.

Note — Users who do not need any communicator other than the default MPI_COMM_WORLD communicator—that is, who do not need any sub- or supersets of processes—can plug in MPI_COMM_WORLD wherever a communicator argument is requested. In these circumstances, users can ignore this section and the associated routines. (These routines can be identified from the listing in Appendix A.)

Data Types

All Sun MPI communication routines have a data type argument. They can be primitive data types, such as integers or floating-point numbers, or they can be userdefined, derived data types that are specified in terms of primitive types.

Derived data types enable users to specify more general, mixed, and noncontiguous communication buffers, such as array sections and structures that contain combinations of primitive data types.

Fortran data types are listed in TABLE 2-5. Data types of Fortran used with the -r8 flag are listed in TABLE 2-6. C data types are listed in TABLE 2-7.

TABLE 2-5 Fortran Data Types

MPI Data Type	Fortran Data Type
MPI_INTEGER	INTEGER INTEGER*4
MPI_INTEGER1	INTEGER*1 (Fortran 90 only)
MPI_INTEGER2	INTEGER*2
MPI_INTEGER4	INTEGER*4
MPI_INTEGER8	INTEGER*8
MPI_REAL	REAL REAL*4
MPI_REAL4	REAL*4
MPI_REAL8	REAL*8
MPI_REAL16	REAL*16
MPI_DOUBLE_PRECISION	REAL*8 DOUBLE PRECISION
MPI_2DOUBLE_PRECISION	Pair of DOUBLE PRECISION variables*
MPI_COMPLEX	COMPLEX
MPI_LOGICAL	LOGICAL
MPI_CHARACTER	CHARACTER(1)
MPI_SIGNED_CHAR	INTEGER*1

 TABLE 2-5
 Fortran Data Types (Continued)

MPI Data Type	Fortran Data Type
MPI_DOUBLE_COMPLEX	DOUBLE COMPLEX
MPI_2REAL	Pair of REALs*
MPI_INTEGER2	INTEGER*2
MPI_INTEGER4	INTEGER*4
MPI_2INTEGER	Pair of INTEGERS*
MPI_BYTE	no corresponding Fortran data type
MPI_PACKED	no corresponding Fortran data type

^{*} For use with MINLOC and MAXLOC

TABLE 2-6 Fortran -r8 Data Types

MPI Data Type	Fortran –r8 Data Type
MPI_INTEGER	INTEGER*4
MPI_INTEGER1	INTEGER*1 (Fortran 90 only)
MPI_INTEGER2	INTEGER*2
MPI_INTEGER4	INTEGER*4
MPI_INTEGER8	INTEGER INTEGER*8
MPI_REAL	REAL*4
MPI_REAL4	REAL*4
MPI_REAL8	REAL REAL*8
MPI_REAL16	REAL*16 DOUBLE PRECISION
MPI_DOUBLE_PRECISION	REAL REAL*8
MPI_2DOUBLE_PRECISION	Pair of REAL*8*
MPI_COMPLEX	COMPLEX*4
MPI_LOGICAL	LOGICAL
MPI_CHARACTER	CHARACTER(1)
MPI_SIGNED_CHAR	INTEGER*1

 TABLE 2-6
 Fortran -r8 Data Types (Continued)

MPI Data Type Fortran –r8 Data Type	
	Fortiali =10 Data Type
MPI_DOUBLE_COMPLEX	COMPLEX
MPI_2REAL	Pair of REAL*4*
MPI_INTEGER2	INTEGER*2
MPI_INTEGER4	INTEGER*4
MPI_2INTEGER	Pair of INTEGER*4
MPI_BYTE	No corresponding Fortran data type
MPI_PACKED	No corresponding Fortran data type

^{*} For use with MINLOC and MAXLOC

 TABLE 2-7
 C Data Types

MPI Data Type	C Data Type
MPI_BYTE	No corresponding C data type
MPI_PACKED	No corresponding C data type
MPI_CHAR	signed char
MPI_SIGNED_CHAR	signed char
MPI_UNSIGNED_CHAR	unsigned char
MPI_SHORT	signed short int
MPI_UNSIGNED_SHORT	unsigned short int
MPI_INT	signed int
MPI_UNSIGNED	unsigned int
MPI_LONG	signed long int
MPI_UNSIGNED_LONG	unsigned long int
MPI_LONG_LONG_INT	long long int
MPI_FLOAT	float
MPI_DOUBLE	double
MPI_LONG_DOUBLE	long double
MPI_WCHAR	wchar_t
MPI_UNSIGNED_LONG_LONG	unsigned long long int
MPI_2INT	Pair of int*

TABLE 2-7 C Data Types (Continued)

MPI Data Type	C Data Type
MPI_FLOAT_INT	float and int*
MPI_DOUBLE_INT	double and int*
MPI_LONG_DOUBLE_INT	long double and int^*
MPI_LONG_INT	long and int*
MPI_SHORT_INT	short and int*

^{*} For use with MINLOC and MAXLOC

Resource Reservation for Batch Processing

If you plan to launch a job that uses the MPI_Comm_spawn or MPI_Comm_spawn_multiple functions, you must first reserve the resources with the resource manager that will run the job. As explained in the mprun.1 man page and the Sun HPC ClusterTools Software User's Guide, you can reserve those resources by adding the -nr flag to the mprun command.

When you launch a job with the mprun command from within a resource manager, the number of processes allocated for that job are stored in the environment variable MPI_UNIVERSE_SIZE. It is the sum of the processes allocated with the mprun command's -np flag, and reserved with its -nr flag.

Programming With Sun MPI

Although there are about 190 non-I/O routines in the Sun MPI library, you can write programs for a wide range of problems using only six routines, as described in TABLE 2-8.

TABLE 2-8 Six Basic MPI Routines

Routine	Description
MPI_Init()	Initializes the MPI library.
<pre>MPI_Finalize()</pre>	Finalizes the MPI library. This includes releasing resources used by the library.
<pre>MPI_Comm_size()</pre>	Determines the number of processes in a specified communicator.
MPI_Comm_rank()	Determines the rank of calling process within a communicator.
MPI_Send()	Sends a message.
MPI_Recv()	Receives a message.

This set of six routines includes the basic send and receive routines. Programs that depend heavily on collective communication might also include $\texttt{MPI_Bcast}()$ and $\texttt{MPI_Reduce}()$.

The functionality of these routines means you can have the benefit of parallel operations without having to learn the whole library at once. As you become more familiar with programming for message passing, you can start learning the more complex and esoteric routines and add them to your programs as needed.

See Appendix A for a complete list of Sun MPI routines.

Fortran Support

Sun MPI provides extended Fortran support, as described in Section 10.2 of the *MPI-2 Standard*. In other words, it provides basic Fortran support, plus additional functions that specifically support Fortran 90:

- MPI_Type_create_f90_complex()
- MPI_Type_create_f90_integer()
- MPI_Type_create_f90_real()
- MPI_Type_match_size()
- MPI_Sizeof()

Basic Fortran support provides the original Fortran bindings and an mpif.h file specified in the *MPI-1 Standard*. The mpif.h file is valid for both fixed- and free-source forms, as specified in the *MPI-2 Standard*.

The MPI interface is known to violate the Fortran standard in several ways, but it causes few problems for FORTRAN 77 programs. Violations of the standard can cause more significant problems for Fortran 90 programs, however, if you do not follow the guidelines recommended in the standard. If you are programming in

Fortran, and particularly if you are using Fortran 90, you should consult Section 10.2 of the *MPI-2 Standard* for detailed information about basic Fortran support in an MPI implementation.

Recommendations for All-to-All and All-to-One Communication

The Sun MPI library uses the TCP protocol to communicate over a variety of networks. MPI depends on TCP to ensure reliable, correct data flow. TCP's reliability compensates for unreliability in the underlying network, as the TCP retransmission algorithms handle any segments that are lost or corrupted. In most cases, this works well with good performance characteristics. However, when doing all-to-all and all-to-one communication over certain networks, a large number of TCP segments can be lost, resulting in poor performance.

You can compensate for this diminished performance over TCP in these ways:

- When writing your own algorithms, avoid flooding one node with a lot of data.
- If you need to do all-to-all or all-to-one communication, use one of the Sun MPI routines to do so. They are implemented in a way that avoids congesting a single node with lots of data. The following routines fall into this category:
 - MPI_Alltoall(), MPI_Alltoallv(), and MPI_Alltoallw() These have been implemented using a pairwise communication pattern, so that every rank is communicating with only one other rank at a given time.
 - MPI_Gather() and MPI_Gatherv() The root process sends ready-to-send packets to each nonroot-rank process to tell the processes to send their data. In this way, the root process can regulate how much data it is receiving at any one time. Using this ready-to-send method is associated with a minor performance cost, however. For this reason, you can override this method by setting the MPI_TCPSAFEGATHER environment variable to 0. (See Appendix B for information about environment variables.)

Signals and MPI

When running the MPI library over TCP, nonfatal SIGPIPE signals can be generated. To handle them, the library sets the signal handler for SIGPIPE to ignore, overriding the default setting (terminate the process). In this way the MPI library can recover in certain situations. You should therefore avoid changing the SIGPIPE signal handler.

The Sun MPI Fortran and C⁺⁺ bindings are implemented as wrappers on top of the C bindings. The profiling interface is implemented using weak symbols. This means a profiling library need contain only a profiled version of C bindings.

The SIGPIPES can occur when a process first starts communicating over TCP. This happens because the MPI library creates connections over TCP only when processes actually communicate with one another. There are some unavoidable conditions where SIGPIPES can be generated when two processes establish a connection. If you want to avoid any SIGPIPES, set the environment variable MPI_FULLCONNINIT, which creates all connections during MPI_Init() and avoids any situations that might generate a SIGPIPE. For more information about environment variables, see Appendix B.

Multithreaded Programming

When you are linked to one of the thread-safe libraries, Sun MPI calls are thread safe, in accordance with basic tenets of thread safety for MPI mentioned in the MPI-2 specification. As a result:

- When two concurrently running threads make MPI calls, the outcome is as if the calls executed in some order.
- Blocking MPI calls block the calling thread only. A blocked calling thread does not prevent progress of other runnable threads on the same process, nor does it prevent them from executing MPI calls. Thus, multiple sends and receives are concurrent.

Use MPI_Init_thread() in place of MPI_Init() to initialize the MPI execution environment with a predetermined level of thread support. Use the MPI_Is_thread_main() function to find out whether a thread is the one that called MPI_Init_thread().

Guidelines for Thread-Safe Programming

Each thread within an MPI process can issue MPI calls; however, threads are not separately addressable. That is, the rank of a send or receive call identifies a process, not a thread, which means that no order is defined for the case in which two threads call MPI_Recv() with the same tag and communicator. Such threads are said to be in conflict.

If threads within the same application post conflicting communication calls, data races will result. You can prevent such data races by using distinct communicators or tags for each thread.

In general, adhere to these guidelines:

- You must not have a request serviced by more than one thread. Although you can have an operation posted in one thread and then completed in another, you cannot have the operation completed in more than one thread.
- A data type or communicator must not be freed by one thread while it is in use by another thread.
- Once MPI_Finalize() is called, subsequent calls in any thread will fail.
- You must ensure that a sufficient number of lightweight processes (LWPs) are available for your multithreaded program. Failure to do so can degrade performance or even result in deadlock.
- You cannot stub the thread calls in your multithreaded program by omitting the threads libraries in the link line. The libmpi.so library automatically calls in the threads libraries, which effectively override any stubs.

The following sections describe more specific guidelines that apply for some routines. They also include some general considerations for collective calls and communicator operations that you should be aware of.

```
MPI_Wait(), MPI_Waitall(), MPI_Waitany(),
MPI_Waitsome()
```

In a program in which two or more threads call one of these routines, you must ensure that they are not waiting for the same request. Similarly, the same request cannot appear in the array of requests of multiple concurrent wait calls.

```
MPI_Cancel()
```

One thread must not cancel a request while that request is being serviced by another thread.

```
MPI_Probe(), MPI_Iprobe()
```

A call to MPI_Probe() or MPI_Iprobe() from one thread on a given communicator should not have a source rank and tags that match those of any other probes or receives on the same communicator. Otherwise, correct matching of message to probe call might not occur.

Collective Calls

Collective calls are matched on a communicator according to the order in which the calls are issued at each processor. All the processes on a given communicator must make the same collective call. You can avoid the effects of this restriction on the threads on a given processor by using a different communicator for each thread.

No process that belongs to the communicator may omit making a particular collective call; that is, none should be left "dangling."

Communicator Operations

Each of the communicator (or intercommunicator) functions operates simultaneously with each of the noncommunicator functions, regardless of what the parameters are and whether the functions are on the same or different communicators. However, if you are using multiple instances of the same communicator function on the same communicator where all parameters are the same, it cannot be determined which threads belong to which resultant communicator. Therefore, when concurrent threads issue such calls, you must ensure that the calls are synchronized in such a way that threads in separate processes participating in the same communicator operation are grouped together. Do this either by using a different base communicator for each call or by making the calls in single-thread mode before actually using them within the separate threads.

Note also these special situations:

- If you are using multiple instances of the same function with differing parameters and multiple threads, you must use separate communicators.
- When using splits with multiple instances of the same function with the same parameters, but with different threads at the split, you must use separate communicators.
 - For example, you might want to produce several communicators in separate sets of threads by performing MPI_Comm_split() on a base communicator. To ensure proper, thread-safe operation, you should replicate the base communicator with MPI_Comm_dup() (in the root thread or in one thread) and then perform MPI_Comm_split() on the resulting duplicate communicators.
- Do not free a communicator in one thread if it is still being used by another thread.

Error Handlers

When an error occurs as a result of an MPI call, the handler might not run on the same thread as the thread that made the error-raising call. In other words, you cannot assume that the error handler will execute in the local context of the thread

that made the error-raising call. The error handler can be executed by another thread on the same process, distinct from the one that returns the error code. Therefore, you cannot rely on local variables for error handling in threads; instead, use global variables from the process.

Profiling Interface

The Sun HPC ClusterTools software suite includes MPProf, a profiling tool to be used with applications that call Sun MPI library routines. When enabled, MPProf collects information about a program's message-passing activities in a set of intermediate files, one file per MPI process. Once the information is collected, you can invoke the MPProf command-line utility mpprof, which generates a report based on the profiling data stored in the intermediate files. You must enable MPProf before starting an MPI program. You do this by setting the environment variable MPI_PROFILE to 1.

If MPProf is enabled, it creates and initializes the intermediate files with header information when the program's MPI_Init call ends. It also creates an index file that contains a map of the intermediate files. mpprof uses this index file to find the intermediate files.

mpprof includes an interface for interacting with loadable protocol modules (loadable PMs). If an MPI program uses a loadable PM, this interface allows MPProf to collect profiling data that is specific to loadable PM activities.

An mpprof report contains the following classes of performance information:

- The percentage of total execution time spent in MPI calls across all processes
- The percentage of time each process spent in MPI calls
- The number of calls, amount of time spent, and number of bytes sent or received per MPI routine, averaged over all processes, with percent variation among processes
- Connectivity statistics (message count and volume) between processor pairs
- The settings of environment variables that have performance implications

You can control aspects of mpprof behavior with the following environment variables:

- MPI_PROFINTERVAL Use this environment variable to specify a data sampling period. When this value is greater than 0, a sequence of snapshots is recorded at the prescribed intervals. Each snapshot represents the MPI activity that occurred since the previous snapshot. The default behavior is to record a single snapshot at the time of the MPI_Finalize call.
- MPI_PROFDATADIR Use this environment variable to specify the location where the intermediate files created for each process rank will be stored.

- MPI_PROFINDEXDIR Use this environment variable to specify the location where the index file created for each profiled job will be stored.
- MPI_PROFMAXFILESIZE Use this environment variable to specify the maximum size of intermediate files.

The Sun HPC ClusterTools software suite also provides a conversion utility, mpdump, which converts the data from each intermediate file into a raw (unevaluated) user-readable format. You can use the ASCII files generated by mpdump as input to a report generator of your choice.

Once you've enabled MPProf profiling by setting MPI_PROFILE to 1 (and run a job using mprun) you will find a file in your working directory of the form

```
mpprof.index.rm.jid
```

Type

% mpprof mpprof.index.rm.jid

to view the profiling report.

Further instructions for using mpprof and mpdump are provided in the *Sun HPC ClusterTools Software User's Guide*.

Sun MPI meets the profiling interface requirements described in Chapter 8 of the MPI-1 Standard. This means you can write your own profiling library or choose from a number of available profiling libraries, such as those in the multiprocessing environment (MPE) from Argonne National Laboratory. (See "MPE: Extensions to the Library" on page 29 for more information.) The User's Guide for mpich, a Portable Implementation of MPI includes more detailed information about using profiling libraries.

FIGURE 2-1 provides a generic illustration of how the software fits together. In this example, the user is linking against a profiling library that collects information on MPI_Send(). No profiling information is being collected for MPI_Recv().

C profiling interfaces are needed even for Fortran programs. If there is profiling for both the Fortran and C version of an MPI function, then a Fortran call will encounter both profilings.

Be sure you make the library dynamic. A static library can experience the linker problems described in Section 8.4.3 of the MPI 1.1 Standard.

For compiling the program, the user's link line would look like this:

```
# cc ..... -1 library-name -1mpi
```

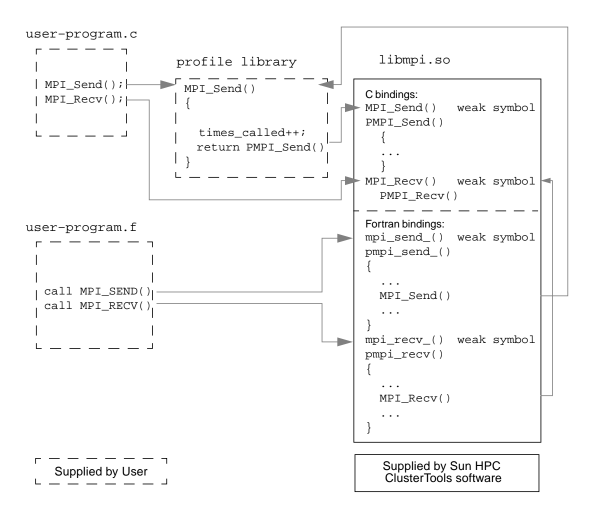


FIGURE 2-1 Sun MPI Profiling Interface

To clarify the layering of PMPI profiling, users need to understand the role of *weak symbols*. A weak symbol is such that, if a user defines the symbol, the user's definition is used. Otherwise, the associated function is used. The relation of weak symbols to associated functions is illustrated in FIGURE 2-2.

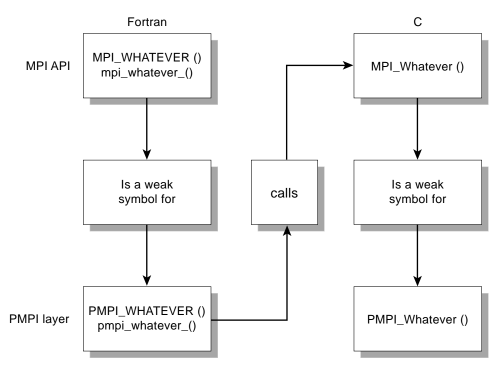


FIGURE 2-2 Layering in PMPI Profiling

MPE: Extensions to the Library

Although the Sun MPI library does not include or support the multiprocessing environment (MPE) available from Argonne National Laboratory (ANL), it is compatible with MPE. If you would like to use these extensions to the MPI library, see the following instructions for downloading them from ANL and building MPE yourself. Note that this procedure may change if ANL makes changes to MPE.

▼ To Obtain and Build the MPE

The MPE software is available from Argonne National Laboratory. The ${\tt mpe.tar.gz}$ file is about 240 Kbytes.

1. Use ftp to obtain the file.

```
ftp://ftp.mcs.anl.gov/pub/mpi/misc/mpe.tar.gz
```

2. Use gunzip and tar to decompress the software.

```
# gunzip mpe.tar.gz
# tar xvf mpe.tar
```

3. Change your current working directory to the mpe directory, and execute configure with the arguments shown.

```
# cd mpe
# configure -cc=cc -fc=f77 -opt=-I/opt/SUNWhpc/include
```

4. Execute a make.

```
# make
```

Note — Sun MPI does not include the MPE error handlers. You must call the debug routines MPE_Errors_call_dbx_in_xterm() and MPE_Signals_call_debugger() yourself.

Refer to the *User's Guide for mpich, a Portable Implementation of MPI* for information on how to use MPE. It is available at the Argonne National Laboratory web site:

http://www.mcs.anl.gov/mpi/mpich/

Getting Started

This chapter explains how to develop, compile and link, execute, and debug a Sun MPI program. The chapter focuses on what is specific to the Sun MPI implementation and, for the most part, does not repeat information that can be found in related documents. Information about programming with the Sun MPI I/O routines is in Chapter 4.

Header Files

Include syntax must be placed at the top of any program that calls Sun MPI routines.

- For C and C++, use #include <mpi.h>
- For Fortran, use

 INCLUDE 'mpif.h'

These lines enable the program to access the Sun MPI version of the mpi header file, which contains the definitions, macros, and function prototypes required when compiling the program. Ensure that you are referencing the *Sun* MPI include file.

The include files are usually found in /opt/SUNWhpc/include/ or /opt/SUNWhpc/include/v9/. If the compiler cannot find them, verify that they exist and are accessible from the machine on which you are compiling your code. The location of the include file is specified by a compiler option (see "Compiling and Linking" on page 34).

Sample Code

Two simple Sun MPI programs are available in /opt/SUNWhpc/examples/mpi and are included here in their entirety. In the same directory you will find the Readme file, which provides instructions for using the examples, and the make file Makefile.

CODE EXAMPLE 3-1 Simple Sun MPI Program in C: connectivity.c

```
* Test the connectivity between all processes.
* /
#pragma ident "@(#)connectivity.c 1.1 99/02/02"
#include <errno.h>
#include <stdio.h>
#include <stdlib.h>
#include <string.h>
#include <netdb.h>
#include <unistd.h>
#include <mpi.h>
int
main(int argc, char **argv)
   MPI Status status;
   int verbose = 0;
   int
              rank;
   int.
               np;
                                    /* number of processes in job */
   int.
               peer;
   int
                i;
   int
                j;
   MPI_Init(&argc, &argv);
   MPI_Comm_rank(MPI_COMM_WORLD, &rank);
   MPI_Comm_size(MPI_COMM_WORLD, &np);
   if (argc>1 && strcmp(argv[1], "-v")==0)
        verbose = 1;
   for (i=0; i<np; i++) {
        if (rank==i) {
            /* rank i sends to and receives from each higher rank */
            for(j=i+1; j<np; j++) {</pre>
                if (verbose)
                  printf("checking connection %4d <-> %-4d\n", i, j);
                MPI_Send(&rank, 1, MPI_INT, j, rank, MPI_COMM_WORLD);
```

CODE EXAMPLE 3-1 Simple Sun MPI Program in C: connectivity.c (Continued)

```
MPI_Recv(&peer, 1, MPI_INT, j, j, MPI_COMM_WORLD, &status);
}
} else if (rank>i) {
    /* receive from and reply to rank i */
    MPI_Recv(&peer, 1, MPI_INT, i, i, MPI_COMM_WORLD, &status);
    MPI_Send(&rank, 1, MPI_INT, i, rank, MPI_COMM_WORLD);
}

MPI_Barrier(MPI_COMM_WORLD);
if (rank==0)
    printf("Connectivity test on %d processes PASSED.\n", np);

MPI_Finalize();
return 0;
}
```

CODE EXAMPLE 3-2 Simple Sun MPI Program in Fortran: monte.f

```
! Estimate pi via Monte-Carlo method.
! Each process sums how many of samplesize random points generated
! in the square (-1,-1),(-1,1),(1,1),(1,-1) fall in the circle of
! radius 1 and center (0,0), and then estimates pi from the formula
! pi = (4 * sum) / samplesize.
! The final estimate of pi is calculated at rank 0 as the average of
! all the estimates.
!
       program monte
        include 'mpif.h'
        double precision drand
        external drand
        double precision x, y, pi, pisum
        integer*4 ierr, rank, np
        integer*4 incircle, samplesize
        parameter(samplesize=2000000)
        call MPI_INIT(ierr)
        call MPI_COMM_RANK(MPI_COMM_WORLD, rank, ierr)
        call MPI_COMM_SIZE(MPI_COMM_WORLD, np, ierr)
        seed random number generator
```

```
x = drand(2 + 11*rank)
        incircle = 0
       do i = 1, samplesize
          x = drand(0)*2.0d0 - 1.0d0 ! generate a random point
           y = drand(0)*2.0d0 - 1.0d0
           if ((x*x + y*y) .lt. 1.0d0) then
              incircle = incircle+1     ! point is in the circle
           endif
        end do
       pi = 4.0d0 * DBLE(incircle) / DBLE(samplesize)
!
       sum estimates at rank 0
        call MPI_REDUCE(pi, pisum, 1, MPI_DOUBLE_PRECISION, MPI_SUM,
                0 , MPI_COMM_WORLD, ierr)
       if (rank .eq. 0) then
           final estimate is the average
!
           pi = pisum / DBLE(np)
             print '(A, I4, A, F8.6, A)', 'Monte-Carlo estimate of pi by ', np,
                 ' processes is ',pi,'.'
       endif
       call MPI_FINALIZE(ierr)
        end
```

Compiling and Linking

Sun MPI programs are compiled with ordinary C, C^{++} , or Fortran compilers, just like any other C, C^{++} , or Fortran program, and linked with the Sun MPI library.

The mpf77, mpf90, mpcc, and mpcc utilities can be used to compile Fortran 77, Fortran 90, C, and C++ programs, respectively. For example, you might use the following entry to compile a Fortran 77 program that uses Sun MPI:

```
% mpf77 -fast -xarch=v8plusa -o a.out a.f -lmpi
```

See the man pages for more information on these utilities.

For performance, the single most important compilation switch is <code>-fast</code>. This is a macro that expands to settings appropriate for high performance for a general set of circumstances. Because its expansion varies from one compiler release to another, you might prefer to specify the underlying switches. To see what <code>-fast</code> expands to, use <code>-v</code> for "verbose" compilation output in Fortran, and <code>-#</code> for C. Also, <code>-fast</code> assumes native compilation, so you should compile on UltraSPARCTM processors.

The next important compilation switch is <code>-xarch</code>. Sun ONE Studio 7 Compiler Collection compilers set <code>-xarch</code> by default when you select <code>-fast</code> for native compilations. If you plan to compile on one type of processor and run the program on another type (nonnative compilation), be sure to use the <code>-xarch</code> flag. Also use it to compile in 64-bit mode. For UltraSPARC II processors, specify:

```
-xarch=v8plusa

or
-xarch=v9a
```

after -fast for 32-bit or 64-bit binaries, respectively. This version is only supported on Solaris 8 software. For UltraSPARC III processors, specify:

```
-xarch=v8plusb

or
-xarch=v9b
```

The v8plusb and v9b flags apply only to running on UltraSPARC III processors, and do not work when running on UltraSparc II processors. For more information, see the *Sun HPC ClusterTools Software Performance Guide* and the documents that came with your compiler.

Sun MPI programs compiled using the Sun ONE Studio 7 Compiler Collection Fortran compiler should be compiled with -xalias=actual. The -xalias=actual workaround requires patch 111718-01 (which requires 111714-01).

This recommendation arises because the MPI Fortran binding is inconsistent with the Fortran 90 standard in several respects. Specifically, this is documented in the MPI 2 standard, which you can find on the World Wide Web:

```
http://www-unix.mcs.anl.gov/mpi/mpi-standard/
mpi-report-2.0/node19.htm#Node19
```

This recommendation applies to the use of high levels of compiler optimization. A highly optimizing Fortran compiler could break MPI codes that use nonblocking operations.

The failure modes are varied and insidious and include the following:

- Silently incorrect answers
- Intermittent and mysterious floating-point exceptions
- Intermittent and mysterious hangs

If you will be using the Prism debugger, you must compile your program with compilers from the ForteTM 6 update 2, or Sun ONE Studio 7 (formerly Forte Development 7 software) Compiler Collections (see "Debugging" on page 39).

TABLE 3-1 Compile and Link Line Options for Sun MPI and Sun MPI I/O

Program	Options
C (nonthreaded example)	Use mpcc (below), or if you prefer: % cc filename.c -o filename \ -I/opt/SUNWhpc/include -L/opt/SUNWhpc/lib \ -R/opt/SUNWhpc/lib -lmpi
C++ Note that <i>x</i> .0 represents the version of your C++ compiler (6.0 for Forte 6 update 2, and 7.0 for Sun ONE Studio 7).	Use mpCC (below), or if you prefer: % CC filename.cc -o filename \ -I/opt/SUNWhpc/include -L/opt/SUNWhpc/lib \ -R/opt/SUNWhpc/lib -L/opt/SUNWhpc/lib/SCx.0 \ -R/opt/SUNWhpc/lib/SCx.y -mt -lmpi++ -lmpi
mpcc, mpCC	% mpcc -o filename filename.c -lmpi % mpcc -o filename filename.cc -mt -lmpi
Fortran 77 (nonthreaded example)	Use mpf77 (below), or if you prefer: % f77 -dalign filename.f -o filename \ -I/opt/SUNWhpc/include -L/opt/SUNWhpc/lib \ -R/opt/SUNWhpc/lib -lmpi
Fortran on a 64-bit system	<pre>% f77 -dalign filename.f -o filename \ -I/opt/SUNWhpc/include/v9 \ -L/opt/SUNWhpc/lib/sparcv9 \ -R/opt/SUNWhpc/lib/sparcv9 -lmpi</pre>
Fortran 90	Replace mpf77 with mpf90, or f77 with f90:
mpf90, mpf95	<pre>% mpf90 -o filename -dalign filename.f -lmpi % mpf95 -o filename -dalign filename.f -lmpi</pre>
Multithreaded programs and programs containing nonblocking MPI I/O routines	To support multithreaded code, replace <code>-lmpi</code> with <code>-lmpi_mt</code> . This change also supports programs with nonblocking MPI I/O routines. Note that <code>-lmpi</code> can be used for programs containing nonblocking MPI I/O routines, but <code>-lmpi_mt</code> must be used for multithreaded programs.

Note – For the Fortran interface, the -dalign option is necessary to avoid the possibility of bus errors. (The underlying C or C++ routines in Sun MPI internals assume that parameters and buffer types passed as REALs are double-aligned.)

Note — If your program has previously been linked to any static libraries, you must relink it to libmpi.so before executing it.

Choosing a Library Path

The paths for the MPI libraries, which you must specify when you are compiling and linking your program, are listed in TABLE 3-2.

TABLE 3-2 Sun MPI Libraries

Category	Description	Path: /opt/SUNWhpc/lib/
32-bit libraries	Default, not thread-safe	libmpi.so
	C^{++} (in addition to libmpi.so)	SC6.0/libmpi++.so
	Thread-safe	libmpi_mt.so
64-bit libraries	Default, not thread-safe	sparcv9/libmpi.so
	C++ (in addition to sparcv9/libmpi.so)	sparcv9/SC6.0/libmpi++.so
	Thread-safe	sparcv9/libmpi_mt.so

Stubbing Thread Calls

The libthread.so libraries are automatically linked into the respective libmpi.so libraries. This means that any thread-function calls in your program can be resolved by the libthread.so library. Simply omitting libthread.so from the link line does not cause thread calls to be stubbed out; you must remove the thread calls yourself. For more information about the libthread.so library, see its man page. (For the location of Solaris man pages at your site, see your system administrator.)

Profiling With mpprof

If you plan to extract MPI profiling information from the execution of a job, you need to set the MPI_PROFILE environment variable to 1 before you start the job execution.

```
% setenv MPI_PROFILE 1
```

If you want to set any other mpprof environment variables, you must set them also before starting the job. See Appendix B for detailed descriptions of the mpprof environment variables.

Basic Job Execution

The CRE environment provides close integration with batch-processing systems, also known as resource managers. You can launch parallel jobs from a batch system to control resource allocation, and continue to use the CRE environment to monitor job status. For a list of currently supported resource managers, see TABLE 3-3.

TABLE 3-3 Currently Supported Resource Managers

Resource manager	Name used with -x option to mprun	Version	Man page
Sun Grid Engine	sge	SGE 5.3	sge_cre.1
PBS	pbs	PBS 2.3.15	pbs_cre.1
	pbs	PBS Pro 5.x.x	pbs_cre.1
LSF	lsf	LSF 4.x	lsf_cre.1

To enable the integration between the CRE environment and the supported resource managers, you must call mprun from a script in the resource manager. Use the -x flag to specify the resource manager, and the -np and -nr flags to specify the resources you need. Instructions and examples for each resource manager are provided in the Sun HPC ClusterTools Software User's Guide.

Before starting your job, you might want to set one or more environment variables, which are also described in Appendix B and in the *Sun HPC ClusterTools Software Performance Guide*.

Executing With CRE

When using CRE software, parallel jobs are launched using the mprun command. For example, to start a job with six processes named mpijob, use this command:

```
% mprun -np 6 mpijob
```

Executing With LSF Suite

Parallel jobs can be either launched by the LSF Parallel Application Manager (PAM) or submitted in queues configured to run PAM as the parallel job starter. LSF's bsub command launches both parallel interactive and batch jobs. For example, to start a batch job named mpijob on four CPUs, use this command:

```
% bsub -n 4 pam mpijob
```

To launch an interactive job, add the -I argument to the command line. For example, to launch an interactive job named earth on a single CPU in the queue named sun, which is configured to launch jobs with PAM, use this command:

```
% bsub -q sun -Ip -n 1 earth
```

Debugging

Debugging parallel programs is notoriously difficult, because you are in effect debugging a program potentially made up of many distinct programs executing simultaneously. Even if the application is an SPMD (single-program, multiple-data) application, each instance can be executing a different line of code at any instant. The Prism development environment eases the debugging process considerably and is recommended for debugging with Sun HPC ClusterTools software.

Debugging With the Prism Environment

Note – To run the graphical version of the Prism environment, you must be running the Solaris 8 operating environment with either OpenWindowsTM software or the Common Desktop Environment (CDE), and with your DISPLAY environment variable set correctly. See the *Prism Software User's Guide* for information.

This section provides a brief introduction to the Prism development environment.

You can use a Prism session to debug more than one Sun MPI job at a time. To debug a child or client program it is necessary to launch an additional Prism session. If the child program is spawned using calls to MPI_Comm_spawn() or MPI_Comm_spawn_multiple(), Prism can (if enabled) debug the child program as well.

However, if an MPI job connects to another job, the current Prism session has control only of the parent or server MPI job. It cannot debug the children or clients of that job. This might occur, for example, when an MPI job sets up a client/server connection to another MPI job with MPI_Comm_accept or MPI_Comm_connect.

With the exception of programs using calls to MPI_Comm_spawn() or MPI_Comm_spawn_multiple(), to use the Prism environment to debug a Sun MPI program the program must be written in the SPMD style. In other words, all processes that make up a Sun MPI program must be running the same executable.

MPI_Comm_spawn_multiple can create multiple executables with only one job ID. Therefore, you can use the Prism environment to debug jobs with various executables that have been spawned with this command.

Starting Up Prism

To start Prism with a Sun MPI program, launch it from within the mprun command. For example,

```
% mprun -np 4 -x lsf prism -np 4 foo
```

launches Prism on executable foo with four processes.

This starts up a graphical version of Prism with your program loaded. You can then debug and visualize data in your Sun MPI program.

You can also attach Prism to running processes. First determine the job ID (not the individual process ID), jobname (or *jid*), using mpps. (See the *Sun HPC ClusterTools Software User's Guide* for further information about mpps.) Then specify the jid at the command line:

```
% prism foo 12345
```

This launches Prism and attaches it to the processes running in job 12345.

One important feature of the Prism environment is that it enables you to debug the Sun MPI program at any level of detail. You can look at the program as a whole, or you can look at subsets of processes within the program (for example, those that have an error condition) or at individual processes, all within the same debugging session. For complete information, see the *Prism Software User's Guide*.

Debugging With TotalView

TotalView is a third-party multiprocess debugger from Etnus that runs on many platforms. Support for using the TotalView debugger on Sun MPI applications includes:

- Making Sun HPC ClusterTools software compatible with the Etnus debugger TotalView
- Allowing Sun MPI jobs to be debugged by TotalView using the Sun Grid Engine (SGE), the Portable Batch System (PBS), and Platform Computing's Load Sharing Facility (LSF)
- Displaying Sun MPI message queues
- Allowing multiple instantiations of TotalView on a single cluster
- Supporting TotalView in Sun HPC ClusterTools software

The following sections provide a brief description of how to use the TotalView debugger with Sun MPI applications, including:

- "Limitations" on page 41
- "Related Documentation" on page 42
- "Starting a New Job Using TotalView" on page 42
- "Attaching to an mprun Job" on page 44
- "Launching Sun MPI Batch Jobs Using TotalView" on page 45

Refer to your TotalView documentation for more information about using TotalView.

Limitations

 Debuggable job restricted according to the license with Etnus. Contact the system administrator who installed TotalView for more details.

- Supports the SPARC platform *only*.
- TotalView 5 supports 32-bit application debugging only. However, TotalView 6 supports both 32-bit and 64-bit application debugging.
- Does *not* support MPI_Comm_spawn and MPI_Comm_spawn_multiple function calls. Use Prism to debug these function calls.
- Displays MPI_COMM_WORLD in the message queue graph only after MPI_Init() has occurred. Displays *neither* collective communicators *nor* MPI_COMM_SELF. Refer to the Etnus TotalView user's guide for more information.
- Does not display any buffer contents for unexpected messages in the message queue window.

Related Documentation

For more information, refer to the following related documentation:

- Sun HPC ClusterTools Software User's Guide
- Sun HPC ClusterTools Software Administrator's Guide
- Sun HPC ClusterTools software man pages
 - lsf_cre(1)
 - lsf cre admin(1M)
 - mpps(1M)
 - mprun(1M)
 - pbs_cre(1)
 - pbs cre admin(1M)
 - sge_cre(1)
 - sge_cre_admin(1M)
 - totalview_mprun(1M)
- **■** Etnus TotalView documentation

Starting a New Job Using TotalView

You can start a new job from the Total View Graphical User Interface (GUI) using:

- GUI method 1
- GUI method 2
- Command-line interface (CLI)

▼ To Start a New Job Using GUI Method 1

1. Type:

```
% totalview mprun [totalview args] -a [mprun args]
```

For example:

```
% totalview mprun -bg blue -a -np 4 /opt/SUNWhpc/mpi/conn.x
```

2. When the GUI appears, type g for go, or click Go in the TotalView window. TotalView may display a dialog box:

```
Process mprun is a parallel job. Do you want to stop the job now?
```

- 3. Click Yes to open the TotalView debugger window with the Sun MPI source window, if compiled with option -g, and to leave all processes in a traced state.
- ▼ To Start a New Job Using GUI Method 2
 - 1. Type:

```
% totalview
```

- 2. Select the menu option File and then New Program.
- 3. Type mprun as the executable name in the dialog box.
- 4. Click OK.

TotalView displays the main debug window.

5. Select the menu option Process and then Startup Parameters, which are the *mprun args*.

▼ To Start a New Job Using the CLI

1. Type:

```
% totalviewcli mprun [totalview args] -a [mprun args]
```

For example:

```
% totalviewcli mprun -a -np 4 /opt/SUNWhpc/mpi/conn.x
```

2. When the job starts, type dgo.

TotalView displays this message:

```
Process mprun is a parallel job. Do you want to stop the job now?
```

3. Type y to start the MPI job, attach TotalView, and leave all processes in a traced state.

Attaching to an mprun Job

This section describes how to attach to an already running mprun job from both the TotalView GUI and CLI.

▼ To Attach to a Running Job from the GUI

1. Find the host name and process identifier (PID) of the mprun job by typing:

```
% mpps -b
```

mprun displays the PID and host name in a similar manner to this example:

T0011111		MDDINI IIO GE
JOBNAME	MPRUN_PID	MPRUN_HOST
cre.99	12345	hpc-u2-9
cre.100	12601	hpc-u2-8

For more information, refer to the mpps(1M) man page, option -b.

2. In the TotalView GUI, select File and then New Program.

- 3. Type the PID in Process ID.
- 4. Type mprun in the field Executable Name.
- 5. Do one of the following:
 - Leave Remote Host blank if TotalView is running on the same node as the mprun job.
 - Enter the host name in Remote Host.
- 6. Click OK.

▼ To Attach to a Running Job From the CLI

1. Find the process identifier (PID) of the launched job.

See the example under the preceding GUI procedure. For more information, refer to the mpps(1M) man page, option -b.

2. Start totalviewcli by typing:

% totalviewcli

3. Attach the executable program to the mprun PID:

% dattach mprun mprun_pid

For example:

% dattach mprun 12601

Launching Sun MPI Batch Jobs Using TotalView

This section describes how to launch Sun MPI batch jobs, including:

- TotalView's control using the GUI only
- Interactive sessions using the GUI
- Interactive sessions using the CLI

This section provides examples of launching batch jobs in Sun Grid Engine (SGE). Refer to Chapter 5 of the *Sun HPC ClusterTools Software User's Guide* for descriptions of launching batch jobs in the Load Sharing Facility (LSF) and the Portable Batch System (PBS).

▼ To Execute Startup in Batch Mode for the TotalView GUI

Executing startup in batch mode for the TotalView CLI is not practical, because there is no controlling terminal for input and output. This procedure describes executing startup in batch mode for the TotalView GUI:

1. Write a batch script, which contains a line similar to the following:

```
% totalview mprun -a -x SGE /opt/SUNWhpc/mpi/conn.x
```

2. Then submit the script to SGE for execution with a command similar to the following:

```
% qsub -1 crfe 4 batch_script
```

The TotalView GUI appears upon successful allocation of resources and execution of the batch script in SGE.

▼ To Use the Interactive Mode

The interactive mode creates an xterm window for your use, so you can use either the TotalView GUI or the CLI.

1. Submit an interactive mode job to SGE with a command similar to the following:

```
% qsh -1 cre 4
```

The system displays an xterm window.

 ${\bf 2.} \ \ {\bf Run\ the\ following,\ or\ an\ equivalent\ path,\ to\ source\ the\ SGE\ environment:}$

```
% source /opt/sge/default/common/settings.csh
```

3. Execute a typical totalview or totalviewcli command.

TotalView GUI example:

```
% totalview mprun -a -x SGE /opt/SUNWhpc/mpi/conn.x
```

TotalView CLI example:

```
% totalviewcli mprun -a -x SGE /opt/SUNWhpc/mpi/conn.x
```

Debugging With MPE

The multiprocessing environment (MPE) available from Argonne National Laboratory includes a debugger that can also be used for debugging at the thread level. For information about obtaining and building MPE, see "MPE: Extensions to the Library" on page 29.

Programming With Sun MPI I/O

File I/O in Sun MPI is fully MPI-2 compliant. MPI I/O is specified as part of that standard, which was published in 1997. Its goal is to provide a library of routines featuring a portable parallel file system interface that is an extension of the MPI framework. See "Related Documentation" on page x for more information about the MPI-2 Standard.

MPI I/O models file I/O on message passing; that is, writing to a file is analogous to sending a message, and reading from a file is analogous to receiving a message. The MPI library provides a high-level way of partitioning data among processes, which saves you from having to specify the details involved in making sure that the right pieces of data go to the right processes. This section describes basic MPI I/O concepts and the Sun MPI I/O routines.

Data Partitioning and Data Types

MPI I/O uses the MPI model of communicators and derived data types to describe communication between processes and I/O devices. MPI I/O determines which processes are communicating with a particular I/O device. Derived data types can be used to define the layout of data in memory and of data in a file on the I/O device. (For more information about derived data types, see "Data Types" on page 17.) Because MPI I/O builds on MPI concepts, it's easy for a knowledgeable MPI programmer to add MPI I/O code to a program.

Data is stored in memory and in the file according to MPI data types. Herein lies one of MPI and MPI I/O's advantages: Because they provide a mechanism whereby you can create your own data types, you have more freedom and flexibility in specifying data layout in memory and in the file.

The library also simplifies the task of describing how your data moves from processor memory to the file and back again. You create derived data types that describe how the data is arranged in the memory of each process and how it should be arranged in that part of the disk file associated with the process.

Three functions are provided to handle the external 22 format. This format, defined by the MPI Forum, represents data in a universal format that is useful for exchanging data between implementations or for writing it to a file. The functions are:

```
MPI_Pack_external()MPI_Unpack_external()MPI_Pack_external_size()
```

The Sun MPI I/O routines are described in "Routines" on page 51. But first, to be able to define a data layout, you will need to understand some basic MPI I/O datalayout concepts. The next section explains some of the fundamental terms and concepts.

Definitions

The following terms are used to describe partitioning data among processes. FIGURE 4-1 illustrates some of these concepts.

- An *elementary data type* (or etype) is the unit of data access and positioning. It can be any MPI basic or derived data type. Data access is performed in elementary-data-type units, and offsets (defined later in this list) are expressed as a count of elementary data types.
- The *file type* (or filetype) is used to partition a file among processes; that is, a file type defines a template for accessing the file. It is either a single elementary data type or a derived MPI data type constructed from elementary data types. A file type can contain "holes," or extents of bytes that are not accessed by this process.
- A file *displacement* (or disp) is an absolute byte position counted from the beginning of a file. The displacement defines the location where a view begins (see FIGURE 4-1).
- A *view* defines the current set of data visible and accessible by a process from an open file in terms of a displacement, an elementary data type, and a file type. The pattern described by a file type is repeated, beginning at the displacement, to define the view.
- An *offset* is a position relative to the current view, expressed as a count of elementary data types. Holes in the view's file type are ignored when calculating this position.

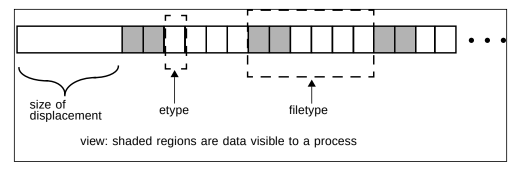


FIGURE 4-1 Displacement, the Elementary Data Type, the File Type, and the View

For a more detailed description of MPI I/O, see Chapter 9, "I/O," of the MPI-2 Standard.

Note for Fortran Users

When writing a Fortran program, you must declare the variable ADDRESS as follows:

```
INTEGER*MPI_ADDRESS_KIND ADDRESS
```

 ${\tt MPI_ADDRESS_KIND} \ is \ a \ constant \ defined \ in \ {\tt mpi.h.} \ This \ constant \ defines \ the length \ of \ the \ declared \ integer.$

Routines

This release of Sun MPI includes all the MPI I/O routines, which are defined in Chapter 9, "I/O," of the MPI-2 Standard.

Code samples that use many of these routines are provided in "Sample Code" on page 59.

File Manipulation

Collective coordination	Noncollective coordination
MPI_File_open()	MPI_File_delete()
MPI_File_close()	MPI_File_get_size()
<pre>MPI_File_set_size()</pre>	<pre>MPI_File_get_group()</pre>
<pre>MPI_File_preallocate()</pre>	<pre>MPI_File_get_amode()</pre>

MPI_File_open() and MPI_File_close() are collective operations that open and close a file, respectively; that is, all processes in a communicator group must together open or close a file. To achieve a single-user, UNIX-like open, set the communicator to MPI_COMM_SELF.

MPI_File_delete() deletes a specified file.

The routines MPI_File_set_size(), MPI_File_get_size(), MPI_File_get_group(), and MPI_File_get_amode() get and set information about a file. When using the collective routine MPI_File_set_size() on a UNIX file, if the size that is set is smaller than the current file size, the file is truncated at the position defined by size. If size is set to be larger than the current file size, the file size becomes size.

When the file size is increased this way with MPI_File_set_size(), new regions are created in the file with displacements between the old file size and the larger, newly set file size. Sun MPI I/O does not necessarily allocate file space for such new regions. You can reserve file space either by using MPI_File_preallocate() or by performing a read or write to unallocated bytes. MPI_File_preallocate() ensures that storage space is allocated for a set quantity of bytes for the specified file; however, its use is very "expensive" in terms of performance and disk space.

The routine ${\tt MPI_File_get_group}()$ returns a communicator group, but it does not free the group.

File Hints

The opaque info object enables you to provide hints for optimization of your code, making it run faster or more efficiently, for example. These hints are set for each file, using the MPI_File_open(), MPI_File_set_view(), MPI_File_set_info(), and MPI_File_delete() routines. MPI_File_set_info() sets new values for the specified file's hints. MPI_File_get_info() returns all the hints that the system currently associates with the specified file.

When using UNIX files, Sun MPI I/O provides four hints for controlling how much buffer space it uses to satisfy I/O requests: noncoll_read_bufsize, noncoll_write_bufsize, coll_read_bufsize, and coll_write_bufsize. These hints can be tuned for your particular hardware configuration and application to improve performance for both noncollective and collective data accesses. For example, if your application uses a single MPI I/O call to request multiple noncontiguous chunks that form a regular strided pattern in the file, you can adjust the noncoll_write_bufsize to match the size of the stride. Note that these hints limit the size of MPI I/O's underlying buffers but do not limit the amount of data a user can read or write in a single request.

File Views

The $\mathtt{MPI_File_set_view}()$ routine changes the view the process has of the data in the file, specifying its displacement, elementary data type, and file type, as well as setting the individual file pointers and shared file pointer to 0.

MPI_File_set_view() is a collective routine; all processes in the group must pass identical values for the file handle and the elementary data type, although the values for the displacement, the file type, and the info object can vary. However, if you use the data-access routines that use file positioning with a shared file pointer, you must also give the displacement and the file type identical values. The data types passed in as the elementary data type and the file type must be committed.

You can also specify the type of data representation for the file. See "File Interoperability" on page 57 for information about registering data representation identifiers.

Note – Displacements within the file type and the elementary data type must be monotonically nondecreasing.

Data Access

The 35 data-access routines are categorized according to file positioning. Data access can be achieved by any of these methods of file positioning:

- Explicit offset
- Individual file pointer
- Shared file pointer

This section discusses each of these methods in more detail.

While blocking I/O calls do not return until the request is completed, nonblocking calls do not wait for the I/O request to complete. A separate "request complete" call, such as MPI_Test() or MPI_Wait(), is necessary to confirm that the buffer is ready to be used again. Nonblocking routines have the prefix MPI_File_i, where the i stands for immediate.

All the nonblocking collective routines for data access are "split" into two routines, each with _begin or _end as a suffix. These *split collective* routines are subject to the semantic rules described in Section 9.4.5 of the *MPI-2 Standard*.

Data Access With Explicit Offsets

Synchronism	Noncollective coordination	Collective coordination
Blocking	<pre>MPI_File_read_at() MPI_File_write_at()</pre>	MPI_File_read_at_all() MPI_File_write_at_all()
Nonblocking or split collective	<pre>MPI_File_iread_at() MPI_File_iwrite_at()</pre>	<pre>MPI_File_read_at_all_begin() MPI_File_read_at_all_end() MPI_File_write_at_all_begin() MPI_File_write_at_all_end()</pre>

To access data at an explicit offset, specify the position in the file where the next data access for each process should begin. For each call to a data-access routine, a process attempts to access a specified number of file types of a specified data type (starting at the specified offset) into a specified user buffer.

The offset is measured in elementary data type units relative to the current view; moreover, holes are not counted when locating an offset. The data is read from (in the case of a read) or written into (in the case of a write) those parts of the file specified by the current view. These routines store the number of buffer elements of a particular data type actually read (or written) in the status object, and all the other fields associated with the status object are undefined. The number of elements that are read or written can be accessed using MPI_Get_count().

MPI_File_read_at() attempts to read from the file by the associated file handle returned from a successful MPI_File_open(). Similarly, MPI_File_write_at() attempts to write data from a user buffer to a file. MPI_File_iread_at() and MPI_File_iwrite_at() are the nonblocking versions of MPI_File_read_at() and MPI_File_write_at(), respectively.

MPI_File_read_at_all() and MPI_File_write_at_all() are collective versions of MPI_File_read_at() and MPI_File_write_at(), in which each process provides an explicit offset. The split collective versions of these nonblocking routines are listed in the preceding table.

Data Access With Individual File Pointers

Synchronism	Noncollective coordination	Collective coordination
Blocking	<pre>MPI_File_read() MPI_File_write()</pre>	<pre>MPI_File_read_all() MPI_File_write_all()</pre>
Nonblocking or split collective	<pre>MPI_File_iread() MPI_File_iwrite()</pre>	<pre>MPI_File_read_all_begin() MPI_File_read_all_end() MPI_File_write_all_begin() MPI_File_write_all_end()</pre>

For each open file, Sun MPI I/O maintains one individual file pointer per process per collective MPI_File_open(). For these data-access routines, MPI I/O implicitly uses the value of the individual file pointer. These routines use and update only the individual file pointers maintained by MPI I/O by pointing to the next elementary data type after the one that most recently has been accessed. The individual file pointer is updated relative to the current view of the file. The shared file pointer is neither used nor updated. (For data access with shared file pointers, see the next section.)

These routines have similar semantics to the explicit-offset data-access routines, except that the offset is defined here to be the current value of the individual file pointer.

MPI_File_read_all() and MPI_File_write_all() are collective versions of MPI_File_read() and MPI_File_write(), with each process using its individual file pointer.

MPI_File_iread() and MPI_File_iwrite() are the nonblocking versions of MPI_File_read() and MPI_File_write(), respectively. The split collective versions of MPI_File_read_all() and MPI_File_write_all() are listed in the preceding table.

Pointer Manipulation

```
MPI_File_seek
MPI_File_get_position
MPI_File_get_byte_offset
```

Each process can call the routine MPI_File_seek() to update its individual file pointer according to the update mode. The update mode has the following possible values:

- MPI_SEEK_SET The pointer is set to the offset.
- MPI SEEK CUR The pointer is set to the current pointer position plus the offset.
- MPI_SEEK_END The pointer is set to the end of the file plus the offset.

The offset can be negative for seeking backwards, but you cannot seek to a negative position in the file. The current position is defined as the elementary data item immediately following the last-accessed data item.

MPI_File_get_position() returns the current position of the individual file pointer relative to the current displacement and file type.

MPI_File_get_byte_offset() converts the offset specified for the current view to the displacement value, or absolute byte position, for the file.

Data Access With Shared File Pointers

Synchronism	Noncollective coordination	Collective coordination
Blocking	<pre>MPI_File_read_shared() MPI_File_write_shared()</pre>	MPI_File_read_ordered() MPI_File_write_ordered() MPI_File_seek_shared() MPI_File_get_position_shared()
Nonblocking or split collective	<pre>MPI_File_iread_shared() MPI_File_iwrite_shared()</pre>	<pre>MPI_File_read_ordered_begin() MPI_File_read_ordered_end() MPI_File_write_ordered_begin() MPI_File_write_ordered_end()</pre>

Sun MPI I/O maintains one shared file pointer per collective MPI_File_open() (shared among processes in the communicator group that opened the file). As with the routines for data access with individual file pointers, you can also use the current value of the shared file pointer to specify the offset of data accesses implicitly. These routines use and update only the shared file pointer; the individual file pointers are neither used nor updated by any of these routines.

These routines have similar semantics to the explicit-offset data-access routines, except:

- The offset is defined here to be the current value of the shared file pointer.
- Multiple calls (one for each process in the communicator group) affect the shared file pointer routines as if the calls were serialized.
- All processes must use the same file view.

After a shared file pointer operation is initiated, it is updated, relative to the current view of the file, to point to the elementary data item immediately following the last one requested, regardless of the number of items actually accessed.

MPI_File_read_shared() and MPI_File_write_shared() are blocking routines that use the shared file pointer to read and write files, respectively. The order of serialization is not deterministic for these noncollective routines, so you need to use other methods of synchronization if you want to impose a particular order.

```
MPI_File_iread_shared() and MPI_File_iwrite_shared() are the
nonblocking versions of MPI_File_read_shared() and
MPI_File_write_shared(), respectively.
```

MPI_File_read_ordered() and MPI_File_write_ordered() are the collective versions of MPI_File_read_shared() and MPI_File_write_shared(). They must be called by all processes in the communicator group associated with the file handle, and the accesses to the file occur in the order determined by the ranks of the processes within the group. After all the processes in the group have issued their respective calls, for each process in the group, these routines determine the position of the shared file pointer after all processes with ranks lower than this process's rank had accessed their data. Then data is accessed (read or written) at that position. The shared file pointer is then updated by the amount of data requested by all processes of the group.

The split collective versions of MPI_File_read_ordered() and MPI_File_write_ordered() are listed in the preceding table.

 $\label{eq:mpi_seek_shared()} \begin{tabular}{ll} MPI_File_seek_shared() is a collective routine, and all processes in the communicator group associated with the particular file handler must call $$ MPI_File_seek_shared()$ with the same file offset and the same update mode. All the processes are synchronized with a barrier before the shared file pointer is updated. $$ $$$

The offset can be negative for seeking backwards, but you cannot seek to a negative position in the file. The current position is defined as the elementary data item immediately following the last-accessed data item, even if that location is a hole.

MPI_File_get_position_shared() returns the current position of the shared file pointer relative to the current displacement and file type.

File Interoperability

```
MPI_Register_datarep()
MPI_File_get_type_extent()
```

Sun MPI I/O supports the basic data representations described in Section 9.5 of the *MPI-2 Standard*:

■ *native* – With native representation, data is stored exactly as in memory, in other words, in Solaris/UltraSPARC data representation. This format offers the highest performance and no loss of arithmetic precision. It should be used only in a

homogeneous environment, that is, on Solaris/UltraSPARC nodes running Sun HPC ClusterTools software. It also can be used when the MPI application will perform the data type conversions itself.

- *internal* With internal representation, data is stored in an implementation-dependent format, such as for Sun MPI.
- external32 With external32 representation, data is stored in a portable format, prescribed by the MPI-2 and IEEE standards.

These data representations, as well as any user-defined representations, are specified as an argument to MPI_File_set_view().

You can create user-defined data representations with MPI_Register_datarep(). Once a data representation has been defined with this routine, you can specify it as an argument to MPI_File_set_view(), so that subsequent data-access operations can call the conversion functions specified with MPI_Register_datarep().

If the file data representation is anything but native, you must be careful when constructing elementary data types and file types. For those functions that accept displacements in bytes, the displacements must be specified in terms of their values in the file for the file data representation being used.

You can use MPI_File_get_type_extent() to calculate the extents of data types in the file. The extent is the same for all processes accessing the specified file. If the current view uses a user-defined data representation, MPI_File_get_type_extent() uses one of the functions specified in setting the data representation to calculate the extent.

File Consistency and Semantics

Noncollective coordination	Collective coordination	
MPI_File_get_atomicity()	<pre>MPI_File_set_atomicity() MPI_File_sync()</pre>	

The routines ending in _atomicity enable you either to set a file's mode as atomic or nonatomic, or to query which mode it is in. In *atomic mode*, all operations within the communicator group that opens a file are completed as if sequentialized into a serial order. In *nonatomic mode*, no such guarantee is made. In nonatomic mode, MPI_File_sync() can be used to ensure weak consistency.

The default mode varies with the number of nodes you are using. If you are running a job on a single node, a file is in *nonatomic* mode by default when it is opened. If you are running a job on more than one node, a file is in *atomic* mode by default.

MPI_File_set_atomicity() is a collective call that sets the consistency semantics for data-access operations. All the processes in the group must pass identical values for both the file handle and the Boolean flag that indicates whether atomic mode is set.

MPI_File_get_atomicity() returns the current consistency semantics for data-access operations. Again, a Boolean flag indicates whether the atomic mode is set.

Note – In some cases, setting atomicity to false can provide better performance. The default atomicity value on a cluster is true. The lack of synchronization among the distributed caches on a cluster can prevent your data from completing in the desired state. In these circumstances, you might suffer performance disadvantages with atomicity set to true, especially when the data accesses overlap.

Sample Code

This section provides sample code to get you started with programming your I/O using Sun MPI. The first example shows how a parallel job can partition file data among its processes. That example is then adapted to use a broad range of other I/O programming styles supported by Sun MPI I/O. The last code sample illustrates the use of the nonblocking MPI I/O routines.

Remember that MPI I/O is part of MPI, so be sure to call $\texttt{MPI_Init()}$ before calling any MPI I/O routines, and call $\texttt{MPI_Finalize()}$ at the end of your program, even if you use only MPI I/O routines.

Partitioned Writing and Reading in a Parallel Job

MPI I/O was designed to enable processes in a parallel job to request multiple data items that are noncontiguous within a file. Typically, a parallel job partitions file data among the processes.

One method of partitioning a file is to derive the offset at which to access data from the rank of the process. The rich set of MPI derived types also makes it easy to partition file data. For example, you could create an MPI vector type as the filetype passed into ${\tt MPI_File_set_view()}$. Because vector types do not end with a hole, you would make a call to either ${\tt MPI_Type_create_resized()}$ or ${\tt MPI_Type_ub()}$ to complete the partition. This call would lengthen the extent to include holes at the end of the type for processes with higher ranks. You can create a partitioned file by passing various displacements to ${\tt MPI_File_set_view()}$. Each

of these displacements would be derived from the process's rank. Consequently, offsets would not need to be derived from the ranks, because only the data in the portion of the partition belonging to the process would be visible to the process.

The following example uses the first method that derives the file offsets directly from the rank of the process. Each process writes and reads NUM_INTS integers starting at the offset rank * NUM_INTS. It passes an explicit offset to the MPI I/O data-access routines MPI_File_write_at() and MPI_File_read_at(). It calls MPI_Get_elements() to find out how many elements were written or read. To verify that the write was successful, it compares the data written and read as well as set up an MPI_Barrier() before calling MPI_File_get_size() to verify that the file is the size expected upon completion of all the writes of the process.

Note that MPI_File_set_view() was called to set the view of the file as essentially an array of integers instead of the UNIX-like view of the file as an array of bytes. Thus, the offsets that are passed to MPI_File_write_at() and MPI_File_read_at() are indices into an array of integers and not a byte offset.

In CODE EXAMPLE 4-1, each process writes and reads NUM_INTS integers to a file using MPI_File_write_at() and MPI_File_read_at(), respectively.

CODE EXAMPLE 4-1 Writing and Reading Integers to a File

```
/* wr_at.c
 * Example to demonstrate use of MPI_File_write_at and MPI_File_read_at
* /
#include <stdio.h>
#include "mpi.h"
#define NUM_INTS 100
void sample_error(int error, char *string)
  fprintf(stderr, "Error %d in %s\n", error, string);
  MPI Finalize();
  exit(-1);
}
void
main( int argc, char **argv )
  char filename[128];
  int i, rank, comm_size;
  int *buff1, *buff2;
```

CODE EXAMPLE 4-1 Writing and Reading Integers to a File (Continued)

```
MPI File fh;
MPI_Offset disp, offset, file_size;
MPI_Datatype etype, ftype, buftype;
MPI Info info;
MPI_Status status;
int result, count, differs;
if(argc < 2) {
  fprintf(stdout, "Missing argument: filename\n");
  exit(-1);
strcpy(filename, argv[1]);
MPI_Init(&argc, &argv);
/* get this processor's rank */
result = MPI_Comm_rank(MPI_COMM_WORLD, &rank);
if(result != MPI_SUCCESS)
  sample_error(result, "MPI_Comm_rank");
result = MPI_Comm_size(MPI_COMM_WORLD, &comm_size);
if(result != MPI SUCCESS)
  sample_error(result, "MPI_Comm_size");
/* communicator group MPI_COMM_WORLD opens file "foo"
   for reading and writing (and creating, if necessary) */
result = MPI_File_open(MPI_COMM_WORLD, filename,
             MPI MODE RDWR | MPI MODE CREATE, (int)NULL, &fh);
if(result != MPI_SUCCESS)
  sample_error(result, "MPI_File_open");
/* Set the file view which tiles the file type MPI_INT, starting
   at displacement 0. In this example, the etype is also MPI_INT. */
disp = 0;
etype = MPI_INT;
ftype = MPI_INT;
info = (MPI Info)NULL;
result = MPI_File_set_view(fh, disp, etype, ftype, (char *)NULL, info);
if(result != MPI_SUCCESS)
  sample_error(result, "MPI_File_set_view");
/* Allocate and initialize a buffer (buff1) containing NUM_INTS integers,
   where the integer in location i is set to i. */
buff1 = (int *)malloc(NUM INTS*sizeof(int));
for(i=0;i<NUM_INTS;i++) buff1[i] = i;</pre>
/* Set the buffer type to also be MPI_INT, then write the buffer (buff1)
```

Writing and Reading Integers to a File (Continued) CODE EXAMPLE 4-1

```
starting at offset 0, i.e., the first etype in the file. */
buftype = MPI_INT;
offset = rank * NUM_INTS;
result = MPI_File_write_at(fh, offset, buff1, NUM_INTS, buftype, &status);
if(result != MPI_SUCCESS)
  sample_error(result, "MPI_File_write_at");
result = MPI_Get_elements(&status, MPI_BYTE, &count);
if(result != MPI SUCCESS)
  sample error(result, "MPI Get elements");
if(count != NUM_INTS*sizeof(int))
  fprintf(stderr, "Did not write the same number of bytes as requested\n");
else
  fprintf(stdout, "Wrote %d bytes\n", count);
/* Allocate another buffer (buff2) to read into, then read NUM_INTS
   integers into this buffer.
buff2 = (int *)malloc(NUM_INTS*sizeof(int));
result = MPI_File_read_at(fh, offset, buff2, NUM_INTS, buftype, &status);
if(result != MPI_SUCCESS)
  sample_error(result, "MPI_File_read_at");
/* Find out how many bytes were read and compare to how many
   we expected */
result = MPI_Get_elements(&status, MPI_BYTE, &count);
if(result != MPI_SUCCESS)
  sample_error(result, "MPI_Get_elements");
if(count != NUM INTS*sizeof(int))
  fprintf(stderr, "Did not read the same number of bytes as requested\n");
else
  fprintf(stdout, "Read %d bytes\n", count);
/* Check to see that each integer read from each location is
   the same as the integer written to that location. */
differs = 0;
for(i=0; i<NUM_INTS; i++) {</pre>
  if(buff1[i] != buff2[i]) {
    fprintf(stderr, "Integer number %d differs\n", i);
    differs = 1;
  }
if(!differs)
  fprintf(stdout, "Wrote and read the same data\n");
MPI_Barrier(MPI_COMM_WORLD);
result = MPI_File_get_size(fh, &file_size);
```

CODE EXAMPLE 4-1 Writing and Reading Integers to a File (Continued)

```
if(result != MPI_SUCCESS)
    sample_error(result, "MPI_File_get_size");

/* Compare the file size with what we expect */
    /* To see a negative response, make the file preexist with a larger
        size than what is written by this program */
    if(file_size != (comm_size * NUM_INTS * sizeof(int)))
        fprintf(stderr, "File size is not equal to the write size\n");

result = MPI_File_close(&fh);
    if(result != MPI_SUCCESS)
        sample_error(result, "MPI_File_close");

MPI_Finalize();

free(buff1);
    free(buff2);
}
```

Data Access Styles

You can adapt CODE EXAMPLE 4-1 to support the I/O programming style that best suits your application. Essentially, there are three dimensions on which to choose an appropriate data access routine for a particular task: file pointer type, collective or noncollective, and blocking or nonblocking.

You need to choose which file pointer type to use: explicit, individual, or shared. The CODE EXAMPLE 4-1 used an explicit pointer and passed it directly as the offset parameter to the MPI_File_write_at() and MPI_File_read_at() routines. Using an explicit pointer is equivalent to calling MPI_File_seek() to set the individual file pointer to offset, and then calling MPI_File_write() or MPI_File_read(), which is directly analogous to calling UNIX lseek() and write() or read(). If each process accesses the file sequentially, individual file pointers save you the effort of recalculating offset for each data access. A different shared file pointer could be used in situations where all the processes needed to cooperatively access a file in a sequential way, such as to write log files.

Collective data-access routines enable you to enforce some implicit coordination among the processes in a parallel job when making data accesses. For example, if a parallel job alternately reads in a matrix and performs computation on it, but cannot progress to the next stage of computation until all processes have completed the last stage, then a coordinated effort between processes when accessing data might be more efficient. In CODE EXAMPLE 4-1, you could easily append the suffix _all to MPI_File_write_at() and MPI_File_read_at() to make the accesses collective. By coordinating the processes, you could achieve greater efficiency in the

MPI library or at the file system level in buffering or caching the next matrix. In contrast, noncollective accesses are used when it is not evident that any benefit would be gained by coordinating disparate accesses by each process. UNIX file accesses are noncollective.

Overlapping I/O With Computation and Communication

MPI I/O also supports nonblocking versions of each of the data-access routines—that is, the data-access routines that have the letter i before write or read in the routine name (i stands for *immediate*). By definition, nonblocking I/O routines return immediately after the I/O request has been issued and do not wait until the I/O request has completed. This functionality enables you to perform computation and communication at the same time as the I/O. Because large I/O requests can take a long time to complete, this provides a way to more efficiently utilize your program's waiting time.

As in the previous example, parallel jobs often partition large matrices stored in files. These parallel jobs can use many large matrices, or matrices that are too large to fit into memory at once. Thus, each process can access the multiple and/or large matrices in stages. During each stage, a process reads in a chunk of data, and then performs a computation on it (which can involve communicating with the other processes in the parallel job). While performing the computation and communication, the process could issue a nonblocking I/O read request for the next chunk of data. Similarly, once the computation on a particular chunk has completed, a nonblocking write request could be issued before performing computation and communication on the next chunk.

The following example code illustrates the use of a nonblocking data-access routine. Note that like nonblocking communication routines, the nonblocking I/O routines require a call to MPI_Wait() to wait for the nonblocking request to complete, or repeated calls to MPI_Test() to determine when the nonblocking data access has completed. Once complete, the write or read buffer is available for use again by the program.

In CODE EXAMPLE 4-2, each process reads and writes *NUM_BYTES* bytes to a file using the nonblocking MPI I/O routines MPI_File_iread_at() and MPI_File_iwrite_at(), respectively. Note the use of MPI_Wait() and MPI_Test() to determine when the nonblocking requests have completed.

CODE EXAMPLE 4-2 Reading and Writing Bytes to a File

```
/* iwr_at.c
 * Example to demonstrate use of MPI_File_iwrite_at and MPI_File_iread_at
* /
#include <stdio.h>
#include "mpi.h"
#define NUM_BYTES 100
void sample_error(int error, char *string)
  fprintf(stderr, "Error %d in %s\n", error, string);
 MPI_Finalize();
  exit(-1);
}
void
main( int argc, char **argv )
  char filename[128];
  char *buff;
  MPI File fh;
 MPI_Offset offset;
 MPI_Request request;
  MPI Status status;
  int i, rank, flag, result;
  if(argc < 2) {
    fprintf(stdout, "Missing argument: filename\n");
    exit(-1);
  strcpy(filename, argv[1]);
  MPI_Init(&argc, &argv);
  result = MPI_Comm_rank(MPI_COMM_WORLD, &rank);
  if(result != MPI_SUCCESS)
    sample_error(result, "MPI_Comm_rank");
  result = MPI_File_open(MPI_COMM_WORLD, filename,
               MPI_MODE_RDWR | MPI_MODE_CREATE,
               (MPI_Info)NULL, &fh);
  if(result != MPI_SUCCESS)
```

CODE EXAMPLE 4-2 Reading and Writing Bytes to a File (Continued)

```
sample error(result, "MPI File open");
buff = (char *)malloc(NUM BYTES*sizeof(char));
for(i=0;i<NUM BYTES;i++) buff[i] = i;</pre>
offset = rank * NUM_BYTES;
result = MPI_File_iread_at(fh, offset, buff, NUM_BYTES,
                 MPI_BYTE, &request);
if(result != MPI_SUCCESS)
  sample_error(result, "MPI_File_iread_at");
/* Perform some useful computation and/or communication */
result = MPI_Wait(&request, &status);
buff = (char *)malloc(NUM_BYTES*sizeof(char));
for(i=0;i<NUM_BYTES;i++) buff[i] = i;</pre>
result = MPI_File_iwrite_at(fh, offset, buff, NUM_BYTES,
                  MPI_BYTE, &request);
if(result != MPI_SUCCESS)
  sample_error(result, "MPI_File_iwrite_at");
/* Perform some useful computation and/or communication */
flag = 0;
i = 0;
while(!flag) {
   result = MPI_Test(&request, &flag, &status);
   i++;
   /* Perform some more computation or communication, if possible */
}
result = MPI File close(&fh);
if(result != MPI SUCCESS)
  sample_error(result, "MPI_File_close");
MPI_Finalize();
fprintf(stdout, "Successful completion\n");
free(buff);
```

Sun MPI and Sun MPI I/O Routines

The tables in this appendix list the Sun MPI and Sun MPI I/O routines, along with the C syntax of the routines and a brief description of each. The routines are found in two sections:

- "Sun MPI Routines" on page 67
- "Sun MPI I/O Routines" on page 95

For more information about the routines, see their online man pages, usually found in

/opt/SUNWhpc/man/

Your system administrator can tell you where they are installed at your site.

Sun MPI Routines

TABLE A-1 lists the Sun MPI routines in alphabetical order. The following sections list the routines by functional category.

Point-to-Point Communication

Blocking Routines

```
MPI_Send()
MPI_Bsend()
MPI_Ssend()
MPI_Rsend()
MPI_Recv()
MPI_Sendrecv()
MPI_Sendrecv()
```

Nonblocking Routines

```
MPI_Isend()
MPI_Ibsend()
MPI_Issend()
MPI_Irsend()
MPI_Irecv()
```

Communication Buffer Allocation

```
MPI_Buffer_attach()
MPI_Buffer_detach()
```

Status Data Structure

```
MPI_Get_count()
MPI_Get_elements()
```

Persistent (Half-Channel) Communication

```
MPI_Send_init()
MPI_Bsend_init()
MPI_Rsend_init()
MPI_Ssend_init()
MPI_Recv_init()
MPI_Start()
MPI_Startall()
```

Completion Tests

```
MPI_Wait()
MPI_Waitany()
MPI_Waitsome()
MPI_Waitall()
MPI_Test()
MPI_Testany()
MPI_Testsome()
MPI_Testall()
MPI_Testall()
MPI_Request_free()
MPI_Cancel()
MPI_Test_cancelled()
```

Probing for Messages (Blocking and Nonblocking)

```
MPI_Probe()
MPI_Iprobe()
```

Packing and Unpacking Functions

```
MPI_Pack()
MPI_Unpack()
MPI_Pack_size()
MPI_Pack_external()
MPI_Unpack_external()
MPI_Pack_external_size()
```

Derived Data Type Constructors and Functions

```
MPI_Address(): Deprecated - Use MPI_Get_address()
MPI_Type_commit()
MPI_Type_contiguous()
MPI_Type_create_f90_complex()
MPI_Type_create_f90_integer()
MPI_Type_create_F90_real()
MPI_Type_match_size()
MPI_Type_Sizeof()
MPI_Type_create_indexed_block()
MPI Type create keyval()
MPI_Type_create_resized()
MPI_Type_delete_attr()
MPI_Type_dup()
MPI_Type_free()
MPI_Type_free_keyval()
MPI_Type_get_attr()
MPI_Type_set_attr()
MPI_Type_get_contents()
MPI_Type_get_envelope()
MPI_Type_get_name()
MPI_Type_get_true_extent()
MPI_Type_set_name()
MPI_Type_hvector(): Deprecated - Use MPI_Type_create_hvector()
MPI_Type_indexed()
MPI_Type_hindexed(): Deprecated - Use MPI_Type_create_hindexed()
MPI_Type_struct(): Deprecated - Use MPI_Type_create_struct()
MPI_Type_lb(): Deprecated - Use MPI_Type_get_extent()
MPI_Type_ub(): Deprecated - Use MPI_Type_get_extent()
MPI_Type_vector()
MPI_Type_extent(): Deprecated - Use MPI_Type_get_extent()
MPI_Type_size()
```

One-Sided Communication

Initialization

```
MPI_Win_create()
MPI_Win_free()
MPI_Win_get_group()
```

Communication Calls

```
MPI_Put()
MPI_Get()
MPI_Accumulate()
```

Synchronization Calls

```
MPI_Win_fence()
MPI_Win_lock()
MPI_Win_unlock()
MPI_Win_start()
MPI_Win_complete()
MPI_Win_post()
MPI_Win_wait()
MPI_Win_test()
```

Collective Communication

Barrier

```
MPI_Barrier()
```

Broadcast

```
MPI_Bcast()
```

Processor Gather and Scatter

```
MPI_Gather()
MPI_Gatherv()
MPI_Allgather()
MPI_Allgatherv()
MPI_Scatter()
MPI_Scatterv()
MPI_Alltoall()
MPI_Alltoallv()
MPI_Alltoallw()
```

Global Reduction and Scan Operations

```
MPI_Reduce()
MPI_Allreduce()
MPI_Reduce_scatter()
MPI_Scan()
MPI Exscan()
MPI_Op_create()
MPI_Op_free()
```

Groups and Communicators

Group Management

Group Accessors

```
MPI_Group_size()
MPI_Group_rank()
MPI_Group_translate_ranks()
MPI_Group_compare()
```

Group Constructors

```
MPI_Comm_group()
MPI_Group_union()
MPI_Group_intersection()
MPI_Group_difference()
MPI_Group_incl()
MPI_Group_excl()
MPI_Group_range_incl()
MPI_Group_range_excl()
MPI_Group_free()
```

Communicator Management

Communicator Accessors

```
MPI_Comm_size()
MPI_Comm_rank()
MPI_Comm_compare()
```

Communicator Constructors

```
MPI_Comm_dup()
MPI_Comm_create()
MPI_Comm_split()
MPI_Comm_free()
```

Intercommunicators

```
MPI_Comm_test_inter()
MPI_Comm_remote_group()
MPI_Comm_remote_size()
MPI_Intercomm_create()
MPI_Intercomm_merge()
```

Communicator Attributes

```
MPI_Keyval_create(): Deprecated - Use MPI_Comm_create_keyval()
MPI_Keyval_free(): Deprecated - Use MPI_Comm_free_keyval()
MPI_Attr_put(): Deprecated - Use MPI_Comm_set_attr()
MPI_Attr_get(): Deprecated - Use MPI_Comm_get_attr()
MPI_Attr_delete(): Deprecated - Use MPI_Comm_delete_attr()
```

Process Topologies

```
MPI_Cart_create()
MPI_Dims_create()
MPI_Graph_create()
MPI_Topo_test()
MPI_Graphdims_get()
MPI_Graph_get()
MPI_Cartdim_get()
MPI_Cart_get()
MPI_Cart_rank()
MPI_Cart_coords()
MPI_Graph_neighbors()
MPI_Graph_neighbors_count()
MPI_Cart_shift()
MPI_Cart_sub()
MPI_Cart_map()
MPI_Graph_map()
```

Process Creation and Management

Establishing Communication

```
MPI_Close_port()
MPI_Comm_accept()
MPI_Comm_connect()
MPI_Comm_disconnect()
MPI_Open_port()
MPI_Comm_join()
```

Name Publishing

```
MPI_Publish_name()
MPI_Unpublish_name()
MPI_Lookup_name()
```

Process Manager Interface

```
MPI_Comm_get_parent()
MPI_Comm_spawn()
MPI_Comm_spawn_multiple()
```

Environmental Inquiry Functions and Profiling

Startup and Shutdown

```
MPI_Init()
MPI_Finalize()
MPI_Finalized()
MPI_Initialized()
MPI_Abort()
MPI_Get_processor_name()
MPI_Get_version()
```

Error Handler Functions

```
MPI_Add_error_class()
MPI_Add_error_code()
MPI_Add_error_string()
MPI_Comm_call_errhandler()
MPI_File_call_errhandler()
MPI_Win_call_errhandler()
MPI_Errhandler_create(): Deprecated - Use MPI_Comm_create_errhandler()
MPI_Errhandler_set(): Deprecated - Use MPI_Comm_set_errhandler()
MPI_Errhandler_get(): Deprecated - Use MPI_Comm_get_errhandler()
MPI_Errhandler_free()
MPI_Error_string()
MPI_Error_class()
```

Info Objects

```
MPI_Info_create()
MPI_Info_delete()
MPI_Info_dup()
MPI_Info_free()
MPI_Info_get()
MPI_Info_get_nkeys()
MPI_Info_get_nthkey()
MPI_Info_get_valuelen()
MPI_Info_set()
```

Timers

```
MPI_Wtime()
MPI_Wtick()
```

Profiling

```
MPI_Pcontrol()
```

Miscellaneous

Associating Information With Status

```
MPI_Status_set_cancelled()
MPI_Status_set_elements()
```

Generalized Requests

```
MPI_Grequest_complete()
MPI_Grequest_start()
```

Naming Objects

```
MPI_Comm_get_name()
MPI_Comm_set_name()
MPI_Type_get_name()
MPI_Type_set_name()
```

Threads

```
MPI_Query_thread()
MPI_Init_thread()
MPI_Is_thread_main()
```

Handle Translation

```
MPI_Comm_c2f()
MPI_Comm_f2c()
MPI_Group_c2f()
MPI_Group_f2c()
MPI_Info_c2f()
MPI_Info_f2c()
MPI_Op_c2f()
MPI_Op_f2c()
MPI_Request_c2f()
MPI_Request_f2c()
MPI_Type_c2f()
MPI_Type_f2c()
```

Status Conversion

```
MPI_Status_c2f()
MPI_Status_f2c()
```

MPI Routines: Alphabetical Listing

TABLE A-1 Sun MPI Routines

Routine and C Syntax	Description
MPI_Abort(MPI_Comm comm, int errorcode)	Terminates MPI execution environment.
MPI_Accumulate(void *origin_addr, int origin_count, MPI_Datatype origin_datatype, int target_rank, MPI_Aint target_disp, int target_count, MPI_Datatype target_datatype, MPI_Op op, MPI_Win win)	Combines the contents of the origin buffer with that of a target buffer.
<pre>MPI_Add_error_class(int *errorclass)</pre>	Creates a new, local error class.
<pre>MPI_Add_error_code(int errorclass, int *errorcode)</pre>	Creates a new error code and associates it with an error class.
<pre>MPI_Add_error_string(int *errorcode, char *string)</pre>	Creates an error string and associates it with an error code or an error class.

 TABLE A-1
 Sun MPI Routines (Continued)

Routine and C Syntax	Description
MPI_Address(void *location, MPI_Aint *address)	Deprecated: Use instead MPI_Get_address(). Gets the address of a location in memory.
MPI_Allgather(void *sendbuf, int sendcount, MPI_Datatype sendtype, void *recvbuf, int recvcount, MPI_Datatype recvtype, MPI_Comm comm)	Gathers data from all processes and distributes it to all.
MPI_Allgatherv(void *sendbuf, int sendcount, MPI_Datatype sendtype, void *recvbuf, int *recvcount, int *displs, MPI_Datatype recvtype, MPI_Comm comm)	Gathers data from all processes and delivers it to all. Each process can contribute a different amount of data.
$\label{eq:mpi_alloc_mem} \begin{split} &\texttt{MPI_Alloc_mem}(\texttt{MPI_Aint} \textit{ size}, \texttt{MPI_Info} \textit{ info}, \texttt{void} \\ &*\textit{baseptr}) \end{split}$	Allocates a specified memory segment.
MPI_Allreduce(void *sendbuf, void *recvbuf, int count, MPI_Datatype datatype, MPI_Op op, MPI_Comm comm)	Combines values from all processes and distributes the result back to all processes.
MPI_Alltoall(void *sendbuf, int sendcount, MPI_Datatype sendtype, void *recvbuf, int recvcount, MPI_Datatype recvtype, MPI_Comm comm)	All processes send data to, and receive data from, all processes.
MPI_Alltoallv(void *sendbuf, int *sendcounts, int *sdispls, MPI_Datatype sendtype, void *recvbuf, int *recvcounts, int *rdispls, MPI_Datatype recvtype, MPI_Comm comm)	All processes send data to, and receive data from, all processes, but user can specify different amounts of data to send and receive.
MPI_Alltoallw(void *sendbuf, int *sendcounts, int *sdispls, MPI_Datatype sendtypes, void *recvbuf, int *recvcounts, int *rdispls, MPI_Datatype recvtypes, MPI_Comm comm)	All processes send data to, and receive data from, all other processes, and user can specify database of individual datablocks of different types.
<pre>MPI_Attr_delete(MPI_Comm comm, int keyval)</pre>	Deprecated: Use instead MPI_Comm_delete_attr(). Deletes attribute value associated with a key.
MPI_Attr_get(MPI_Comm comm, int keyval, void *attribute_val, int *flag)	Deprecated: Use instead MPI_Comm_get_attr(). Retrieves attribute value by key.
MPI_Attr_put(MPI_Comm comm, int keyval, void *attribute_val)	Deprecated: Use instead MPI_Comm_set_attr(). Stores attribute value associated with a key.
<pre>MPI_Barrier(MPI_Comm comm)</pre>	Blocks until all processes have reached this routine.

 TABLE A-1
 Sun MPI Routines (Continued)

Routine and C Syntax	Description
MPI_Bcast(void *buffer, int count, MPI_Datatype datatype, int root, MPI_Comm comm)	Broadcasts a message from the process with rank root to all other processes of the group.
MPI_Bsend(void *buf, int count, MPI_Datatype datatype, int dest, int tag, MPI_Comm comm)	Basic send with user-specified buffering.
MPI_Bsend_init(void *buf, int count, MPI_Datatype datatype, int dest, int tag, MPI_Comm comm, MPI_Request *request)	Builds a handle for a buffered send.
MPI_Buffer_attach(void *buf, int size)	Attaches a user-defined buffer for sending.
MPI_Buffer_detach(void *buf, int *size)	Removes an existing buffer (for use in MPI_Bsend(), etc.).
MPI_Cancel(MPI_Request *request)	Cancels a communication request.
MPI_Cart_coords(MPI_Comm comm, int rank, int maxdims, int *coords)	Determines process coordinates in Cartesian topology given rank in group.
<pre>MPI_Cart_create(MPI_Comm comm_old, int ndims, int *dims, int *periods, int reorder, MPI_Comm *comm_cart)</pre>	Makes a new communicator to which Cartesian topology information has been attached.
<pre>MPI_Cart_get(MPI_Comm comm, int maxdims, int *dims, int *periods, int *coords)</pre>	Retrieves Cartesian topology information associated with a communicator.
MPI_Cart_map(MPI_Comm comm, int ndims, int *dims, int *periods, int *newrank)	Maps process to Cartesian topology information.
<pre>MPI_Cart_rank(MPI_Comm comm, int *coords, int *rank)</pre>	Determines process rank in communicator given Cartesian location.
<pre>MPI_Cart_shift(MPI_Comm comm, int direction, int disp, int *rank_source, int *rank_dest)</pre>	Returns the shifted source and destination ranks, given a shift direction and amount.
MPI_Cart_sub(MPI_Comm comm, int *remain_dims, MPI_Comm *comm_new)	Partitions a communicator into subcommunicators that form lower-dimensional Cartesian subgrids.
<pre>MPI_Cartdim_get(MPI_Comm comm, int *ndims)</pre>	Retrieves Cartesian topology information associated with a communicator.
<pre>MPI_Close_port(char *port_name)</pre>	Releases the specified network address.

 TABLE A-1
 Sun MPI Routines (Continued)

Routine and C Syntax	Description
MPI_Comm_accept(char *port_name, MPI_Info info, int root, MPI_Comm comm, MPI_Comm *newcomm)	Establishes communication with a client (collective).
MPI_Comm_c2f(MPI_Comm comm)	Translates a C handle into a Fortran handle.
MPI_Comm_compare(MPI_Comm comm1, MPI_Comm comm2, int *result)	Compares two communicators.
MPI_Comm_connect(char *port_name, MPI_Info info, int root, MPI_Comm comm, MPI_Comm *newcomm)	Establishes communication with a server (collective).
$\label{eq:mpi_comm_create} \begin{split} \texttt{MPI_Comm} & \ \textit{comm}, \ \texttt{MPI_Group} \\ & \textit{group}, \ \texttt{MPI_Comm} \ *\textit{newcomm}) \end{split}$	Creates a new communicator from a group.
MPI_Comm_create_errhandler(MPI_Comm_errhandler_fn *function, MPI_Errhandler *errhandler)	Creates an error handler that can be attached to communicators.
MPI_Comm_create_keyval(MPI_Comm_copy_attr_function *comm_copy_attr_fn, MPI_Comm_delete_attr_function *comm_delete_attr_fn, int *comm_keyval, void *extra_state)	Generates a new attribute key.
<pre>MPI_Comm_delete_attr(MPI_Comm comm, int comm_keyval)</pre>	Deletes attribute value associated with a key.
<pre>MPI_Comm_disconnect(MPI_Comm *comm)</pre>	De-allocates communicator object and sets handle to MPI_COMM_NULL (collective).
<pre>MPI_Comm_dup(MPI_Comm comm, MPI_Comm *newcomm)</pre>	Duplicates an existing communicator with all its cached information.
MPI_Comm_f2c(MPI_Fint comm)	Translates a Fortran handle into a C handle.
<pre>MPI_Comm_free(MPI_Comm *comm)</pre>	Marks the communicator object for de-allocation.
<pre>MPI_Comm_free_keyval(int *comm_keyval)</pre>	Frees attribute key for communicator cache attribute.
MPI_Comm_get_attr(MPI_Comm comm, int comm_keyval, void *attribute_val, int *flag)	Retrieves attribute value by key.
MPI_Comm_get_errhandler(MPI_Comm comm, MPI_Errhandler *errhandler)	Retrieves error handler associated with a communicator.

 TABLE A-1
 Sun MPI Routines (Continued)

Routine and C Syntax	Description
MPI_Comm_get_name(MPI_Comm comm, char *comm_name, int *resultlen)	Returns the name that was most recently associated with a communicator.
<pre>MPI_Comm_get_parent(MPI_Comm *parent)</pre>	Returns the parent intercommunicator of current spawned process.
MPI_Comm_group(MPI_Comm comm, MPI_Group *group)	Accesses the group associated with a communicator.
<pre>MPI_Comm_join(int fd, MPI_Comm *intercomm)</pre>	Creates an intercommunicator from the union of two MPI processes connected by a socket.
<pre>MPI_Comm_rank(MPI_Comm comm, int *rank)</pre>	Determines the rank of the calling process in a communicator.
<pre>MPI_Comm_remote_group(MPI_Comm comm, MPI_Group *group)</pre>	Accesses the remote group associated with an intercommunicator.
<pre>MPI_Comm_remote_size(MPI_Comm comm, int size)</pre>	Determines the size of the remote group associated with an intercommunicator.
<pre>MPI_Comm_set_attr(MPI_Comm comm, int comm_keyval, void *attribute_val)</pre>	Stores attribute value associated with a key.
<pre>MPI_Comm_set_errhandler(MPI_Comm comm, MPI_Errhandler *errhandler)</pre>	Attaches a new error handler to a communicator.
<pre>MPI_Comm_set_name(MPI_Comm comm, char *comm_name)</pre>	Associates a name with a communicator.
<pre>MPI_Comm_size(MPI_Comm comm, int *size)</pre>	Determines the size of the group associated with a communicator.
MPI_Comm_spawn(char *command, char *argv[], int maxprocs, MPI_Info info, int root, MPI_Comm comm, MPI_Comm *intercomm, int array_of_errcodes[])	Spawns a number of identical binaries.
MPI_Comm_spawn_multiple(int count, char *array_of_commands[], char **array_of_argv[], int array_of_maxprocs[], MPI_Info array_of_info[], int root, MPI_Comm comm, MPI_Comm *intercomm, int array_of_errcodes[])	Spawns multiple binaries, or the same binary with multiple sets of arguments.
<pre>MPI_Comm_split(MPI_Comm comm, int color, int key, MPI_Comm *newcomm)</pre>	Creates new communicators based on colors and keys.
<pre>MPI_Comm_test_inter(MPI_Comm comm, int *flag)</pre>	Tests whether a communicator is an intercommunicator.

 TABLE A-1
 Sun MPI Routines (Continued)

Routine and C Syntax	Description
MPI_Dims_create(int <i>nnodes</i> , int <i>ndims</i> , int *dims)	Creates a division of processors in a Cartesian grid.
MPI_Errhandler_create(MPI_Handler_function *function, MPI_Errhandler *errhandler)	Deprecated: Use instead MPI_Comm_create_errhandler (). Creates an MPI error handler.
<pre>MPI_Errhandler_free(MPI_Errhandler *errhandler)</pre>	Frees an MPI error handler.
MPI_Errhandler_get(MPI_Comm comm, MPI_Errhandler *errhandler)	Deprecated: Use instead MPI_Comm_get_errhandler(). Gets the error handler for a communicator.
MPI_Errhandler_set(MPI_Comm comm, MPI_Errhandler errhandler)	Deprecated: Use instead MPI_Comm_set_errhandler(). Sets the error handler for a communicator.
<pre>MPI_Error_class(int errorcode, int *errorclass)</pre>	Converts an error code into an error class.
<pre>MPI_Error_string(int errorcode, char *string, int *resultlen)</pre>	Returns a string for a given error code.
MPI_Exscan(void *sendbuf, void *recvbuf, int count, MPI_Datatype datatype, MPI_Op op, MPI_Comm, comm)	Performs an exclusive prefix reduction on data distributed across the calling processes.
MPI_Finalize()	Terminates MPI execution environment.
MPI_Finalized(int *flag)	Checks whether $\mathtt{MPI_Finalize}()$ has completed.
MPI_Free_mem(void *base)	Frees memory that has been allocated using MPI_Alloc_mem.
MPI_Gather(void *sendbuf, int *sendcount, MPI_Datatype sendtype, void *recvbuf, int recvcount, MPI_Datatype recvtype, int root, MPI_Comm comm)	Gathers values from a group of processes.
MPI_Gatherv(void *sendbuf, int sendcount, MPI_Datatype sendtype, void *recvbuf, int *recvcounts, int *displs, MPI_Datatype recvtype, int root, MPI_Comm comm)	Gathers into specified locations from all processes in a group. Each process can contribute a different amount of data.
MPI_Get(void *origin_addr, int origin_count, MPI_Datatype origin_datatype, int target_rank, MPI_Aint target_disp, int target_count, MPI_Datatype target_datatype, MPI_Win win)	Copies data from the target memory to the origin.

 TABLE A-1
 Sun MPI Routines (Continued)

Routine and C Syntax	Description
<pre>MPI_Get_address(void *location, MPI_Aint *address)</pre>	Gets the address of a location in memory.
<pre>MPI_Get_count(MPI_Status *status, MPI_Datatype datatype, int *count)</pre>	Gets the number of top-level elements received.
MPI_Get_elements(MPI_Status *status, MPI_Datatype datatype, int *count)	Returns the number of basic elements in a data type.
<pre>MPI_Get_processor_name(char *name, int *resultlen)</pre>	Gets the name of the processor.
<pre>MPI_Get_version(int *version, int *subversion)</pre>	Returns the version of the standard corresponding to the current implementation.
MPI_Graph_create(MPI_Comm comm_old, int nnodes, int *index, int *edges, int reorder, MPI_Comm *comm_graph)	Makes a new communicator to which graph topology information has been attached.
<pre>MPI_Graph_get(MPI_Comm comm, int maxindex, int maxedges, int *index, int *edges)</pre>	Retrieves graph topology information associated with a communicator.
<pre>MPI_Graph_map(MPI_Comm comm, int nnodes, int *index, int *edges, int *newrank)</pre>	Maps process to graph topology information.
<pre>MPI_Graph_neighbors(MPI_Comm comm, int rank, int maxneighbors, int *neighbors)</pre>	Returns the neighbors of a node associated with a graph topology.
<pre>MPI_Graph_neighbors_count(MPI_Comm comm, int rank, int *nneighbors)</pre>	Returns the number of neighbors of a node associated with a graph topology.
<pre>MPI_Graphdims_get(MPI_Comm comm, int *nnodes, int *nedges)</pre>	Retrieves graph topology information associated with a communicator.
<pre>MPI_Grequest_complete(MPI_Request request)</pre>	Reports that a generalized request is complete.
MPI_Grequest_start(MPI_Grequest_query_function *query_fn, MPI_Grequest_free_function *free_fn, MPI_Grequest_cancel_function *cancel_fn, void *extra_state, MPI_Request *request)	Starts a generalized request and returns a handle to it.
<pre>MPI_Group_c2f(MPI_Group group)</pre>	Translates a C handle into a Fortran handle.
<pre>MPI_Group_compare(MPI_Group group1, MPI_Group group2, int *result)</pre>	Compares two groups.

 TABLE A-1
 Sun MPI Routines (Continued)

Routine and C Syntax	Description
MPI_Group_difference(MPI_Group group1, MPI_Group group2, MPI_Group *group_out)	Makes a group from the difference of two groups.
MPI_Group_excl(MPI_Group group, int n, int *ranks, MPI_Group *newgroup)	Produces a group by reordering an existing group and taking only unlisted members.
MPI_Group_f2c(MPI_Fint group)	Translates a Fortran handle into a C handle.
MPI_Group_free(MPI_Group *group)	Frees a group.
MPI_Group_incl(MPI_Group group, int n, int *ranks, MPI_Group *group_out)	Produces a group by reordering an existing group and taking only listed members.
<pre>MPI_Group_intersection(MPI_Group group1, MPI_Group group2, MPI_Group *group_out)</pre>	Produces a group at the intersection of two existing groups.
MPI_Group_range_excl(MPI_Group group, int n, int ranges[][3], MPI_Group *newgroup)	Produces a group by excluding ranges of processes from an existing group.
MPI_Group_range_incl(MPI_Group group, int n, int ranges[][3], MPI_Group *newgroup)	Creates a new group from ranges of ranks in an existing group.
MPI_Group_rank(MPI_Group group, int *rank)	Returns the rank of this process in the given group.
MPI_Group_size(MPI_Group group, int *size)	Returns the size of a group.
MPI_Group_translate_ranks(MPI_Group group1, int n, int *ranks1, MPI_Group group2, int *ranks2)	Translates the ranks of processes in one group to those in another group.
MPI_Group_union(MPI_Group group1, MPI_Group group2, MPI_Group *group_out)	Produces a group by combining two groups.
MPI_Ibsend(void *buf, int count, MPI_Datatype datatype, int dest, int tag, MPI_Comm comm, MPI_Request *request)	Starts a nonblocking buffered send.
MPI_Info_c2f(MPI_Info info)	Translates a C handle into a Fortran handle.
MPI_Info_create(MPI_Info *info)	Creates a new info object.
MPI_Info_delete(MPI_Info *info, char *key, char *value)	Deletes a key/value pair from info.
MPI_Info_dup(MPI_Info info, MPI_Info *newinfo)	Duplicates an info object.
MPI_Info_f2c(MPI_Fint info)	Translates a Fortran handle into a C handle.

 TABLE A-1
 Sun MPI Routines (Continued)

Routine and C Syntax	Description
MPI_Info_free(MPI_Info *info)	Frees <i>info</i> and sets it to MPI_INFO_NULL.
<pre>MPI_Info_get(MPI_Info *info, char *key, char *value)</pre>	Retrieves key value for an <i>info</i> object.
<pre>MPI_Info_get_nkeys(MPI_Info info, int *nkeys)</pre>	Returns the number of currently defined keys in info.
$\label{eq:mpi_mpi_mpi_mpi_mpi_n} \begin{split} \texttt{MPI_Info_get_nthkey}(\texttt{MPI_Info} \ \textit{info}, \ \text{int} \ \textit{n}, \ \text{char} \\ *\textit{key}) \end{split}$	Returns the <i>n</i> th defined key in <i>info</i> .
<pre>MPI_Info_get_valuelen(MPI_Info info, char *key, int *valuelen, int *flag)</pre>	Retrieves the length of the key value associated with an <i>info</i> object.
<pre>MPI_Info_set(MPI_Info *info, char *key, char *value)</pre>	Adds a key/value pair to info.
<pre>MPI_Init(int *argc, char ***argv)</pre>	Initializes the MPI execution environment.
<pre>MPI_Initialized(int *flag)</pre>	Indicates whether ${\tt MPI_Init()}$ has been called.
<pre>MPI_Init_thread(int *argc, char ***argv, int required, int *provided)</pre>	Initializes the MPI execution environment with a predetermined level of thread support. Thread that calls this function becomes the main thread. Call instead of MPI_Init(), not in addition to.
<pre>MPI_Intercomm_create(MPI_Comm local_comm, int local_leader, MPI_Comm peer_comm, int remote_leader, int tag, MPI_Comm *newintercomm)</pre>	Creates an intercommunicator.
<pre>MPI_Intercomm_merge(MPI_Comm intercomm, int high, MPI_Comm *newintracomm</pre>	Creates an intracommunicator from an intercommunicator.
<pre>MPI_Iprobe(int source, int tag, MPI_Comm comm, int *flag, MPI_Status *status)</pre>	Nonblocking test for a message.
MPI_Irecv(void *buf, int count, MPI_Datatype datatype, int source, int tag, MPI_Comm comm, MPI_Request *request)	Begins a nonblocking receive.
MPI_Irsend(void *buf, int count, MPI_Datatype datatype, int dest, int tag, MPI_Comm comm, MPI_Request *request)	Begins a nonblocking ready send.
MPI_Isend(void *buf, int count, MPI_Datatype datatype, int dest, int tag, MPI_Comm comm, MPI_Request *request)	Begins a nonblocking send.

 TABLE A-1
 Sun MPI Routines (Continued)

Routine and C Syntax	Description
MPI_Issend(void *buf, int count, MPI_Datatype datatype, int dest, int tag, MPI_Comm comm, MPI_Request *request)	Begins a nonblocking synchronous send.
${\tt MPI_Is_thread_main} (\text{int } * \textit{flag})$	Called by a thread to determine whether it is the main thread. See MPI_Init_thread().
MPI_Keyval_create(MPI_Copy_function *copy_fn, MPI_Delete_function *delete_fn, int *keyval, void *extra_state)	Deprecated: Use instead MPI_Comm_create_keyval(). Generates a new attribute key.
MPI_Keyval_free(int *keyval)	Deprecated: Use instead MPI_Comm_free_keyval(). Frees attribute key for communicator cache attribute.
MPI_Lookup_name (char *service-name, MPI_Info info, char *port-name)	Retrieves the port name associated with a <i>service-name</i> published by MPI_Publish_name().
MPI_Op_c2f(MPI_Op op)	Translates a C handle into a Fortran handle.
MPI_Op_create(MPI_User_function *function, int commute, MPI_Op *op)	Creates a user-defined combination function handle.
MPI_Op_f2c(MPI_Fint op)	Translates a Fortran handle into a C handle.
MPI_Op_free(MPI_Op *op)	Frees a user-defined combination function handle.
MPI_Open_port(MPI_Info info, char *port_name)	Establishes a network address for a server to accept connections from clients.
MPI_Pack(void *inbuf, int incount, MPI_Datatype datatype, void *outbuf, int outsize, int *position, MPI_Comm comm)	Packs data of a given data type into contiguous memory.
MPI_Pack_external(char *datarep, void *inbuf, int incount, MPI_Datatype datatype, void *outbuf, MPI_Aint *position)	Packs data into the external32 format, used to exchange data between MPI implementations, or when writing data to a file.
MPI_Pack_external_size(char *datarep int incount, MPI_Datatype datatype, MPI_Aint *size)	Returns the upper bound on the amount of space necessary to pack a message in the external32 format.
MPI_Pack_size(int incount, MPI_Datatype datatype, MPI_Comm comm, int *size)	Returns the upper bound on the amount of space necessary to pack a message.

 TABLE A-1
 Sun MPI Routines (Continued)

Routine and C Syntax	Description
MPI_Pcontrol(int level,)	Controls profiling.
MPI_Probe(int source, int tag, MPI_Comm comm, MPI_Status *status)	Blocking test for a message.
<pre>MPI_Publish_name (char *service-name, MPI_Info info, char *port-name)</pre>	Publishes the pair (service-name, port-name) so that applications can use MPI_Lookup_name() to find port-name.
MPI_Put(void *origin_addr, int origin_count, MPI_Datatype origin_datatype, int target_rank, MPI_Aint target_disp, int target_count, MPI_Datatype target_datatype, MPI_Win win)	Copies data from the origin memory to the target.
<pre>MPI_Query_thread(int *provided)</pre>	Returns the current level of thread support.
MPI_Recv(void *buf, int count, MPI_Datatype datatype, int source, int tag, MPI_Comm comm, MPI_Status *status)	Performs a standard receive.
<pre>MPI_Recv_init(void *buf, int count, MPI_Datatype datatype, int source, int tag, MPI_Comm comm, MPI_Request *request)</pre>	Builds a persistent receive request handle.
MPI_Reduce(void *sendbuf, void *recvbuf, int count, MPI_Datatype datatype, MPI_Op op, int root, MPI_Comm comm)	Reduces values on all processes to a single value.
MPI_Reduce_scatter(void *sendbuf, void *recvbuf, int *recvcounts, MPI_Datatype datatype, MPI_Op op, MPI_Comm comm)	Combines values and scatters the results.
<pre>MPI_Request_c2f(MPI_Request request)</pre>	Translates a C handle into a Fortran handle.
<pre>MPI_Request_f2c(MPI_Fint request)</pre>	Translates a Fortran handle into a C handle.
<pre>MPI_Request_free(MPI_Request *request)</pre>	Frees a communication request object.
<pre>MPI_Request_get_status(MPI_Request request, int *flag, MPI_Status *status)</pre>	Accesses information associated with a request without freeing the request.
MPI_Rsend(void *buf, int count, MPI_Datatype datatype, int dest, int tag, MPI_Comm comm)	Performs a ready send.
MPI_Rsend_init(void *buf, int count, MPI_Datatype datatype, int dest, int tag, MPI_Comm comm, MPI_Request *request)	Builds a persistent ready send request handle.

 TABLE A-1
 Sun MPI Routines (Continued)

Routine and C Syntax	Description
MPI_Scan(void *sendbuf, void *recvbuf, int count, MPI_Datatype datatype, MPI_Op op, MPI_Comm comm)	Computes the scan (partial reductions) of data on a collection of processes.
MPI_Scatter(void *sendbuf, int sendcount, MPI_Datatype sendtype, void *recvbuf, int recvcount, MPI_Datatype recvtype, int root, MPI_Comm comm)	Sends data from one job to all other processes in a group.
MPI_Scatterv(void *sendbuf, int *sendcounts, int *displs, MPI_Datatype sendtype, void *recvbuf, int recvcount, MPI_Datatype recvtype, int root, MPI_Comm comm)	Scatters a buffer in parts to all processes in a group.
MPI_Send(int *buf, int count, MPI_Datatype datatype, int dest, int tag, MPI_Comm comm)	Performs a standard send.
MPI_Send_init(void *buf, int count, MPI_Datatype datatype, int dest, int tag, MPI_Comm comm, MPI_Request *request)	Builds a persistent send request handle.
MPI_Sendrecv(void *sendbuf, int sendcount, MPI_Datatype sendtype, int dest, int sendtag, void *recvbuf, int recvcount, MPI_Datatype recvtype, int source, int recvtag, MPI_Comm comm, MPI_Status *status)	Sends and receives two messages at the same time.
MPI_Sendrecv_replace(void *buf, int count, MPI_Datatype datatype, int dest, int sendtag, int source, int recvtag, MPI_Comm comm, MPI_Status *status)	Sends and receives using a single buffer.
<pre>MPI_Size_of(x, size, ierror)</pre>	Fortran only. Returns the size, in bytes, of the datatype of variable x .
MPI_Ssend(void *buf, int count, MPI_Datatype datatype, int dest, int tag, MPI_Comm comm)	Performs a synchronous send.
MPI_Ssend_init(void *buf, int count, MPI_Datatype datatype, int dest, int tag, MPI_Comm comm, MPI_Request *request)	Builds a persistent synchronous send request handle.
<pre>MPI_Start(MPI_Request *request)</pre>	Initiates a communication using a persistent request handle.
MPI_Startall(int count, MPI_Request array_of_requests[])	Starts a collection of requests.
MPI_Status_c2f(MPI_Status *c_status, MPI_Fint *f_status)	Translates a C status into a Fortran status.
MPI_Status_f2c(MPI_Fint *f_status, MPI_Status *c_status)	Translates a Fortran status into a C status.

 TABLE A-1
 Sun MPI Routines (Continued)

Routine and C Syntax	Description
<pre>MPI_Status_set_cancelled(MPI_Status*status, int flag)</pre>	Sets <i>status</i> to indicate that a request has been cancelled.
MPI_Status_set_elements(MPI_Status *status, MPI_Datatype datatype, int count)	Modifies opaque part of <i>status</i> to enable MPI_Get_elements() to return <i>count</i> .
MPI_Test(MPI_Request *request, int *flag, MPI_Status *status)	Tests for the completion of a send or receive.
MPI_Test_cancelled(MPI_Status *status, int *flag)	Tests whether a request was canceled.
MPI_Testall(int <i>count</i> , MPI_Request array_of_requests, int *flag, MPI_Status *array_of_statuses)	Tests for the completion of all the given communications.
MPI_Testany(int count, MPI_Request array_of_requests[], int *index, int *flag, MPI_Status status)	Tests for completion of any of the given communications.
MPI_Testsome(int incount, MPI_Request array_of_requests[], int *outcount, int *array_of_indices, MPI_Status *array_of_statuses)	Tests for some given communications to complete.
MPI_Topo_test(MPI_Comm comm, int *top_type)	Determines the type of topology (if any) associated with a communicator.
MPI_Type_c2f(MPI_Datatype datatype)	Translates a C handle into a Fortran handle.
MPI_Type_commit(MPI_Datatype *datatype)	Commits a data type.
MPI_Type_contiguous(int <i>count</i> , MPI_Datatype <i>oldtype</i> , MPI_Datatype * <i>newtype</i>)	Creates a contiguous data type.
MPI_Type_create_darray(int size, int rank, int ndims, int array_of_gsizes[], int array_of_distribs[], int array_of_dargs[], int array_of_psizes[], int order, MPI_Datatype oldtype, MPI_Datatype *newtype)	Creates an array of data types.
MPI_Type_create_f90_complex(int p, int r, MPI_Datatype *newtype)	Returns a bounded MPI complex datatype.
MPI_Type_create_f90_integer(int r, MPI_Datatype *newtype)	Returns a bounded MPI integer datatype.
MPI_Type_create_f90_complex(int p, int r; MPI_Datatype *newtype)	Returns a bounded MPI real datatype.

 TABLE A-1
 Sun MPI Routines (Continued)

Routine and C Syntax	Description
MPI_Type_create_hindexed(int count, int array_of_blocklengths, MPI_Aint array_of_displacements[], MPI_Datatype oldtype, MPI_Datatype *newtype)	Creates an indexed data type with offsets in bytes.
MPI_Type_create_hvector(int count, int blocklength, MPI_Aint stride, MPI_Datatype oldtype, MPI_Datatype *newtype)	Creates a vector (strided) data type with offset in bytes.
MPI_Type_create_indexed_block(int count, int blocklength, int array_of_displacements[], MPI_Datatype oldtype, MPI_Datatype *newtype)	Creates an indexed block.
MPI_Type_create_keyval(MPI_Type_copy_attr_function *type_copy_attr_fn, MPI_Type_delete_attr_function *type_delete_attr_fn, int *type_keyval, void *extra_state)	Generates a new attribute key.
MPI_Type_create_resized(MPI_Datatype oldtype, MPI_Aint lb, MPI_Aint extent, MPI_Datatype *newtype)	Returns a new data type with new extent and upper and lower bounds.
MPI_Type_create_struct(int count, int array_of_blocklengths[], MPI_Aint array_of_displacements[], MPI_Datatype array_of_types[], MPI_Datatype *newtype)	Creates a struct data type.
MPI_Type_create_subarray(int ndims, int array_of_sizes[], int array_of_subsizes[], int array_of_starts[], int order, MPI_Datatype oldtype, MPI_Datatype *newtype)	Creates a data type describing a subarray of an array.
<pre>MPI_Type_delete_attr(MPI_Datatype type, int type_keyval)</pre>	Deletes attribute value associated with a key.
MPI_Type_dup(MPI_Datatype type, MPI_Datatype *newtype)	Duplicates a data type with associated key values.
MPI_Type_extent(MPI_Datatype datatype, MPI_Aint *extent)	Deprecated: Use instead MPI_Type_get_extent(). Returns the extent of a data type, the difference between the upper and lower bounds of the data type.
MPI_Type_f2c(MPI_Fint datatype)	Translates a Fortran handle into a C handle.
MPI_Type_free(MPI_Datatype *datatype)	Frees a data type.
MPI_Type_free_keyval(int *type_keyval)	Frees an attribute key.
MPI_Type_get_attr(MPI_Datatype type, int type_keyval, void *attribute_val, int *flag)	Returns the attribute associated with a data type.

 TABLE A-1
 Sun MPI Routines (Continued)

Routine and C Syntax	Description
MPI_Type_get_contents(MPI_Datatype datatype, int max_integers, int max_addresses, int max_datatypes, int array_of_integers[], MPI_Aint array_of_addresses[], MPI_Datatype array_of_datatypes[])	Returns information about arguments used in creation of a data type.
MPI_Type_get_envelope(MPI_Datatype datatype, int *num_integers, int *num_addresses, int *num_datatypes, int *combiner)	Returns information about input arguments associated with a data type.
MPI_Type_get_extent(MPI_Datatype datatype, MPI_Aint *lb, MPI_Aint *extent)	Returns the lower bound and extent of a data type.
MPI_Type_get_name(MPI_Datatype type, char *type_name, int *resultlen)	Gets the name of a data type.
MPI_Type_get_true_extent(MPI_Datatype datatype, MPI_Aint *true_lb, MPI_Aint *true_extent)	Returns the true lower bound and extent of a data type's corresponding type map, ignoring MPI_UB and MPI_LB markers.
MPI_Type_hindexed(int count, int *array_of_blocklengths, MPI_Aint *array_of_displacements, MPI_Datatype oldtype, MPI_Datatype *newtype)	Deprecated: Use instead MPI_Type_create_hindexed(). Creates an indexed data type with offsets in bytes.
MPI_Type_hvector(int count, int blocklength, MPI_Aint stride, MPI_Datatype oldtype, MPI_Datatype *newtype)	Deprecated: Use instead MPI_Type_create_hvector(). Creates a vector (strided) data type with offset in bytes.
MPI_Type_indexed(int count, int *array_of_blocklengths, int *array_of_displacements, MPI_Datatype oldtype, MPI_Datatype *newtype)	Creates an indexed data type.
MPI_Type_1b(MPI_Datatype datatype, MPI_Aint *displacement)	Deprecated: Use instead MPI_Type_get_extent(). Returns the lower bound of a data type.
MPI_Type_match_size(int typeclass, int size, MPI_Datatype *type)	Returns an MPI data type of a given type and size.
MPI_Type_set_attr(MPI_Datatype type, int type_keyval, void *attribute_val)	Stores attribute value associated with a key.
MPI_Type_set_name(MPI_Comm comm, char *type_name)	Sets the name of a data type.
MPI_Type_size(MPI_Datatype datatype, int *size)	Returns the number of bytes occupied by entries in the data type.

 TABLE A-1
 Sun MPI Routines (Continued)

Routine and C Syntax	Description
***may_of_types, MPI_Datatype ***array_of_types, MPI_Datatype ***ray_of_types, MPI_Datatype ***newtype ***newt	Deprecated: Use instead MPI_Type_create_struct(). Creates a struct data type.
MPI_Type_ub(MPI_Datatype datatype, MPI_Aint *displacement)	Deprecated: Use instead MPI_Type_get_extent(). Returns the upper bound of a data type.
MPI_Type_vector(int count, int blocklength, int stride, MPI_Datatype oldtype, MPI_Datatype *newtype)	Creates a vector (strided) data type.
MPI_Unpack(void *inbuf, int insize, int *position, void *outbuf, int outcount, MPI_Datatype datatype, MPI_Comm comm)	Unpacks a data type into contiguous memory.
MPI_Unpack_external(void *inbuf, int insize, int *position, void *outbuf, int outcount, MPI_Datatype datatype, MPI_Comm comm)	Unpacks into contiguous memory a data type packed in the external32 format.
MPI_Unpublish_name (char *service-name, MPI_Info info, char *port-name)	Removes the pair (service-name, port-name) published by MPI_Publish_name(), so that applications can no longer use MPI_Lookup_name() to find portname.
<pre>MPI_Wait(MPI_Request *request, MPI_Status *status)</pre>	Waits for an MPI send or receive to complete.
<pre>MPI_Waitall(int count, MPI_Request array_of_requests[], MPI_Status array_of_statuses[])</pre>	Waits for all of the given communications to complete.
<pre>MPI_Waitany(int count, MPI_Request array_of_requests[], int *index, MPI_Status *status)</pre>	Waits for any of the given communications to complete.
<pre>MPI_Waitsome(int incount, MPI_Request array_of_requests[], int *outcount, int array_of_indices[], MPI_Status array_of_statuses[])</pre>	Waits for some given communications to complete.
MPI_Win_c2f(MPI_Win win)	Translates a C handle into a Fortran handle.
MPI_Win_create(void *base, MPI_Aint size, int disp_unit, MPI_Info info, MPI_Comm comm, MPI_Win *win)	Opens a communication window in memory.
MPI_Win_create_errhandler(MPI_Win_errhandler_fn *function, MPI_Errhandler *errhandler)	Creates an error handler that can be attached to windows.

 TABLE A-1
 Sun MPI Routines (Continued)

Routine and C Syntax	Description
MPI_Win_create_keyval(MPI_Win_copy_attr_function *win_copy_attr_fn, MPI_Win_delete_attr_function *win_delete_attr_fn, int *win_keyval, void *extra_state)	Creates a caching attribute that can be associated with a window.
<pre>MPI_Win_delete_attr(MPI_Win win, int win_keyval)</pre>	Deletes the attribute created with MPI_Win_create_keyval.
MPI_Win_f2c(MPI_Fint win)	Translates a Fortran handle into a C handle.
MPI_Win_fence(int assert, MPI_Win win)	Synchronizes RMA calls on a window.
<pre>MPI_Win_free(MPI_Win *win)</pre>	Frees the window object and returns a null handle.
<pre>MPI_Win_free_keyval(int *win_keyval)</pre>	Releases a window attribute.
<pre>MPI_Win_get_attr(MPI_Win win, int win_keyval, void *attribute_val, int *flag)</pre>	Obtains the value of a window attribute.
<pre>MPI_Win_get_errhandler(MPI_Win win, MPI_Errhandler *errhandler)</pre>	Retrieves the error handler currently associated with a window.
<pre>MPI_Win_get_group(MPI_Win win, MPI_Group *group)</pre>	Returns a duplicate of the group of the communicator used to create the window.
<pre>MPI_Win_get_name(MPI_Win win, char *win_name, int *resultlen)</pre>	Returns the last name associated with a window object.
MPI_Win_lock(int lock_type, int rank, int assert, MPI_Win win)	Starts an RMA access epoch, during which only the window at the process with the specified rank can be accessed.
<pre>MPI_Win_post(MPI_Group group, int assert, MPI_Win win)</pre>	Starts an RMA exposure epoch for the local window associated with <i>win</i> .
<pre>MPI_Win_set_attr(MPI_Win win, int win_keyval, void *attribute_val)</pre>	Associates an attribute with a window.
<pre>MPI_Win_test(MPI_Win win, int *flag)</pre>	Attempts to complete an RMA exposure epoch; a nonblocking version of MPI_Win_wait.
MPI_Win_set_errhandler(MPI_Win win, MPI_Errhandler errhandler)	Attaches a new error handler to a window.
<pre>MPI_Win_set_name(MPI_Win win, char *win_name)</pre>	Assigns a name to a window.

 TABLE A-1
 Sun MPI Routines (Continued)

Routine and C Syntax	Description
MPI_Win_start(MPI_Group group, int assert, MPI_Win win)	Starts an RMA access epoch for win.
MPI_Win_unlock(int rank, MPI_Win win)	Completes an RMA access epoch started by a call to MPI_Win_lock().
double MPI_Wtick()	Returns the resolution of MPI_Wtime().
<pre>double MPI_Wtime()</pre>	Returns an elapsed time on the calling processor.

Sun MPI I/O Routines

TABLE A-2 lists the Sun MPI I/O routines in alphabetical order. The following sections list the routines by functional category.

File Manipulation

Collective coordination	Noncollective coordination
MPI_File_open()	<pre>MPI_File_delete()</pre>
MPI_File_close()	MPI_File_get_size()
MPI_File_set_size()	<pre>MPI_File_get_group()</pre>
<pre>MPI_File_preallocate()</pre>	<pre>MPI_File_get_amode()</pre>

File Info

Noncollective coordination	Collective coordination
MPI_File_get_info()	MPI_File_set_info()

Data access

Data Access With Explicit Offsets

Synchronism	Noncollective coordination	Collective coordination
Blocking	<pre>MPI_File_read_at() MPI_File_write_at()</pre>	<pre>MPI_File_read_at_all() MPI_File_write_at_all()</pre>
Nonblocking or split collective	MPI_File_iread_at()	<pre>MPI_File_read_at_all_begin() MPI_File_read_at_all_end()</pre>
	<pre>MPI_File_iwrite_at()</pre>	<pre>MPI_File_write_at_all_begin() MPI_File_write_at_all_end()</pre>

Data Access With Individual File Pointers

Synchronism	Noncollective coordination	Collective coordination
Blocking	<pre>MPI_File_read() MPI_File_write()</pre>	MPI_File_read_all() MPI_File_write_all()
Nonblocking or split collective	<pre>MPI_File_iread()</pre>	<pre>MPI_File_read_all_begin() MPI_File_read_all_end()</pre>
	<pre>MPI_File_iwrite()</pre>	<pre>MPI_File_write_all_begin() MPI_File_write_all_end()</pre>

Data Access With Shared File Pointers

Synchronism	Noncollective coordination	Collective coordination
Blocking	<pre>MPI_File_read_shared() MPI_File_write_shared()</pre>	<pre>MPI_File_read_ordered() MPI_File_write_ordered() MPI_File_seek_shared() MPI_File_get_position_shared()</pre>
Nonblocking or split collective	<pre>MPI_File_iread_shared() MPI_File_iwrite_shared()</pre>	<pre>MPI_File_read_ordered_begin() MPI_File_read_ordered_end() MPI_File_write_ordered_begin() MPI_File_write_ordered_end()</pre>

Pointer Manipulation

```
MPI_File_seek()
MPI_File_get_position()
MPI_File_get_byte_offset()
```

File Interoperability

```
MPI_Register_datarep()
MPI_File_get_type_extent()
```

File Consistency and Semantics

```
MPI_File_set_atomicity()
MPI_File_get_atomicity()
MPI_File_sync()
```

Handle Translation

```
MPI_File_f2c()
MPI_File_c2f()
```

MPI I/O Routines: Alphabetical Listing

TABLE A-2 Sun MPI I/O Routines

Routine and C Syntax	Description
MPI_File_c2f(MPI_File file)	Translates a C handle into a Fortran handle.
<pre>MPI_File_close(MPI_File *fh)</pre>	Closes a file (collective).
MPI_File_create_errhandler(MPI_File_errhandler_fn *function, MPI_Errhandler *errhandler)	Creates an MPI-style error handler that can be attached to a file.
MPI_File_delete(char *filename, MPI_Info info)	Deletes a file.
MPI_File_f2c(MPI_File file)	Translates a Fortran handle into a C handle.
<pre>MPI_File_get_amode(MPI_File fh, int *amode)</pre>	Returns mode associated with open file.
<pre>MPI_File_get_atomicity(MPI_File fh, int *flag)</pre>	Returns current consistency semantics for data-access operations.
<pre>MPI_File_get_byte_offset(MPI_File fh, MPI_Offset offset, MPI_Offset *disp)</pre>	Converts a view-relative offset into an absolute byte position.
MPI_File_get_errhandler(MPI_Comm file, MPI_Errhandler *errhandler)	Gets the error handler for a file.
<pre>MPI_File_get_group(MPI_File fh, MPI_Group *group)</pre>	Returns the process group of file.

TABLE A-2 Sun MPI I/O Routines (Continued)

Routine and C Syntax	Description
MPI_File_get_info(MPI_File fh, MPI_Info *info_used)	Returns a new info object containing hints.
MPI_File_get_position(MPI_File fh, MPI_Offset *offset)	Returns current position of individual file pointer.
<pre>MPI_File_get_position_shared(MPI_File fh, MPI_Offset *offset)</pre>	Returns current position of the shared file pointer (collective).
MPI_File_get_size(MPI_File fh, MPI_Offset *size)	Returns current size of file.
MPI_File_get_type_extent(MPI_File fh, MPI_Datatype datatype, MPI_Aint *extent)	Returns the extent of the data type in a file.
MPI_File_get_view(MPI_File fh, MPI_Offset *disp, MPI_Datatype *etype, MPI_Datatype *filetype, char *datarep)	Returns process's view of data in file.
MPI_File_iread(MPI_File fh, void *buf, int count, MPI_Datatype datatype, MPI_Request *request)	Reads a file starting at the location specified by the individual file pointer (nonblocking, noncollective).
MPI_File_iread_at(MPI_File fh, MPI_Offset offset, void *buf, int count, MPI_Datatype datatype, MPI_Request *request)	Reads a file at an explicitly specified offset (nonblocking, noncollective).
MPI_File_iread_shared(MPI_File fh, void *buf, int count, MPI_Datatype datatype, MPI_Request *request)	Reads a file using the shared file pointer (nonblocking, noncollective).
MPI_File_iwrite(MPI_File fh, void *buf, int count, MPI_Datatype datatype, MPI_Request *request)	Writes a file starting at the location specified by the individual file pointer (nonblocking, noncollective).
MPI_File_iwrite_at(MPI_File fh, MPI_Offset offset, void *buf, int count, MPI_Datatype datatype, MPI_Request *request)	Writes a file at an explicitly specified offset (nonblocking, noncollective).
MPI_File_iwrite_shared(MPI_File fh, void *buf, int count, MPI_Datatype datatype, MPI_Request *request)	Writes a file using the shared file pointer (nonblocking, noncollective).
MPI_File_open(MPI_Comm comm, char *filename, init amode, MPI_Info info, MPI_File *fh)	Opens a file (collective).
<pre>MPI_File_preallocate(MPI_File fh, MPI_Offset size)</pre>	Preallocates storage space for a portion of a file (collective).

TABLE A-2 Sun MPI I/O Routines (Continued)

Routine and C Syntax	Description
MPI_File_read(MPI_File fh, void *buf, int count, MPI_Datatype datatype, MPI_Status *status)	Reads a file starting at the location specified by the individual file pointer.
MPI_File_read_all(MPI_File fh, void *buf, int count, MPI_Datatype datatype, MPI_Status *status)	Reads a file starting at the locations specified by individual file pointers (collective).
MPI_File_read_all_begin(MPI_File fh, void *buf, int count, MPI_Datatype datatype)	Reads a file starting at the locations specified by individual file pointers; the beginning part of a split collective routine (nonblocking).
MPI_File_read_all_end(MPI_File fh, void *buf, MPI_Status *status)	Reads a file starting at the locations specified by individual file pointers; the ending part of a split collective routine (blocking).
MPI_File_read_at(MPI_File fh, MPI_Offset offset, void *buf, int count, MPI_Datatype datatype, MPI_Status *status)	Reads a file at an explicitly specified offset.
MPI_File_read_at_all(MPI_File fh, MPI_Offset offset, void *buf, int count, MPI_Datatype datatype, MPI_Status *status)	Reads a file at explicitly specified offsets (collective).
MPI_File_read_at_all_begin(MPI_File fh, MPI_Offset offset, void *buf, int count, MPI_Datatype datatype)	Reads a file at explicitly specified offsets; the beginning part of a split collective routine (nonblocking).
MPI_File_read_at_all_end(MPI_File fh, void *buf, MPI_Status *status)	Reads a file at explicitly specified offsets; the ending part of a split collective routine (blocking).
MPI_File_read_ordered(MPI_File fh, void *buf, int count, MPI_Datatype datatype, MPI_Status *status)	Reads a file at a location specified by a shared file pointer (collective).
MPI_File_read_ordered_begin(MPI_File fh, void *buf, int count, MPI_Datatype datatype)	Reads a file at a location specified by a shared file pointer; the beginning part of a split collective routine (nonblocking).
MPI_File_read_ordered_end(MPI_File fh, void *buf, MPI_Status *status)	Reads a file at a location specified by a shared file pointer; the ending part of a split collective routine (blocking).
MPI_File_read_shared(MPI_File fh, void *buf, int count, MPI_Datatype datatype, MPI_Status *status)	Reads a file using the shared file pointer (blocking, noncollective).
MPI_File_seek(MPI_File fh, MPI_Offset offset, int whence)	Updates individual file pointers.

 TABLE A-2
 Sun MPI I/O Routines (Continued)

Routine and C Syntax	Description
MPI_File_seek_shared(MPI_File fh, MPI_Offset offset, int whence)	Updates the global shared file pointer (collective).
<pre>MPI_File_set_atomicity(MPI_File fh, int flag)</pre>	Sets consistency semantics for data-access operations (collective).
MPI_File_set_errhandler(MPI_File file, MPI_Errhandler errhandler)	Sets the error handler for a file.
MPI_File_set_info(MPI_File fh, MPI_Info info)	Sets new values for hints (collective).
<pre>MPI_File_set_size(MPI_File fh, MPI_Offset size)</pre>	Resizes a file (collective).
MPI_File_set_view(MPI_File fh, MPI_Offset disp, MPI_Datatype etype, MPI_Datatype filetype, char *datarep, MPI_Info info)	Changes the process's view of data in file (collective).
MPI_File_sync(MPI_File fh)	Makes semantics consistent for data-access operations (collective).
MPI_File_write(MPI_File fh, void *buf, int count, MPI_Datatype datatype, MPI_Status *status)	Writes a file starting at the location specified by the individual file pointer.
MPI_File_write_all(MPI_File fh, void *buf, int count, MPI_Datatype datatype, MPI_Status *status)	Writes a file starting at the locations specified by individual file pointers (collective).
MPI_File_write_all_begin(MPI_File fh, void *buf, int count, MPI_Datatype datatype)	Writes a file starting at the locations specified by individual file pointers; the beginning part of a split collective routine (nonblocking).
MPI_File_write_all_end(MPI_File fh, void *buf, MPI_Status *status)	Writes a file starting at the locations specified by individual file pointers;the ending part of a split collective routine (blocking).
<pre>MPI_File_write_at(MPI_File fh, MPI_Offset offset, void *buf, int count, MPI_Datatype datatype, MPI_Status *status)</pre>	Writes a file at an explicitly specified offset.
MPI_File_write_at_all(MPI_File fh, MPI_Offset offset, void *buf, int count, MPI_Datatype datatype, MPI_Status *status)	Writes a file at explicitly specified offsets (collective).
MPI_File_write_at_all_begin(MPI_File fh, MPI_Offset offset, void *buf, int count, MPI_Datatype datatype)	Writes a file at explicitly specified offsets; the beginning part of a split collective routine (nonblocking).

TABLE A-2 Sun MPI I/O Routines (Continued)

Routine and C Syntax	Description
MPI_File_write_at_all_end(MPI_File fh, void *buf, MPI_Status *status)	Writes a file at explicitly specified offsets; the ending part of a split collective routine (blocking).
MPI_File_write_ordered(MPI_File fh, void *buf, int count, MPI_Datatype datatype, MPI_Status *status)	Writes a file at a location specified by a shared file pointer (collective).
MPI_File_write_ordered_begin(MPI_File fh, void *buf, int count, MPI_Datatype datatype)	Writes a file at a location specified by a shared file pointer; the beginning part of a split collective routine (nonblocking).
<pre>MPI_File_write_ordered_end(MPI_File fh, void *buf, MPI_Status *status)</pre>	Writes a file at a location specified by a shared file pointer; the ending part of a split collective routine (blocking).
<pre>MPI_File_write_shared(MPI_File fh, void *buf, int count, MPI_Datatype datatype, MPI_Status *status)</pre>	Writes a file using the shared file pointer (blocking, noncollective).
MPI_Register_datarep(char *datarep, MPI_Datarep_conversion_function *read_conversion_fn, MPI_Datarep_conversion_function *write_conversion_fn, MPI_Datarep_extent_function *dtype_file_extent_fn, void *extra_state)	Defines data representation.

Environment Variables

Many environment variables are available for fine-tuning your Sun MPI environment. All Sun MPI environment variables are listed here with brief descriptions. The same descriptions are also available on the MPI man page. If you want to return to the default setting after setting a variable, simply unset it (using unsetenv). The effects of some of the variables are explained in more detail in the Sun HPC ClusterTools Software Performance Guide.

The environment variables are listed here in six groups:

- "Informational Variables" on page 103
- "General Performance Tuning" on page 104
- "Tuning Memory for Point-to-Point Performance" on page 106
- "Numerics" on page 109
- "Synchronization of One-Sided Communications" on page 109
- "MPProf" on page 112
- "Miscellaneous" on page 113

Informational Variables

MPI_PRINTENV

When set to 1, this variable causes other environment variables and the hpc.conf parameters associated with the MPI job to be printed. The default value is 0.

MPI QUIET

If set to 1, this variable suppresses Sun MPI warning messages. The default value is 0.

MPI SHOW ERRORS

If set to 1, the MPI ERRORS RETURN error handler prints the error message and returns the error. The default value is 0.

MPI SHOW INTERFACES

When set to 1, 2, or 3, information regarding which interfaces are being used by an MPI application is printed to stdout. Set MPI_SHOW_INTERFACES to 1 to print the selected internode interface. Set it to 2 to print all the interfaces and their rankings. Set it to 3 for verbose output. The default value, 0, does not print information to stdout.

General Performance Tuning

MPI POLLALL

When this variable is set to 1, the default value, all connections are polled for receives, also known as full polling. When set to 0, only those connections are polled where receives are posted. Full polling helps drain system buffers, lessening the chance of deadlock for "unsafe" codes. Well-written codes should set MPI_POLLALL to 0 for best performance.

Note – If MPI_POLLALL is set to 0 (zero) and your program performs an MPI_Send/MPI_Cancel without a corresponding MPI call on the receiving process, the MPI_Cancel may not succeed. Your program may hang.

MPI PROCBIND

Binds each MPI process to its own processor. The system administrator can allow or disable processor binding by setting the allow_pbind parameter in the CREOptions section of the hpc.conf file. If this parameter is set, the MPI_PROCBIND environment variable is disabled. Performance can be enhanced with processor binding, but very poor performance will result if processor binding is used for multithreaded jobs or for more than one job at a time.

By default, MPI_PROCBIND is set to 0, which turns off processor binding. To turn on processor binding, set the value to 1. With processor binding turned on, the processes in a job are assigned to all CPUs that are not already bound to other processes.

You can further control CPU binding by using these values for MPI_PROCBIND:

```
MPI_PROCBIND \{P|L|T\} [list | range]
```

- L Only the first thread in each process is bound to a CPU. Any additional threads created by the job are not bound.
- P Every thread created by the process is bound to a CPU.
- T Every thread in every process that is part of the same task is bound to a CPU.
 (See the mprun -P option.)

To specify the CPUs to which the threads are bound, you use either list or range.

■ *list* – Provides explicit binding and can be a single CPU ID or a list of CPU IDs separated by commas. For example, to assign a thread to the fifth CPU in a 24-CPU node, use:

```
% setenv MPI_PROCBIND L4
```

If a node had fewer CPUs than the number you specified, CRE would assign the thread to the CPU that was the modulus of the number you specified, divided by the number of CPUs. For example, a list of L12 in a node with only 8 CPUs would result in the process being assigned to CPU number 3.

When you use a list, the CRE environment does not check to see whether those CPU's are already bound. As a result, you could have two threads bound to the same CPU.

range – Provides automatic binding and assigns threads to all available (unbound)
 CPUs within the range. The range is expressed as:

 $N-M\times I$

In other words, specify the range with a starting number (N), an ending number (M), and a counting interval (I). The counting interval (I) is optional, and its default value is 1.

For example:

% setenv MPI_PROCBIND L0-11

The preceding setting would search through the first 12 CPUs, and assign processes to those that are unbound. If the number of available CPUs is less than the number of processes, the extra processes would remain unbound.

The mprun -v option (verbose) prints the CPU assignments that have been made. You can also run pbind(1M) on each node to verify their CPU bindings.

MPI SPIN

This variable sets the spin policy. The default value is 0, which causes MPI processes to spin nonaggressively, allowing best performance when the load is at least as great as the number of CPUs. A value of 1 causes MPI processes to spin aggressively, leading to best performance if extra CPUs are available on each node to handle system daemons and other background activities.

Tuning Memory for Point-to-Point Performance

MPI RSM CPOOLSIZE

This is the requested size, in bytes, to be allocated per stripe for buffers for each RSM connection. This value can be overridden when connections are established on the basis of the size of the segment allocated. The default value is 256 Kbytes.

MPI_RSM_NUMPOSTBOX

This is the number of postboxes per stripe per RSM connection. The maximum number of postboxes depends on the value of rsm_maxsegsize. The default is 128 postboxes.

MPI RSM PIPESIZE

This limits the size (in bytes) of a message that can be sent over remote shared memory through the buffer list of one postbox per stripe. The default is 64 Kbytes. This size also depends on the block size used for sending data. The maximum size is equal to $\min (cpoolsize/2, (10 * max(blk1sz, blk2sz)))$.

MPI_RSM_SBPOOLSIZE

If set, this variable specifies the requested size in bytes of each RSM send buffer pool. An RSM send buffer pool is the pool of buffers on a node that a remote process would use to send to processes on the node. A multiple of 1024 must be used. If unset, the size of buffer pool is equal to *cpoolsize* times processes per node. The max value allowed is *maxsegsize* minus the memory used for postboxes.

MPI RSM SHORTMSGSIZE

This variable specifies the maximum size, in bytes, of a message that will be sent by means of remote shared memory without using buffers. The default value is 3918 bytes. The upper limit is determined by the number of postboxes available.

MPI_RSM_STRONGPARTITION

If set to 1, the RSM protocol module uses strong partition to manage memory. Every connection has a set of blocks in the buffer pool reserved for communication. Otherwise, the pool is shared by all the receivers. The default value is 0.

MPI SHM CPOOLSIZE

This variable represents the amount of memory, in bytes, that can be allocated to each connection pool. When MPI_SHM_SBPOOLSIZE is not set, the default value is 24,576 bytes. Otherwise, the default value is MPI_SHM_SBPOOLSIZE.

MPI SHM CYCLESIZE

This variable represents the limit, in bytes, on the portion of a shared-memory message that will be sent via the buffer list of a single postbox during a cyclic transfer. The default value is 8192 bytes. A multiple of 1024 that is at most MPI_SHM_CPOOLSIZE/2 must be used.

MPI SHM CYCLESTART

Shared-memory transfers that are larger than MPI_SHM_CYCLESTART bytes are cyclic. The default value is 24,576 bytes.

MPI_SHM_NUMPOSTBOX

This variable represents the number of postboxes dedicated to each shared-memory connection. The default value is 16.

MPI SHM PIPESIZE

This variable represents the limit, in bytes, on the portion of a shared-memory message that will be sent via the buffer list of a single postbox during a pipeline transfer. The default value is 8192 bytes. The value must be a multiple of 1024.

MPI SHM PIPESTART

This variable represents the size, in bytes, at which shared-memory transfers starts to be pipelined. The default value is 2048. Multiples of 1024 must be used.

MPI_SHM_SBPOOLSIZE

If set, this variable represents the size, in bytes, of the pool of shared-memory buffers dedicated to each sender. A multiple of 1024 must be used. If unset, then pools of shared-memory buffers are dedicated to connections rather than to senders.

MPI_SHM_SHORTMSGSIZE

This variable represents the size (in bytes) of the section of a postbox that contains either data or a buffer list. The default value is 256 bytes.

Note — If MPI_SHM_PIPESTART, MPI_SHM_PIPESIZE, or MPI_SHM_CYCLESIZE is increased to a size larger than 31,744 bytes, then MPI_SHM_SHORTMSGSIZE might also have to be increased. See the *Sun HPC ClusterTools Software Performance Guide* for more information.

Numerics

MPI_CANONREDUCE

Prevents reduction operations from using any optimizations that take advantage of the physical location of processors. This can provide more consistent results in the case of floating-point addition, for example. However, the operation can take longer to complete. The default value is 0, meaning that optimizations are allowed. To prevent optimizations, set the value to 1.

Synchronization of One-Sided Communications

MPI_USE_AGENT_THREAD

If the environmental variable MPI_USE_AGENT_THREAD is set to 1, upon the first call to MPI_Win_create the Sun MPI library creates one agent thread for processes that need such a thread. (If MPI_USE_AGENT_THREAD is not set, or it is set to 0 [zero], no such thread is created.)

The two purposes of MPI_USE_AGENT_THREAD are to ensure progress in passive target RMA synchronization and to perform MPI RMA operations on local window memory on behalf of other processes when those processes do not have direct (shared-memory or RSM) access to window memory.

The agent thread does not run user code. Thread-safety in the non-thread-safe MPI library is achieved by a monitor around MPI communication calls. If no windows requiring the use of an agent thread are active, the agent thread is suspended If MPI_USE_AGENT_THREAD is not set, one-sided MPI operations can be delayed till the next synchronization call.

MPI RSM PUTSIZE

Use the environmental variable MPI RSM PUTSIZE to control the method used to put small buffers over the RSM PM. Buffers passed to MPI Put that are smaller than MPI_RSM_PUTSIZE are transferred to the remote side using a message protocol that has low latency but requires an implicit synchronization with the remote side, which may delay the sending side and reduce performance. The delay is usually small if both sides perform frequent small transfers. Buffers larger than MPI_RSM_PUTSIZE are transferred using a direct one-sided protocol which does not require synchronization, but has higher latency. The cost of the higher latency is not significant for large enough messages. Choose a value of 0 (zero) to always use the direct protocol, and a large value to always use the message protocol. The default value is 16 Kbytes.

MPI RSM GETSIZE

Use the environmental variable MPI RSM GETSIZE to control the method used to get small buffers over the RSM PM. Buffers passed to MPI Get that are larger than MPI RSM GETSIZE are transferred using a message protocol that achieves high bandwidth but requires an implicit synchronization with the remote side, which may delay the sending side and reduce performance. For large enough messages, the higher bandwidth offsets the cost of the synchronization delay. Buffers passed to MPI Get that are smaller than MPI RSM GETSIZE are transferred using a direct one-sided protocol which does not require synchronization, but has lower bandwidth. For small enough messages, the lower bandwidth has little impact on performance. Choose a value of 0 (zero) to always use the message protocol, and a large value to always use the direct protocol. The default value is 16Kbytes.

Tuning Rendezvous

MPI EAGERONLY

When this variable is set to 1, the default, only the eager protocol is used. When it is set to 0, both eager and rendezvous protocols are used.

MPI_RSM_RENDVSIZE

Messages communicated by remote shared memory that are greater than this size use the rendezvous protocol unless the environment variable MPI_EAGERONLY is set to 1. The default value is 256 Kbytes.

MPI_SHM_RENDVSIZE

Messages communicated by shared memory that are greater than this size use the rendezvous protocol unless the environment variable MPI_EAGERONLY is set. The default value is 24,576 bytes.

MPI_TCP_RENDVSIZE

Messages communicated by TCP that contain data of this size and greater use the rendezvous protocol unless the environment variable MPI_EAGERONLY is set. The default value is 49,152 bytes.

MPProf

MPI PROFILE

Setting this variable to 1 enables a profiling session. When profiling is enabled, profiling data for the MPI process ranks are written to a set of intermediate files, one file per process rank. MPProf also creates an index file of the form: mpprof.index.rm.jid (where rm is the resource manager and jid is the job ID) that contains pointers to the intermediate files of the form mpprof.n.rm.jid (where n is the process rank, rm is the resource manager, and jid is the job ID). If MPI PROFILE is not set, program execution proceeds without generating profiling data.

MPI PROFDATADIR

By default, the temporary files generated during MPProf profiling are located in /usr/tmp/. Set an alternative location as a value for environment variable MPI PROFDATADIR.

MPI PROFINDEXDIR

By default, the index file for MPProf profiling is located in the current directory. Set an alternative, nondefault location as a value for MPI_PROFINDEXDIR.

MPI PROFINTERVAL

The variable MPI_PROFINTERVAL can be used to specify a time interval for controlling when snapshots of the profiling data will be written to the intermediate files.

Setting MPI_PROFINTERVAL to 0 forces a snapshot for every MPI call that is made. Setting MPI PROFINTERVAL to Inf causes only one snapshot to be recorded at MPI Finalize time. If MPI PROFINTERVAL is unset or has no value, the default value of 60 seconds will be used.

If time intervals are used and an MPI program terminates before the MPI_Finalize call, any snapshots that were recorded can be used by mpprof to generate a profile of program operations up to the point of termination.

MPI_PROFMAXFILESIZE

This variable can be used to specify the maximum size, in Kbytes, that can be written to the intermediate files.

The default intermediate file size limit files is 51,200 Kbytes (50 Mbytes). If a process records data that exceeds the file size limit, that write operation completes, but it cannot record additional profiling data. Other intermediate files that have not reached the limit can continue to receive data. The file size limit can be removed altogether by setting MPI_PROFMAXFILESIZE to unlimited.

Miscellaneous

MPI_COSCHED

This variable specifies the user's preference regarding use of the spind daemon for coscheduling. The value can be 0 (prefer no use) or 1 (prefer use). This preference can be overridden by the system administrator's policy. This policy is set in the hpc.conf file and can be 0 (forbid use), 1 (require use), or 2 (no policy). If no policy is set and no user preference is specified, coscheduling is not used.

Note – If no user preference is specified, the value 2 is displayed when environment variables are printed with MPI_PRINTENV.

MPI_CHECK_ARGS

When this variable is set to 1, argument checking is performed on MPI calls, and errors are printed when they occur. The default is 0.

MPI FLOWCONTROL

This variable limits the number of unexpected messages that can be queued from a particular connection. Once this quantity of unexpected messages has been received, polling the connection for incoming messages stops. The default value, 0, indicates that no limit is set. To limit flow, set the value to an integer greater than 0.

MPI FULLCONNINIT

This variable ensures that all connections are established during initialization. By default, connections are established lazily. However, you can override this default by setting the environment variable MPI FULLCONNINIT to 1, forcing full-connection initialization mode. The default value is 0.

MPI MAXFHANDLES

This variable representst the maximum number of Fortran handles for objects other than requests. MPI MAXFHANDLES specifies the upper limit on the number of concurrently allocated Fortran handles for MPI objects other than requests. This variable is ignored in the default 32-bit library. The default value is 1024. Users should take care to free MPI objects that are no longer in use. There is no limit on handle allocation for C codes.

MPI MAXPROCS

This variable overrides the value specified by maxprocs_default in hpc.conf; it cannot exceed the value specified by maxprocs_limit in hpc.conf. If the value does exceed the maxprocs_limit value, the job aborts with an error when the program calls MPI_Init. Note that the upper limit of support for RSM communication is 2048 processes.

MPI MAXREOHANDLES

This variable representst the maximum number of Fortran request handles. MPI MAXREQHANDLES specifies the upper limit on the number of concurrently allocated MPI request handles. Users must take care to free up request handles by properly completing requests. The default value is 1024. This variable is ignored in the default 32-bit library.

MPI OPTCOLL

The MPI collectives are implemented using a variety of optimizations. Some of these optimizations can inhibit performance of point-to-point messages for "unsafe" programs. The default value of this variable, 1, indicates that optimized collectives are used. The optimizations can be turned off by setting the value to 0.

MPI RSM MAXSTRIPE

This variable specifies the maximum number of interfaces that can be used for striping data during communication over RSM. The value cannot be higher than 64 and the number of installed interfaces. The default is 4.

MPI SHM BCASTSIZE

On SMPs, MPI_Bcast() is implemented for large messages using a double-buffering scheme. The size of each buffer (in bytes) is settable by using this environment variable. The default value is 32,768 bytes.

MPI SHM GBPOOLSIZE

This variable representst the amount of memory available, in bytes, to the general buffer pool for use by collective operations. The default value is 20,971,520 bytes.

MPI SHM REDUCESIZE

On SMPs, calling $\texttt{MPI}_\texttt{Reduce}()$ causes all processors to participate in the reduce. Each processor works on a piece of data equal to the $\texttt{MPI}_\texttt{SHM}_\texttt{REDUCESIZE}$ setting. The default value is 256 bytes.

You must take care when setting this variable because the system reserves $MPI_SHM_REDUCESIZE*np*np$ memory to execute the reduce.

MPI_SPINDTIMEOUT

When coscheduling is enabled, this variable limits the length of time (in milliseconds) a message remains in the poll waiting for the spind daemon to return. If the timeout occurs before the daemon finds any messages, the process reenters the polling loop. The default value is 1000 milliseconds. A default can also be set by a system administrator in the hpc.conf file.

MPI TCP CONNLOOP

This variable sets the number of times MPI_TCP_CONNTIMEOUT occurs before signaling an error. The default value for this variable is 0, meaning that the program aborts on the first occurrence of MPI TCP CONNTIMEOUT.

MPI TCP CONNTIMEOUT

This variable sets the timeout value in seconds that is used for an accept() call. The default value for this variable is 600 seconds (10 minutes). This timeout can be triggered in both full- and lazy-connection initialization. After the timeout is reached, a warning message is printed. If MPI_TCP_CONNLOOP is set to 0, then the first timeout causes the program to abort.

MPI_TCP_SAFEGATHER

This variable allows use of a congestion-avoidance algorithm for MPI_Gather() and MPI Gatherv() over TCP. By default, MPI TCP SAFEGATHER is set to 1, which meansthat use of this algorithm is on. If you know that your underlying network can handle gathering large amounts of data on a single node, you might want to override this algorithm by setting MPI_TCP_SAFEGATHER to 0.

Troubleshooting

This appendix describes some common problem situations, resulting error messages, and suggestions for fixing the problems. It includes the following topics:

- "MPI Messages" on page 117
- "MPI I/O Error Handling" on page 120
- "Adjusting System V Shared-Memory Limits" on page 122

Sun MPI error reporting, including I/O, follows the MPI-2 Standard. By default, errors are reported in the form of standard error classes. These classes and their meanings are listed in TABLE C-1 (for non-I/O MPI) and TABLE C-2 (for MPI I/O) and are also available on the MPI man page.

Three predefined error handlers are available in Sun MPI:

- MPI_ERRORS_RETURN The default; returns an error code if an error occurs.
- MPI_ERRORS_ARE_FATAL I/O errors are fatal, and no error code is returned.
- MPI_THROW_EXCEPTION A special error handler to be used only with C++.

MPI Messages

You can make changes to and get information about the error handler by using any of the following routines:

- MPI_Comm_call_errhandler
- MPI_File_call_errhandler
- MPI_Win_call_errhandler
- MPI_Comm_create_errhandler
- MPI_Comm_get_errhandler
- MPI_Comm_set_errhandler
- MPI Add error class
- MPI_Add_error_code
- MPI_Add-error-string

Messages resulting from an MPI program fall into two categories:

- Error messages Error messages stem from within MPI. Usually an error message explains why your program cannot complete, and the program aborts.
- Warning messages Warnings stem from the environment in which you are running your MPI program and are usually sent by MPI_Init(). They are not associated with an aborted program; that is, programs continue to run despite warning messages.

Error Messages

Sun MPI error messages use a standard format:

[xyz] Error in function name: errclass string:intern(a): description: unixerrstring

Where

- $[x \ y \ z]$ is the process communication identifier, which is present in every error message, and:
 - \blacksquare *x* is the job ID (or jid).
 - y is the name of the communicator if a name exists; otherwise it is the address of the opaque object.
 - *z* is the rank of the process.
- function name is the name of the associated MPI function. It is present in every error message.
- errclass string is the string associated with the MPI error class. It is present in every error message.
- *intern* is an internal function. It is optional.
- *a* is a system call if one is the cause of the error. It is optional.
- description is a description of the error. It is optional.
- unixerrstring is the UNIX error string that describes system call a. It is optional.

Warning Messages

Sun MPI warning messages also use a standard format:

[x y z] Warning message

Where *message* is a description of the error.

Standard Error Classes

TABLE C-1 lists the error return classes you can encounter in your MPI programs. Error values may also be found in mpi.h (for C), mpif.h (for Fortran), and mpi++.h (for C++).

MPI I/O messages are listed separately, in TABLE C-2.

TABLE C-1 Sun MPI Standard Error Classes

Error Code	Value	Meaning
MPI_SUCCESS	0	Successful return code.
MPI_ERR_BUFFER	1	Invalid buffer pointer.
MPI_ERR_COUNT	2	Invalid count argument.
MPI_ERR_TYPE	3	Invalid datatype argument.
MPI_ERR_TAG	4	Invalid tag argument.
MPI_ERR_COMM	5	Invalid communicator.
MPI_ERR_RANK	6	Invalid rank.
MPI_ERR_ROOT	7	Invalid root.
MPI_ERR_GROUP	8	Null group passed to function.
MPI_ERR_OP	9	Invalid operation.
MPI_ERR_TOPOLOGY	10	Invalid topology.
MPI_ERR_DIMS	11	Illegal dimension argument.
MPI_ERR_ARG	12	Invalid argument.
MPI_ERR_UNKNOWN	13	Unknown error.
MPI_ERR_TRUNCATE	14	Message truncated on receive.
MPI_ERR_OTHER	15	Other error; use Error_string.
MPI_ERR_INTERN	16	Internal error code.
MPI_ERR_IN_STATUS	17	Look in status for error value.
MPI_ERR_PENDING	18	Pending request.
MPI_ERR_REQUEST	19	Illegal MPI_Request() handle.
MPI_ERR_KEYVAL	36	Illegal key value.

TABLE C-1 Sun MPI Standard Error Classes (Continued)

Error Code	Value	Meaning
MPI_ERR_INFO	37	Invalid info object.
MPI_ERR_INFO_KEY	38	Illegal info key.
MPI_ERR_INFO_NOKEY	39	No such key.
MPI_ERR_INFO_VALUE	40	Illegal info value.
MPI_ERR_TIMEDOUT	41	Timed out.
MPI_ERR_RESOURCES	42	Out of resources.
MPI_ERR_TRANSPORT	43	Transport layer error.
MPI_ERR_HANDSHAKE	44	Error accepting/connecting.
MPI_ERR_SPAWN	45	Error spawning.
MPI_ERR_WIN	46	Invalid window.
MPI_ERR_BASE	47	Invalid base.
MPI_ERR_SIZE	48	Invalid size.
MPI_ERR_DISP	49	Invalid displacement.
MPI_ERR_LOCKTYPE	50	Invalid lock type.
MPI_ERR_ASSERT	51	Invalid assert.
MPI_ERR_RMA_CONFLICT	52	Conflicting accesses to window.
MPI_ERR_RMA_SYNC	53	Erroneous RMA synchronization.
MPI_ERR_NO_MEM	54	Memory exhausted.
MPI_ERR_LASTCODE	55	Last error code.

MPI I/O Error Handling

Sun MPI I/O error reporting follows the MPI-2 Standard. By default, errors are reported in the form of standard error codes (found in

/opt/SUNWhpc/include/mpi.h). Error classes and their meanings are listed in TABLE C-2. You can also find them in mpif.h (for Fortran) and mpi++.h (for C++). You can change the default error handler by specifying MPI_FILE_NULL as the file handle with the routine MPI_File_set_errhandler(), even if no file is currently open. Or, you can use the same routine to change a specific file's error handler.

TABLE C-2 Sun MPI I/O Error Classes

Error Class	Value	Meaning
MPI_ERR_FILE	20	Bad file handle.
MPI_ERR_NOT_SAME	21	Collective argument not identical on all processes.
MPI_ERR_AMODE	22	Unsupported amode passed to open.
MPI_ERR_UNSUPPORTED_DATAREP	23	Unsupported datarep passed to MPI_File_set_view().
MPI_ERR_UNSUPPORTED_OPERATION	24	Unsupported operation, such as seeking on a file that supports only sequential access.
MPI_ERR_NO_SUCH_FILE	25	File (or directory) does not exist.
MPI_ERR_FILE_EXISTS	26	File exists.
MPI_ERR_BAD_FILE	27	Invalid file name (for example, path name too long).
MPI_ERR_ACCESS	28	Permission denied.
MPI_ERR_NO_SPACE	29	Not enough space.
MPI_ERR_QUOTA	30	Quota exceeded.
MPI_ERR_READ_ONLY	31	Read-only file system.
MPI_ERR_FILE_IN_USE	32	File operation could not be completed, because the file is currently open by a process.
MPI_ERR_DUP_DATAREP	33	Conversion functions could not be registered because a data representation identifier that was already defined was passed to MPI_REGISTER_DATAREP.
MPI_ERR_CONVERSION	34	An error occurred in a user-supplied data-conversion function.
MPI_ERR_IO	35	I/O error.
MPI_ERR_INFO	37	Invalid info object.

TABLE C-2 Sun MPI I/O Error Classes (Continued)

Error Class	Value	Meaning
MPI_ERR_INFO_KEY	38	Illegal info key.
MPI_ERR_INFO_NOKEY	39	No such key.
MPI_ERR_INFO_VALUE	40	Illegal info value.
MPI_ERR_LASTCODE	55	Last error code.

Adjusting System V Shared-Memory Limits

When using the Sun MPI library on a cluster equipped with RSM links, memory allocated from System V shared memory is used for process communication.

Given that the maximum number of such shared-memory segments is set to a conservative value by default, calls to allocate new memory segments can fail due to these system-imposed limits. When such calls fail, check the values specified for the following variables in /etc/system:

- shmsys:shminfo_shmseg
- shmsys:shminfo_shmmni
- shmsys:shminfo shmmax

If the values are too low, increase them by editing /etc/system and rebooting the system.

Index

A Argonne National Laboratory and MPE, 47 array sections, 17 attributes, with communicators, 16	and MPI I/O, 49 and multithreaded programming, 25 and process topologies, 14 compiling, 34 to 37 with profiling library, 27 See also include syntax. context, defined, 16
В	
blocking routines. See routines, blocking.	D -dalign option, 36 data type
C caching, with communicators, 16	possible values for C, 19 derived (user-defined), 17, 49
Cartesian topology. <i>See</i> topology, Cartesian. code samples. <i>See</i> sample programs. collective communication. <i>See</i> communication, collective.	possible values for Fortran, 17, 18 primitive, 17 debugging, 39 to 47 with mpe, 47
communication buffers, 17 collective, 12, 16, 21 in multithreaded programs, 25	See also Prism. displacement (disp), 50, 53
restrictions, 13 half-channel, 14 interprocess, 16 persistent request, defined, 14 point-to-point, 16 port, 14 communicator default, 16 defined, 16	elementary data type (etype), 50 environmental inquiry functions, 15 error handling, 15 and MPE, 30 and multithreaded programming, 25 error messages standard error classes (Sun MPI I/O), 120 standard error values (Sun MPI), 119

F	Mississippi State University URL, x
features, 1	Standards
file type (filetype), 50	profiling, 27
Fortran	URL, x
compiling with -dalign option, 36	University of New Mexico URL, x
compiling with -xalias option, 35	MPI I/O, 49
companing man manner option, or	Sun MPI implementation.
	See Sun MPI I/O.
_	MPI_COMM_GROUP, 16
G	MPI_COMM_WORLD
graph topology. See topology, graph.	as default communicator, 16
grid topology. <i>See</i> topology, Cartesian. group, defined, 16	multiprocessing environment (MPE), 29 to 30 and debugging, 47
	multithreaded programming, 23 to 26 stubbing thread calls, 37
Н	
header files, 31	N I
"holes" (in an MPI I/O file type), $50,54$	N
	nonblocking routines. See routines, nonblocking
I	0
I/O. See Sun MPI I/O, MPI I/O.	offset, 50
include syntax, 31	
intercommunicator, defined, 16	one-sided communication, 5
intracommunicator, 14	options
defined, 16	-dalign, 36 -xalias, 35
	-xallas, 33
L	_
libraries	Р
libthread.so, 37	persistent communication request. See
linking, 34 to 37	communication, persistent request.
linking with profiling library, 27	point-to-point
linking, 34 to 37	communication. <i>See</i> communication, point-to-point.
	routines. <i>See</i> routines, point-to-point.
NA.	Prism, 39 to 41
М	compilers to use, 36
man pages	process
Solaris, location, 37	relation to group, 16
Sun MPI, location, xi	process topologies, 14
modes for point-to-point communication, 4	profiling, 27 to 28
MPI	
Forum, URL, x	

R	defined, 39
rank, of a process, 14, 16	standard mode. See modes for point-to-point
ready mode. See modes for point-to-point	communication.
communication.	starting up, 15
receive. <i>See</i> routines, receive.	static libraries, and relinking, 37
routines	Sun MPI I/O, 49 to 66
all-gather, 12	synchronous mode. See modes for point-to-point
all-to-all, 12	communication.
blocking, 4, 12	
broadcast, 12	
collective, 12, 16	_
in multithreaded programs, 25	Т
for constructing communicators, 16	thread safety. See multithreaded programming.
data access (MPI I/O), 53 to 57	timers, 15
pointer manipulation, 55 to 56	topology
with explicit offsets, 54	Cartesian, 14
with individual file pointers, 55	graph, 14
with shared file pointers, 56 to 57	virtual, defined, 14
error-handling, 15	See also process topologies.
file consistency (MPI I/O), 58	1 1 0
file manipulation (MPI I/O), 52	
gather, 12	
for constructing groups, 16	V
local, 16	view, 50
nonblocking, 4	
point-to-point, 4	
receive, 4, 21	V
reduction, 12	X
scan, 12	-xaliasoption, 35
scatter, 12	
semantics (MPI I/O), 58	
send, 4, 21	
Sun MPI	
listed alphabetically, 78 to 95	
listed by functional category, 67 to 78 Sun MPI I/O	
listed alphabetically, 98 to 102 listed by functional category, 95 to 98	
isted by functional category, 33 to 30	
_	
S	
sample programs	
Sun MPI, 32 to 34	
Sun MPI I/O, 59 to 66	
send. See routines, send.	
shutting down, 15	

SPMD programs