SPECIAL ISSUE

MPI: A MESSAGE-PASSING INTERFACE STANDARD

The standard, called the Message Passing Interface (MPI), provides a common interface for distributed memory concurrent computers and networks of workstations. MPI functionality includes point-to-point and collective communication routines, as well as support for process groups, communication contexts, and application topologies. While making use of new ideas, the MPI standard is based largely on current practice, such as Express, PVM, NX/2 Vertex, and P4.

The main advantages of establishing a message passing interface are portability and ease of use; a standard for message passing is a key component in building a concurrent computing environment in which applications, software libraries, and tools can be transparently ported between different machines. Furthermore, the definition of a standard provides vendors with a clearly defined set of routines that they can implement efficiently, or in some cases provide hardware or low-level system support for, thereby enhancing scalability.

The MPI standardization effort involved about 60 people from 40 organizations, mainly from the United States and Europe. Most of the major vendors of concurrent computers have been involved in MPI, along with researchers from universities, government laboratories and industry. MPI is intended to be a standard message passing interface for applications running on MIMD distributed memory concurrent computers and workstation networks.
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PREFACE

Jack Dongarra

The Message Passing Interface effort began in the summer of 1991 when a small group of researchers started discussions at a mountain retreat in Austria. Out of that discussion came a Workshop on Standards for Message Passing in a Distributed Memory Environment held on April 29–30, 1992, in Williamsburg, Virginia. At this workshop the basic features essential to a standard message-passing interface were discussed, and a working group established to continue the standardization process. More formal meetings and discussions began in January 1993 and continued with meetings every six weeks with discussions via e-mail. The MPI Standard was completed in March of 1994. The MPI effort involved about 60 people from 40 organizations, mainly from the United States and Europe. Most of the major vendors of concurrent computers were involved in MPI, along with researchers from universities, government laboratories, and industry.

This effort defines the syntax and semantics of a core of library routines useful to a wide range of users writing portable message-passing programs in Fortran or C. The MPI effort was conducted in a similar spirit to the High-Performance Fortran Forum (HPFF).

MPI provides parallel hardware vendors with a clearly defined base set of routines that can be efficiently implemented. As a result, hardware vendors can build upon this collection of standard low-level routines to create higher-level routines for the distributed-memory communication environments supplied with their parallel machines. MPI provides a simple-to-use portable interface for the basic user, yet is powerful enough to allow programmers to use the high-performance message-passing operations available on advanced machines.

As an effort to create a “true” standard for message passing, researchers incorporated into MPI the most useful features of several systems, rather than choosing one system to adopt as a standard. Features were used from systems by IBM, Intel, nCUBE, PVM, Express, P4, and PARMACS. The message paradigm, we believe, will be attractive because of its wide portability and can be used in

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communications for distributed-memory and shared-memory multiprocessors, networks of workstations, and any combination of these elements. The paradigm will not be made obsolete by increases in network speeds or by architectures combining shared and distributed-memory components. As this standard is printed, we have implementations of MPI on various platforms with more expected in the months to come.

MPI operated on a very tight budget (in reality, it had no budget when the first meeting was announced). ARPA and NSF have supported research at various institutions and have made a contribution toward travel for the U.S. academics. Support for several European participants was provided by ESPRIT.

This issue of the Journal is also available in Postscript and HTML forms over the Internet. To retrieve the postscript file you can anonymous ftp to netlib2.cs.utk.edu; cd mpi; get mpi-report.ps. The HTML form can be found by the URL: http://www.mcs.anl.gov/mpi/mpi-report/mpi-report.html. An up-to-date list of errata for this document is being maintained. To receive a copy, send an e-mail message to netlib@ornl.gov with contents: send mpi.errata.ps from mpi.
INTRODUCTION TO MPI

1.1 Overview and Goals

Message passing is a paradigm used widely on certain classes of parallel machines, especially those with distributed memory. Although there are many variations, the basic concept of processes communicating through messages is well understood. Over the last ten years, substantial progress has been made in casting significant applications in this paradigm. Each vendor has implemented its own variant. More recently, several systems have demonstrated that a message-passing system can be efficiently and portably implemented. It is thus an appropriate time to try to define both the syntax and semantics of a core of library routines that will be useful to a wide range of users and efficiently implementable on a wide range of computers.

In designing MPI we have sought to make use of the most attractive features of a number of existing message-passing systems, rather than selecting one of them and adopting it as the standard. Thus, MPI has been strongly influenced by work at the IBM T.J. Watson Research Center [1, 2], Intel's NX/2 [23], Express [22], nCUBE's Vertex [21], p4 [7, 6], and PARMACS [5, 8]. Other important contributions have come from Zipcode [24, 25], Chimp [14, 15], PVM [4, 11], Chameleon [19], and PICL [18].

The MPI standardization effort involved about 60 people from 40 organizations mainly from the United States and Europe. Most of the major vendors of concurrent computers were involved in MPI, along with researchers from universities, government laboratories, and industry. The standardization process began with the Workshop on Standards for Message Passing in a Distributed Memory Environment, sponsored by the Center for Research on Parallel Computing, held April 29-30, 1992, in Williamsburg, Virginia [29]. At this workshop the basic features essential to a standard message-passing interface were discussed, and a working group established to continue the standardization process.
A preliminary draft proposal, known as MPII, was put forward by Dongarra, Hempel, Hey, and Walker in November 1992, and a revised version was completed in February 1993 [12]. MPII embodied the main features that were identified at the Williamsburg workshop as being necessary in a message-passing standard. Since MPII was primarily intended to promote discussion and “get the ball rolling,” it focused mainly on point-to-point communications. MPII brought to the forefront a number of important standardization issues, but did not include any collective communication routines and was not thread-safe.

In November 1992, a meeting of the MPI working group was held in Minneapolis, at which it was decided to place the standardization process on a more formal footing, and to generally adopt the procedures and organization of the High Performance Fortran Forum. Subcommittees were formed for the major component areas of the standard, and an e-mail discussion service established for each. In addition, the goal of producing a draft MPI standard by the Fall of 1993 was set. To achieve this goal the MPI working group met every 6 weeks for two days throughout the first 9 months of 1993, and presented the draft MPI standard at the Supercomputing 93 conference in November 1993. These meetings and the e-mail discussion together constituted the MPI Forum, membership of which has been open to all members of the high performance computing community.

The main advantages of establishing a message-passing standard are portability and ease-of-use. In a distributed memory communication environment in which the higher level routines and/or abstractions are built upon lower level message-passing routines the benefits of standardization are particularly apparent. Furthermore, the definition of a message-passing standard, such as that proposed here, provides vendors with a clearly defined base set of routines that they can implement efficiently, or in some cases provide hardware support for, thereby enhancing scalability.

The goal of the Message Passing Interface simply stated is to develop a widely used standard for writing message-passing programs. As such the interface should establish a practical, portable, efficient, and flexible standard for message passing.

A complete list of goals follows.

- Design an application programming interface (not necessarily for compilers or a system implementation library).
- Allow efficient communication: Avoid memory-to-memory copying and allow overlap of computation and communication and offload to communication co-processor, where available.
- Allow for implementations that can be used in a heterogeneous environment.
- Allow convenient C and Fortran 77 bindings for the interface.
- Assume a reliable communication interface: the user need not cope with communication failures. Such failures are dealt with by the underlying communication subsystem.
• Define an interface that is not too different from current practice, such as PVM, NX, Express, p4, etc., and provides extensions that allow greater flexibility.
• Define an interface that can be implemented on many vendors’ platforms, with no significant changes in the underlying communication and system software.
• Semantics of the interface should be language independent.
• The interface should be designed to allow for thread-safety.

1.2 Who Should Use This Standard?

This standard is intended for use by all those who want to write portable message-passing programs in Fortran 77 and C. This includes individual application programmers, developers of software designed to run on parallel machines, and creators of environments and tools. In order to be attractive to this wide audience, the standard must provide a simple, easy-to-use interface for the basic user while not semantically precluding the high-performance message-passing operations available on advanced machines.

1.3 What Platforms Are Targets For Implementation?

The attractiveness of the message-passing paradigm at least partially stems from its wide portability. Programs expressed this way may run on distributed-memory multiprocessors, networks of workstations, and combinations of all of these. In addition, shared-memory implementations are possible. The paradigm will not be made obsolete by architectures combining the shared- and distributed-memory views, or by increases in network speeds. It thus should be both possible and useful to implement this standard on a great variety of machines, including those “machines” consisting of collections of other machines, parallel or not, connected by a communication network.

The interface is suitable for use by fully general MIMD programs, as well as those written in the more restricted style of SPMD. Although no explicit support for threads is provided, the interface has been designed so as not to prejudice their use. With this version of MPI no support is provided for dynamic spawning of tasks.

MPI provides many features intended to improve performance on scalable parallel computers with specialized interprocessor communication hardware. Thus, we expect that native, high-performance implementations of MPI will be provided on such machines. At the same time, implementations of MPI on top of standard Unix interprocessor communication protocols will provide portability to workstation clusters and heterogenous networks of workstations. Several proprietary, native implementations of MPI, and a public domain, portable implementation of MPI are in progress at the time of this writing [17, 13].
1.4 What is Included in the Standard?

The standard includes:

- Point-to-point communication
- Collective operations
- Process groups
- Communication contexts
- Process topologies
- Bindings for Fortran 77 and C
- Environmental Management and inquiry
- Profiling interface

1.5 What Is Not Included in the Standard?

The standard does not specify:

- Explicit shared-memory operations
- Operations that require more operating system support than is currently standard; for example, interrupt-driven receives, remote execution, or active messages
- Program construction tools
- Debugging facilities
- Explicit support for threads
- Support for task management
- I/O functions

There are many features that have been considered and not included in this standard. This happened for a number of reasons, one of which is the time constraint that was self-imposed in finishing the standard. Features that are not included can always be offered as extensions by specific implementations. Perhaps future versions of MPI will address some of these issues.

1.6 Organization of This Document

The following is a list of the remaining chapters in this document, along with a brief description of each.

- Chapter 2, MPI Terms and Conventions, explains notational terms and conventions used throughout the MPI document.
- Chapter 3, Point-to-Point Communication, defines the basic, pairwise communication subset of MPI. send and receive are found here, along with many associated functions designed to make basic communication powerful and efficient.
- Chapter 4, Collective Communication, defines process-group collective communication operations. Well-known examples of this are barrier and broadcast over a group of processes (not necessarily all the processes).
• Chapter 5, Groups, Contexts, and Communicators, shows how groups of processes are formed and manipulated, how unique communication contexts are obtained, and how the two are bound together into a communicator.

• Chapter 6, Process Topologies, explains a set of utility functions meant to assist in the mapping of process groups (a linearly ordered set) to richer topological structures such as multi-dimensional grids.

• Chapter 7, MPI Environmental Management, explains how the programmer can manage and make inquiries of the current MPI environment. These functions are needed for the writing of correct, robust programs, and are especially important for the construction of highly portable message-passing programs.

• Chapter 8, Profiling Interface, explains a simple name-shifting convention that any MPI implementation must support. One motivation for this is the ability to put performance profiling calls into MPI without the need for access to the MPI source code. The name shift is merely an interface; it says nothing about how the actual profiling should be done and in fact, the name shift can be useful for other purposes.

• Annex A, Language Binding, gives specific syntax in Fortran 77 and C, for all MPI functions, constants, and types.

• The MPI Function Index is a simple index showing the location of the precise definition of each MPI function, together with both C and Fortran bindings.

2.2 Procedure Specification

MPI procedures are specified using a language-independent notation. The arguments of procedure calls are marked as IN, OUT, or INOUT. The meanings of these are:

• the call uses but does not update an argument marked IN,
• the call may update an argument marked OUT,
• the call both uses and updates an argument marked INOUT.

There is one special case where an argument is a handle to an opaque object (these terms are defined in section 2.4.1), and the object is updated by the procedure call, then the argument is marked OUT. It is marked this way even
CHAPTER 2

MPI TERMS AND CONVENTIONS

This chapter explains notational terms and conventions used throughout the MPI document, some of the choices that have been made, and the rationale behind those choices.

2.1 Document Notation

Rationale. Throughout this document, the rationale for the design choices made in the interface specification is set off in this format. Some readers may wish to skip these sections, while readers interested in interface design may want to read them carefully. (End of rationale.)

Advice to users. Throughout this document, material that speaks to users and illustrates usage is set off in this format. Some readers may wish to skip these sections, while readers interested in programming in MPI may want to read them carefully. (End of advice to users.)

Advice to implementors. Throughout this document, material that is primarily commentary to implementors is set off in this format. Some readers may wish to skip these sections, while readers interested in MPI implementations may want to read them carefully. (End of advice to implementors.)

2.2 Procedure Specification

MPI procedures are specified using a language-independent notation. The arguments of procedure calls are marked as IN, OUT or INOUT. The meanings of these are:

- the call uses but does not update an argument marked IN,
- the call may update an argument marked OUT,
- the call both uses and updates an argument marked INOUT.

There is one special case—if an argument is a handle to an opaque object (these terms are defined in Section 2.4.1), and the object is updated by the procedure call, then the argument is marked OUT. It is marked this way even
though the handle itself is not modified—we use the OUT attribute to denote that what the handle references is updated.

The definition of MPI tries to avoid, to the largest possible extent, the use of INOUT arguments, because such use is error-prone, especially for scalar arguments.

A common occurrence for MPI functions is an argument that is used as IN by some processes and OUT by other processes. Such argument is, syntactically, an INOUT argument and is marked as such, although, semantically, it is not used in one call both for input and for output.

Another frequent situation arises when an argument value is needed only by a subset of the processes. When an argument is not significant at a process then an arbitrary value can be passed as argument.

Unless specified otherwise, an argument of type OUT or type INOUT cannot be aliased with any other argument passed to an MPI procedure. An example of argument aliasing in C appears below. If we define a C procedure like this,

```c
void copyIntBuffer( int *pin, int *pout, int len )
{
    int i;
    for (i=0; i<len; ++i) *pout++ = *pin++;
}
```

then a call to it in the following code fragment has aliased arguments.

```c
int a[10];
copyIntBuffer( a, a+3, 7);
```

Although the C language allows this, such usage of MPI procedures is forbidden unless otherwise specified. Note that Fortran prohibits aliasing of arguments.

All MPI functions are first specified in the language-independent notation. Immediately below this, the ANSI C version of the function is shown, and below this, a version of the same function in Fortran 77.

### 2.3 Semantic Terms

When discussing MPI procedures the following semantic terms are used. The first two are usually applied to communication operations.

**nonblocking** If the procedure may return before the operation completes, and before the user is allowed to re-use resources (such as buffers) specified in the call.

**blocking** If return from the procedure indicates the user is allowed to re-use resources specified in the call.

**local** If completion of the procedure depends only on the local executing process. Such an operation does not require communication with another user process.

**non-local** If completion of the operation may require the execution of some
MPI procedure on another process. Such an operation may require communication occurring with another user process.

**collective** If all processes in a process group need to invoke the procedure.

### 2.4 Data Types

#### 2.4.1 OPAQUE OBJECTS

MPI manages system memory that is used for buffering messages and for storing internal representations of various MPI objects such as groups, communicators, datatypes, etc. This memory is not directly accessible to the user, and objects stored there are **opaque**: their size and shape is not visible to the user. Opaque objects are accessed via **handles**, which exist in user space. MPI procedures that operate on opaque objects are passed handle arguments to access these objects. In addition to their use by MPI calls for object access, handles can participate in assignment and comparisons.

In Fortran, all handles have type **INTEGER**. In C, a different handle type is defined for each category of objects. These should be types that support assignment and equality operators.

In Fortran, the handle can be an index to a table of opaque objects in system table; in C it can be such index or a pointer to the object. More bizarre possibilities exist.

Opaque objects are allocated and deallocated by calls that are specific to each object type. These are listed in the sections where the objects are described. The calls accept a handle argument of matching type. In an allocate call this is an OUT argument that returns a valid reference to the object. In a call to deallocate this is an INOUT argument which returns with an “invalid handle” value. MPI provides an “invalid handle” constant for each object type. Comparisons to this constant are used to test for validity of the handle.

A call to deallocate invalidates the handle and marks the object for deallocation. The object is not accessible to the user after the call. However, MPI need not deallocate the object immediately. Any operation pending (at the time of the deallocate) that involves this object will complete normally; the object will be deallocated afterwards.

An opaque object and its handle are significant only at the process where the object was created, and cannot be transferred to another process.

MPI provides certain predefined opaque objects and predefined, static handles to these objects. Such objects may not be destroyed.

**Rationale.** This design hides the internal representation used for MPI data structures, thus allowing similar calls in C and Fortran. It also avoids conflicts with the typing rules in these languages, and easily allows future extensions of functionality. The mechanism for opaque objects used here loosely follows the POSIX Fortran binding standard.

The explicit separating of handles in user space, objects in system space, allows space-reclaiming, deallocation calls to be made at appropriate points
in the user program. If the opaque objects were in user space, one would have to be very careful not to go out of scope before any pending operation requiring that object completed. The specified design allows an object to be marked for deallocation, the user program can then go out of scope, and the object itself still persists until any pending operations are complete.

The requirement that handles support assignment/comparison is made since such operations are common. This restricts the domain of possible implementations. The alternative would have been to allow handles to have been an arbitrary, opaque type. This would force the introduction of routines to do assignment and comparison, adding complexity, and was therefore ruled out. (End of rationale.)

Advice to users. A user may accidentally create a dangling reference by assigning to a handle the value of another handle, and then deallocating the object associated with these handles. Conversely, if a handle variable is deallocated before the associated object is freed, then the object becomes inaccessible (this may occur, for example, if the handle is a local variable within a subroutine, and the subroutine is exited before the associated object is deallocated). It is the user’s responsibility to avoid adding or deleting references to opaque objects, except as a result of calls that allocate or deallocate such objects. (End of advice to users.)

Advice to implementors. The intended semantics of opaque objects is that each opaque object is separate from each other; each call to allocate such an object copies all the information required for the object. Implementations may avoid excessive copying by substituting referencing for copying. For example, a derived datatype may contain references to its components, rather then copies of its components; a call to MPI.COMM.GROUP may return a reference to the group associated with the communicator, rather than a copy of this group. In such cases, the implementation must maintain reference counts, and allocate and deallocate objects such that the visible effect is as if the objects were copied. (End of advice to implementors.)

2.4.2 ARRAY ARGUMENTS

An MPI call may need an argument that is an array of opaque objects, or an array of handles. The array-of-handles is a regular array with entries that are handles to objects of the same type in consecutive locations in the array. Whenever such an array is used, an additional len argument is required to indicate the number of valid entries (unless this number can be derived otherwise). The valid entries are at the beginning of the array; len indicates how many of them there are, and need not be the entire size of the array. The same approach is followed for other array arguments.
2.4.3 STATE

MPI procedures use at various places arguments with state types. The values of such data type are all identified by names, and no operation is defined on them. For example, the MPI.ERRHANDLER_SET routine has a state type argument with values MPI_ERRORS.ARE_FATAL, MPI_ERRORS.RETURN, etc.

2.4.4 NAMED CONSTANTS

MPI procedures sometimes assign a special meaning to a special value of a basic type argument; e.g. tag is an integer-valued argument of point-to-point communication operations, with a special wild-card value, MPI.ANY_TAG. Such arguments will have a range of regular values, which is a proper subrange of the range of values of the corresponding basic type; special values (such as MPI.ANY_TAG) will be outside the regular range. The range of regular values can be queried using environmental inquiry functions (Chapter 7).

2.4.5 CHOICE

MPI functions sometimes use arguments with a choice (or union) data type. Distinct calls to the same routine may pass by reference actual arguments of different types. The mechanism for providing such arguments will differ from language to language. For Fortran, the document uses <type> to represent a choice variable, for C, we use (void *).

2.4.6 ADDRESSES

Some MPI procedures use address arguments that represent an absolute address in the calling program. The datatype of such an argument is an integer of the size needed to hold any valid address in the execution environment.

2.5 Language Binding

This section defines the rules for MPI language binding in general and for Fortran 77 and ANSI C in particular. Defined here are various object representations, as well as the naming conventions used for expressing this standard. The actual calling sequences are defined elsewhere.

It is expected that any Fortran 90 and C++ implementations use the Fortran 77 and ANSI C bindings, respectively. Although we consider it premature to define other bindings to Fortran 90 and C++, the current bindings are designed to encourage, rather than discourage, experimentation with better bindings that might be adopted later.

Since the word PARAMETER is a keyword in the Fortran language, we use the word "argument" to denote the arguments to a subroutine. These are normally referred to as parameters in C, however, we expect that C programmers will understand the word "argument" (which has no specific meaning in C), thus allowing us to avoid unnecessary confusion for Fortran programmers.
double precision a
integer b
...
call MPI_send(a,...)
call MPI_send(b,...)

Fig. 2.1 An example of calling a routine with mismatched formal and actual arguments.

There are several important language binding issues not addressed by this standard. This standard does not discuss the interoperability of message passing between languages. It is fully expected that many implementations will have such features, and that such features are a sign of the quality of the implementation.

2.5.1 FORTRAN 77 BINDING ISSUES

All MPI names have an MPI_ prefix, and all characters are capitals. Programs must not declare variables or functions with names beginning with the prefix, MPI_. This is mandated to avoid possible name collisions.

All MPI Fortran subroutines have a return code in the last argument. A few MPI operations are functions, which do not have the return code argument. The return code value for successful completion is MPI_SUCCESS. Other error codes are implementation dependent; see Chapter 7.

Handles are represented in Fortran as INTEGERs. Binary-valued variables are of type LOGICAL.

Array arguments are indexed from one.

Unless explicitly stated, the MPI F77 binding is consistent with ANSI standard Fortran 77. There are several points where this standard diverges from the ANSI Fortran 77 standard. These exceptions are consistent with common practice in the Fortran community. In particular:

- MPI identifiers are limited to thirty, not six, significant characters.
- MPI identifiers may contain underscores after the first character.
- An MPI subroutine with a choice argument may be called with different argument types. An example is shown in Figure 2.1. This violates the letter of the Fortran standard, but such a violation is common practice. An alternative would be to have a separate version of MPI_SEND for each data type.
- Although not required, it is strongly suggested that named MPI constants (PARAMETERS) be provided in an include file, called mpi.h. On systems that do not support include files, the implementation should specify the values of named constants.
- Vendors are encouraged to provide type declarations in the mpi.h file on Fortran systems that support user-defined types. One should define, if possible, the type MPI_ADDRESS, which is an INTEGER of the size needed...
to hold an address in the execution environment. On systems where type
definition is not supported, it is up to the user to use an \texttt{INTEGER}
of the right kind to represent addresses (i.e., \texttt{INTEGER*4} on a 32 bit machine,
\texttt{INTEGER*8} on a 64 bit machine, etc.).

2.5.2 C BINDING ISSUES

We use the ANSI C declaration format. All \texttt{MPI} names have an \texttt{MPI}
 prefixed, defined constants are in all capital letters, and defined types and functions have one
capital letter after the prefix. Programs must not declare variables or functions
with names beginning with the prefix, \texttt{MPI}. This is mandated to avoid possible
name collisions.

The definition of named constants, function prototypes, and type definitions
must be supplied in an include file \texttt{mpi.h}.

Almost all C functions return an error code. The successful return code will be
\texttt{MPI\_SUCCESS}, but failure return codes are implementation dependent. A few
C functions do not return values, so that they can be implemented as macros.

Type declarations are provided for handles to each category of opaque objects. Either a pointer or an integer type is used.

Array arguments are indexed from zero.

Logical flags are integers with value 0 meaning “false” and a non-zero value
meaning “true.”

Choice arguments are pointers of type \texttt{void*}.

Address arguments are of \texttt{MPI} defined type \texttt{MPI\_Aint}. This is defined to be
an int of the size needed to hold any valid address on the target architecture.

2.6 Processes

An \texttt{MPI} program consists of autonomous processes, executing their own code,
in an MIMD style. The codes executed by each process need not be identical.
The processes communicate via calls to \texttt{MPI} communication primitives. Typically, each process executes in its own address space, although shared-memory
implementations of \texttt{MPI} are possible. This document specifies the behavior of a
parallel program assuming that only \texttt{MPI} calls are used for communication. The
interaction of an \texttt{MPI} program with other possible means of communication
(e.g., shared memory) is not specified.

\texttt{MPI} does not specify the execution model for each process. A process can
be sequential, or can be multi-threaded, with threads possibly executing concurrently. Care has been taken to make \texttt{MPI} “thread-safe,” by avoiding the use
of implicit state. The desired interaction of \texttt{MPI} with threads is that concurrent
threads be all allowed to execute \texttt{MPI} calls, and calls be reentrant; a blocking
\texttt{MPI} call blocks only the invoking thread, allowing the scheduling of another
thread.

\texttt{MPI} does not provide mechanisms to specify the initial allocation of processes
to an \texttt{MPI} computation and their binding to physical processors. It is expected
that vendors will provide mechanisms to do so either at load time or at run time. Such mechanisms will allow the specification of the initial number of required processes, the code to be executed by each initial process, and the allocation of processes to processors. Also, the current proposal does not provide for dynamic creation or deletion of processes during program execution (the total number of processes is fixed), although it is intended to be consistent with such extensions. Finally, we always identify processes according to their relative rank in a group, that is, consecutive integers in the range $0 \ldots \text{groupsize}-1$.

### 2.7 Error Handling

MPI provides the user with reliable message transmission. A message sent is always received correctly, and the user does not need to check for transmission errors, time-outs, or other error conditions. In other words, MPI does not provide mechanisms for dealing with failures in the communication system. If the MPI implementation is built on an unreliable underlying mechanism, then it is the job of the implementor of the MPI subsystem to insulate the user from this unreliability, or to reflect unrecoverable errors as failures. Whenever possible, such failures will be reflected as errors in the relevant communication call. Similarly, MPI itself provides no mechanisms for handling processor failures. The error handling facilities described in Section 7.2 can be used to restrict the scope of an unrecoverable error, or design error recovery at the application level.

Of course, MPI programs may still be erroneous. A **program error** can occur when an MPI call is called with an incorrect argument (non-existing destination in a send operation, buffer too small in a receive operation, etc.). This type of error would occur in any implementation. In addition, a **resource error** may occur when a program exceeds the amount of available system resources (number of pending messages, system buffers, etc.). The occurrence of this type of error depends on the amount of available resources in the system and the resource allocation mechanism used; this may differ from system to system. A high-quality implementation will provide generous limits on the important resources so as to alleviate the portability problem this represents.

Almost all MPI calls return a code that indicates successful completion of the operation. Whenever possible, MPI calls return an error code, if an error occurred during the call. By default, an error detected during the execution of the MPI library causes the parallel computation to abort. However, MPI provides mechanisms for users to change this default and to handle recoverable errors. The user may specify that no error is fatal, and handle error codes returned by MPI calls by himself or herself. Also, the user may provide his or her own error-handling routines, which will be invoked whenever an MPI call returns abnormally. The MPI error handling facilities are described in Section 7.2.

Several factors limit the ability of MPI calls to return with meaningful error codes when an error occurs. MPI may not be able to detect some errors; other errors may be too expensive to detect in normal execution mode; finally some
errors may be "catastrophic" and may prevent MPI from returning control to the caller in a consistent state.

Another subtle issue arises because of the nature of asynchronous communications: MPI calls may initiate operations that continue asynchronously after the call returned. Thus, the operation may return with a code indicating successful completion, yet later cause an error exception to be raised. If there is a subsequent call that relates to the same operation (e.g., a call that verifies that an asynchronous operation has completed) then the error argument associated with this call will be used to indicate the nature of the error. In a few cases, the error may occur after all calls that relate to the operation have completed, so that no error value can be used to indicate the nature of the error (e.g., an error in a send with the ready mode). Such an error must be treated as fatal, since information cannot be returned for the user to recover from it.

This document does not specify the state of a computation after an erroneous MPI call has occurred. The desired behavior is that a relevant error code be returned, and the effect of the error be localized to the greatest possible extent. E.g., it is highly desirable that an erroneous receive call will not cause any part of the receiver's memory to be overwritten, beyond the area specified for receiving the message.

Implementations may go beyond this document in supporting in a meaningful manner MPI calls that are defined here to be erroneous. For example, MPI specifies strict type matching rules between matching send and receive operations: it is erroneous to send a floating point variable and receive an integer. Implementations may go beyond these type matching rules, and provide automatic type conversion in such situations. It will be helpful to generate warnings for such nonconforming behavior.

2.8 Implementation Issues

There are a number of areas where an MPI implementation may interact with the operating environment and system. While MPI does not mandate that any services (such as I/O or signal handling) be provided, it does strongly suggest the behavior to be provided if those services are available. This is an important point in achieving portability across platforms that provide the same set of services.

2.8.1 INDEPENDENCE OF BASIC RUNTIME ROUTINES

MPI programs require that library routines that are part of the basic language environment (such as data and write in Fortran and print and malloc in ANSI C) and are executed after MPI_INIT and before MPI_FINALIZE operate independently and that their completion is independent of the action of other processes in an MPI program.

Note that this in no way prevents the creation of library routines that provide parallel services whose operation is collective. However, the following program
is expected to complete in an ANSI C environment regardless of the size of
_MPI_COMM_WORLD (assuming that I/O is available at the executing nodes).

    int rank;
    MPI_Init( argc, argv );
    MPI_Comm_rank( MPI_COMM_WORLD, &rank );
    if (rank == 0) printf( "Starting program\n" );
    MPI_Finalize();

The corresponding Fortran 77 program is also expected to complete.

An example of what is not required is any particular ordering of the action
of these routines when called by several tasks. For example, MPI makes neither
requirements nor recommendations for the output from the following program
(again assuming that I/O is available at the executing nodes).

    MPI_Comm_rank( MPI_COMM_WORLD, &rank );
    printf( "Output from task rank %d\n", rank );

In addition, calls that fail because of resource exhaustion or other error are
not considered a violation of the requirements here (however, they are required
to complete, just not to complete successfully).

2.8.2 INTERACTION WITH SIGNALS IN POSIX

MPI does not specify either the interaction of processes with signals, in a UNIX
environment, or with other events that do not relate to MPI communication.
That is, signals are not significant from the viewpoint of MPI, and implementors
should attempt to implement MPI so that signals are transparent: an MPI call
suspended by a signal should resume and complete after the signal is handled.
Generally, the state of a computation that is visible or significant from the viewpoint
of MPI should only be affected by MPI calls.

The intent of MPI to be thread and signal safe has a number of subtle effects.
For example, on Unix systems, a catchable signal such as SIGALRM (an alarm
signal) must not cause an MPI routine to behave differently than it would have
in the absence of the signal. Of course, if the signal handler issues MPI calls or
changes the environment in which the MPI routine is operating (for example,
consuming all available memory space), the MPI routine should behave as appropriate
for that situation (in particular, in this case, the behavior should be
the same as for a multi-threaded MPI implementation).

A second effect is that a signal handler that performs MPI calls must not
interfere with the operation of MPI. For example, an MPI receive of any type that
occurs within a signal handler must not cause erroneous behavior by the MPI
implementation. Note that an implementation is permitted to prohibit the use
of MPI calls from within a signal handler, and is not required to detect such use.

It is highly desirable that MPI not use SIGALRM, SIGFPE, or SIGIOT. An imple-
mentation is required to clearly document all of the signals that the MPI imple-
mentation uses; a good place for this information is a Unix ‘man’ page on MPI.
3.1 Introduction

Sending and receiving of messages by processes is the basic MPI communication mechanism. The basic point-to-point communication operations are send and receive. Their use is illustrated in the example below.

```c
#include "mpi.h"
main( argc, argv )
int argc;
char **argv;
{
    char message[20];
    int myrank;
    MPI_Status status;
    MPI_Init( &argc, &argv );
    MPI_Comm_rank( MPI_COMM_WORLD, &myrank );
    if (myrank == 0) /* code for process zero */
    {
        strcpy(message,"Hello, there");
        MPI_Send( message, strlen(message), MPI_CHAR, 1, 99, MPI_COMM_WORLD);
    }
    else /* code for process one */
    {
        MPI_Recv( message, 20, MPI_CHAR, 0, 99, MPI_COMM_WORLD, &status);
        printf("received :\%s:\n", message);
    }
    MPI_Finalize();
}
```

In this example, process zero (myrank = 0) sends a message to process one using the send operation MPI_SEND. The operation specifies a send buffer in the sender memory from which the message data is taken. In the example above, the send buffer consists of the storage containing the variable message in the
memory of process zero. The location, size, and type of the send buffer are
specified by the first three parameters of the send operation. The message sent
will contain the 15 characters of this variable. In addition, the send operation
associates an envelope with the message. This envelope specifies the message
destination and contains distinguishing information that can be used by the
receive operation to select a particular message. The last three parameters of
the send operation specify the envelope for the message sent.

Process one (myrank = 1) receives this message with the receive operation
MPI_RECV. The message to be received is selected according to the value of
its envelope, and the message data is stored into the receive buffer. In the
example above, the receive buffer consists of the storage containing the string
message in the memory of process one. The first three parameters of the receive
operation specify the location, size, and type of the receive buffer. The next three
parameters are used for selecting the incoming message. The last parameter is
used to return information on the message just received.

The next sections describe the blocking send and receive operations. We dis-
cuss send, receive, blocking communication semantics, type matching require-
ments, type conversion in heterogeneous environments, and more general com-
munication modes. Nonblocking communication is addressed next, followed by
channel-like constructs and send-receive operations. We then consider general
datatypes that allow one to transfer efficiently heterogeneous and noncontiguous
data. We conclude with the description of calls for explicit packing and
unpacking of messages.

3.2 Blocking Send and Receive Operations
3.2.1 BLOCKING SEND
The syntax of the blocking send operation is given below.

MPI_SEND(buf, count, datatype, dest, tag, comm)

| IN  | buf          | initial address of send buffer (choice) |
| IN  | count        | number of elements in send buffer (nonnegative integer) |
| IN  | datatype     | datatype of each send buffer element (handle) |
| IN  | dest         | rank of destination (integer) |
| IN  | tag          | message tag (integer) |
| IN  | comm         | communicator (handle) |

int MPI_Send(void* buf, int count, MPI_Datatype datatype, int dest, int tag,
MPI_Comm comm)

MPI_SEND(BUF, COUNT, DATATYPE, DEST, TAG, COMM, IERROR)

<type> BUF(*)
INTEGER COUNT, DATATYPE, DEST, TAG, COMM, IERROR

The blocking semantics of this call are described in Section 3.4.