

**Installation and User's Guide to MPICH,
a Portable Implementation of MPI
Version 1.2.3
The globus2 device for Grids**

by

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Abstract

MPI (Message-Passing Interface) is a standard specification for message-passing libraries. MPICH is a portable implementation of the full MPI-1 specification for a wide variety of parallel and distributed computing environments. MPICH contains, along with the MPI library itself, a programming environment for working with MPI programs. The programming environment includes a portable startup mechanism, several profiling libraries for studying the performance of MPI programs, and an X interface to all of the tools. This document describes how to install and use the MPICH implementation of MPI.

This document describes how to obtain, install, and use MPICH [9], the portable implementation of the MPI Message-Passing Standard. This document describes version 1.2.3.

1 Introduction

MPICH is a freely available implementation of the MPI standard that runs on a wide variety of systems. The details of the MPICH implementation are described in [9]; related papers include [6] and [7].

Major Features of MPICH:

- Full MPI 1.2 compliance, *including* cancel of sends.
- MPMD programs (see Section 3.5) and heterogeneous clusters are supported.
- The MPI-2 standard C++ bindings are available for the MPI-1 functions.
- Both Fortran 77 and Fortran 90 bindings, including both ‘`mpif.h`’ and an MPI module.
- A Windows NT version is available as open source. The installation and use for this version is different; this manual covers only the Unix version of MPICH.
- Supports a wide variety of environments, including clusters of SMPs and massively parallel computers.
- Follows many (but not yet all) of GNU-recommended build and install targets, including VPATH.
- Parts of MPI-2 are also supported:
 - Most of MPI-IO is supported through the ROMIO implementation (See ‘`romio/README`’ for details).
 - Support for `MPI_INIT_THREAD` (but only for `MPI_THREAD_SINGLE` and `MPI_THREAD_FUNNELLED`).
 - Miscellaneous new `MPI_Info` and `MPI_Datatype` routines.
- MPICH also includes components of a parallel programming environment, including
 - Tracing and logfile tools based on the MPI profiling interface, including a scalable logfile format (SLOG).

- Parallel performance visualization tools (`upshot` and `jumpshot`).
- Extensive correctness and performance tests.
- Both large and small application examples.

1.1 Features of recent releases

Version 1.2.3 of MPICH is primarily a bug fix and increased portability release, particularly for Linux-based clusters.

New and improved in 1.2.3:

- Reorganized manuals, arranged by device and including a frequently asked questions section.
- Further improvements to the `ch_p4mpd` device.
- Improvements to the SMP support in the `ch_p4` device, thanks to Pete Wykcoff.
- Support for IA64 systems running Linux and for Mac OS X.
- Support for version 2 of Globus in the `globus2` device.
- Many bug fixes and code improvements. See www.mcs.anl.gov/mpi/mpich/r1_2_3changes.html for a complete list of changes.

New and improved in 1.2.2:

- A greatly improved `ch_p4mpd` device.
- Improved support for assorted Fortran 77 and Fortran 90 compilers, including compile-time evaluation of Fortran constants used in the MPICH implementation.
- An improved `globus2` device, providing better performance.
- A new `bproc` mode for the `ch_p4` device supports Scyld Beowulfs.
- Many TCP performance improvements for the `ch_p4` and `ch_p4mpd` devices, as well as
- Many bug fixes and code improvements. See www.mcs.anl.gov/mpi/mpich/r1_2_2changes.html for a complete list of changes.

1.2 Choosing the correct device and release

MPICH is designed to be ported and optimized for a variety of systems through implementations of an *abstract device interface* (ADI). Many different implementations of the ADI exist; each is called a *device*.

Workstation Networks, Beowulf Clusters, and Individual Workstations. The most important devices are `ch_p4` and `ch_p4mpd`. The `ch_p4` device is the most general and supports SMP nodes, MPMD programs, and heterogeneous collections of systems. The `ch_p4mpd` device (so far) supports only homogenous clusters of uniprocessors, but provides for far faster and more scalable startup.

Grids. The `globus2` uses Globus (www.globus.org) to provide a grid-enabled implementation of MPI. This device is appropriate for systems on which Globus is installed.

Symmetric Multiprocessors. The `ch_shmem` device is appropriate for a single shared-memory system, such as a SGI Origin or Sun E10000. This device uses shared memory to pass messages between processes, and makes use of facilities provided by the operating system such as System V shared memory or anonymous mmap regions for data and System V semaphores or OS-specific mutex routines for synchronization. The `ch_lfshmem` device is a version of `ch_shmem` that uses a lock-free approach and that we developed for the NEC SX-4 [6]. The `ch_shmem` is appropriate for most systems; the `ch_lfshmem` requires special assembly language coding but can be ported to most systems. Currently, however, the `ch_lfshmem` supports only the NEC SX-4.

Massively Parallel Processors (MPPs). MPICH was originally developed to provide an implementation of MPI for many of the massively parallel processors, each of which had its own, proprietary, message-passing system. We still include the devices `ch_meiko`, `ch_nx` and `ch_mpl` with MPICH releases, though most of these systems have now disappeared.

Others. MPICH was designed to enable other groups to use it when developing their own MPI implementations. Both vendors and research groups have used MPICH as the basis for their implementation. One important one in the cluster environment is MPICH-GM, for Myrinet switch-connected clusters; it is available directly from Myricom.

To make it easier to build and use MPICH, customized versions of this manual are constructed for each major device.

1.3 Citations and References

The name MPICH is derived from MPI and Chameleon; Chameleon both because MPICH can run (adapt its color) on a wide range of environments and because the initial implementation of MPICH used the Chameleon [13] message-passing portability system. MPICH is pronounced “Em Pee Eye See Aych,” not “Emm Pitch.” The proper references for MPICH are the paper in the journal *Parallel Computing* [9] and this manual [8]. BibTeX versions of these citations are available at www.mcs.anl.gov/mpi/mpich/mpich-prefer-ref.html.

2 Quick Start

Here is a set of steps for setting up and minimally testing MPICH. Details and instructions for a more thorough tour of MPICH’s features, including installing, validating, benchmarking, and using the performance evaluation tools, are given in the following sections.

2.1 Downloading MPICH

The first step is to download MPICH and install any necessary patches.

1. The easiest way to get MPICH is to use the web page www.mcs.anl.gov/mpi/mpich/download.html; you can also use anonymous ftp from [ftp.mcs.anl.gov](ftp://ftp.mcs.anl.gov) in directory 'pub/mpi'. Get the file 'mpich.tar.gz'. (If that file is too big, try getting the pieces from pub/mpi/mpisplit and catting them together.)
2. Unpack the 'mpich.tar.gz' file into a build directory. We recommend using a locally mounted partition rather than an NFS (network file system) partition. For example, on many systems, '/tmp' or '/sandbox' are locally mounted. Make sure that there is enough space available (100MB should be more than enough). To unpack, assuming that 'mpich.tar.gz' has been downloaded into '/tmp', use

```
% cd /tmp
% tar zxovf mpich.tar.gz
```

If your tar does not accept the z option, use

```
% cd /tmp
% gunzip -c mpich.tar.gz | tar zxovf -
```

3. Apply any patches. Check the web page www.mcs.anl.gov/mpi/mpich/buglist-tbl.html for any patches that need to be applied. Normally, versions of MPICH that already have these important patches applied are available; they are indicated by a fourth number in the release name (e.g., 1.2.2.3). But in some cases, a patch is made available early. The patch page has instructions on applying the patches.

Now you are ready to build MPICH.

2.2 Configuring, Making, and Installing

Before you can use MPICH, you must configure and make it. The configuration process analyzes your system and determines the correct options and settings; it also creates the 'Makefile's that are used to make MPICH.

1. Decide where you want to install MPICH. This step is not strictly necessary; however, installing MPICH (which can be done without any privileges in a user directory) both makes it easier to manage updates to MPICH and allows you to reduce the amount disk space that MPICH takes up, since the installed version contains only the libraries, header files, documentation, and supporting programs. We recommend an install path that contains the version number of MPICH. For example, if you are installing MPICH for others to use, and you have the required access rights, you could choose '/usr/local/mpich-1.2.3/'. If you are installing it just for your own use, you could use something like '/home/me/software/mpich-1.2.3'.

2. Invoke `configure` with the appropriate `prefix`:

```
% ./configure --prefix=/usr/local/mpich-1.2.3 |& tee c.log
```

This will configure MPICH for the default device; this is usually the appropriate choice. Section 4.1 discusses the options that can be given to `configure` to customize MPICH.

The output of `configure` is piped to `tee`; this program both writes the output to the file specified by its argument (here `'c.log'`) and to standard output. If you have trouble with the `configure` or `make` step, the file `'c.log'` will help identify any problems.

3. Make MPICH:

```
% make |& tee make.log
```

This may take a while, depending on the load on your system and on your file server, it may take anywhere from a few minutes to an hour or more.

2.3 Running examples

1. (Optional) Build and run a simple test program:

```
% cd examples/basic
% make cpi
% ../../bin/mpirun -np 4 cpi
```

At this point you have run an MPI program on your system. If you have trouble, see Section 6.

2. (Optional) Put the distribution through its complete acceptance test (See Section 4.11 for how to do this).
3. (Optional) If you wish to install MPICH in a public place so that others may use it, use

```
% make install
```

to install MPICH into the directory specified by the `--prefix` option to `configure`. Installation will consist of an `'include'`, `'lib'`, `'bin'`, `'sbin'`, `'www'`, and `'man'` directories and a small `'examples'` directory. Should you wish to remove the installation, you can run the script `sbin/mpiuninstall`.

4. (Optional) At this point you can announce to your users how to compile and run MPI programs, using the installation you have just built in `'/usr/local/mpich-1.2.3/'` (or wherever you have installed it). See Section 3 for commands they can use. They can also copy the `'Makefile'` in `'/usr/local/mpich-1.2.3/examples'` and adapt it for their own use.

2.4 Sample MPI programs

The MPICH distribution contains a variety of sample programs, which are located in the MPICH source tree. Most of these will work with any MPI implementation, not just MPICH.

examples/basic contains a few short programs in Fortran, C, and C++ for testing the simplest features of MPI.

examples/test contains multiple test directories for the various parts of MPI. Enter “make testing” in this directory to run our suite of function tests.

examples/perftest Performance benchmarking programs. See the script `runmpptest` for information on how to run the benchmarks. These are relatively sophisticated.

mpe/contrib/mandel A Mandelbrot program that uses the MPE graphics package that comes with mpich. It should work with any other MPI implementation as well, but we have not tested it. This is a good demo program if you have a fast X server and not too many processes.

mpe/contrib/mastermind A program for solving the Mastermind puzzle in parallel. It can use graphics (`gmm`) or not (`mm`).

Additional examples from the book *Using MPI* [10] are available at www.mcs.anl.gov/mpi/using. Tutorial material on MPI can also be found at www.mcs.anl.gov/mpi.

3 Programming Tools

The MPICH implementation comes with a variety of tools for building, running, debugging, and analyzing MPI programs. This section details these tools.

3.1 Compiling, linking, and running programs

The MPICH implementation provides four commands for compiling and linking C (`mpicc`), C++ (`mpiCC`), Fortran 77 (`mpif77`), and Fortran 90 (`mpif90`) programs.

Use these commands just like the usual C, Fortran 77, C++, or Fortran compilers. For example,

```
mpicc -c foo.c
mpif77 -c foo.f
mpiCC -c foo.C
mpif90 -c foo.f
```

and

```
mpicc -o foo foo.o
mpif77 -o foo foo.o
mpiCC -o foo foo.o
mpif90 -o foo foo.o
```

Commands for the linker may include additional libraries. For example, to use routines from the C math library, use

```
mpicc -o foo foo.o -lm
```

Combining compilation and linking in a single command, as shown here,

```
mpicc -o foo foo.c
mpif77 -o foo foo.f
mpiCC -o foo foo.C
mpif90 -o foo foo.f
```

may also be used (on most systems).

Note that while the suffixes `.c` for C programs and `.f` for Fortran-77 programs are standard, there is no consensus for the suffixes for C++ and Fortran-90 programs. The ones shown here are accepted by many but not all systems. MPICH tries to determine the accepted suffixes, but may not always be able to.

You can override the choice of compiler by specifying the environment variable `MPICH_CC`, `MPICH_F77`, `MPICH_CCC`, or `MPICH_F90`. However, be warned that this will work only if the alternate compiler is compatible with the default one (by compatible, we mean that it uses the same sizes for all datatypes and layouts, and generates object code that can be used with the MPICH libraries). If you wish to override the linker, use the environment variables `MPICH_CLINKER`, `MPICH_F77LINKER`, `MPICH_CCLINKER`, or `MPICH_F90LINKER`.

If you want to see the commands that would be used without actually running them, add the command line argument `-show`.

In addition, the following special options are supported for accessing some of the features of the MPE environment for monitoring MPI calls from within an application:

-mpilog Build version that generates MPE log files.

-mpitrace Build version that generates traces.

-mpianim Build version that generates real-time animation.

These are described in more detail in Section 3.7.

3.1.1 Compiling and Linking without the Scripts

In some cases, it is not possible to use the scripts supplied by MPICH for compiling and linking programs. For example, another tool may have its own compilation scripts. In this

case, you can use `-compile_info` and `-link_info` to have the MPICH compilation scripts indicate the compiler flags and linking libraries that are required for correct operation of the MPICH routines. For example, when using the `ch_shmem` device on Solaris systems, the library `thread` (`-lthread`) must be linked with the application. If the `thread` library is not provided, the application will still link, but essential routines will be replaced with dummy versions contained within the Solaris C library, causing the application to fail.

For example, to determine the flags used to compile and link C programs, you can use these commands, whose output for the `ch_p4` device on a Linux workstation is shown.

```
% mpicc -compile_info
cc -DUSE_STDARG -DHAVE_STDLIB_H=1 -DHAVE_STRING_H=1 -DHAVE_UNISTD_H=1
-DHAVE_STDARG_H=1 -DUSE_STDARG=1 -DMALLOC_RET_VOID=1
-I/usr/local/mpich/include -c

% mpicc -link_info
cc -DUSE_STDARG -DHAVE_STDLIB_H=1 -DHAVE_STRING_H=1 -DHAVE_UNISTD_H=1
-DHAVE_STDARG_H=1 -DUSE_STDARG=1 -DMALLOC_RET_VOID=1
-L/usr/local/mpich/lib -lmpich
```

3.2 Running programs with `mpirun`

To run an MPI program, use the `mpirun` command, which is located in `‘/usr/local/mpich-1.2.3/bin’`. For almost all systems, you can use the command

```
mpirun -np 4 a.out
```

to run the program `‘a.out’` on four processors. The command `mpirun -help` gives you a complete list of options, which may also be found in Appendix E.

On exit, `mpirun` returns the status of one of the processes, usually the process with rank zero in `MPI_COMM_WORLD`.

3.3 `mpirun` and Globus

In the section we describe how to run MPI programs using the MPICH `globus2` device in a Globus-enabled distributed computing environment. It is assumed that (a) Globus has been installed and the appropriate Globus daemons are running on the machines you wish to launch your MPI application, (b) you have already acquired your Globus ID and security credentials, (c) your Globus ID has been registered on all machines, and (d) you have a valid (unexpired) Globus proxy. See <http://www.globus.org> for information on those topics.

Every `mpirun` command under the `globus2` device submits a Globus *Resource Specification Language Script*, or simply *RSL script*, to a Globus-enabled grid of computers. Each RSL script is composed of one or more RSL *subjobs*, typically one subjob for each machine in the computation. You may supply your own RSL script¹ explicitly to `mpirun` (using

¹See www.globus.org for the syntax and semantics of the Globus Resource Specification Language.

the `-globusrsl <rslfilename>` option) in which case you would not specify any other options to `mpirun`, or you may have `mpirun` construct an RSL script for you based on the arguments you pass to `mpirun` and the contents of your *machines* file (discussed below). In either case it is important to remember *communication between nodes in different subjobs is always facilitated over TCP/IP and the more efficient vendor-supplied MPI is used only among nodes within the same subjob*.

3.4 Using `mpirun` To Construct An RSL Script For You

You would use this method if you wanted to launch a *single* executable file, which implies a set of one or more binary-compatible machines that all share the same filesystem (i.e., they can all access the executable file).

Using `mpirun` to construct an RSL script for you requires a *machines* file. The `mpirun` command determines which machines file to use as follows:

1. If a `-machinefile <machinefilename>` argument is specified on the `mpirun` command, it uses that; otherwise,
2. it looks for a file `'machines'` in the directory in which you typed `mpirun`; and finally,
3. it looks for `'/usr/local/mpich/bin/machines'` where `'/usr/local/mpich'` is the MPICH installation directory.

If it cannot find a machines file from any of those places then `mpirun` fails.

The machines file is used to list the computers upon which you wish to run your application. Computers are listed by naming the Globus “service” on that machine. For most applications the default service can be used, which requires specifying only the fully qualified domain name. Consult your local Globus administrator or the Globus web site www.globus.org for more information regarding special Globus services. For example, consider the following pair of binary-compatible machines, `{m1,m2}.utech.edu`, that have access to the same filesystem. Here is what a machines file that uses default Globus services might look like.

```
"m1.utech.edu" 10
"m2.utech.edu" 5
```

The number appearing at the end of each line is optional (default=1). It specifies the maximum number of nodes that can be created in a single RSL subjob on each machine. `mpirun` uses the `-np` specification by “wrapping around” the machines file. For example, using the machines file above `mpirun -np 8` creates an RSL with a single subjob with 8 nodes on `m1.utech.edu`, while `mpirun -np 12` creates two subjobs where the first subjob has 10 nodes on `m1.utech.edu` and the second has 2 nodes on `m2.utech.edu`, and finally `mpirun -np 17` creates three subjobs with 10 nodes on `m1.utech.edu` followed by 5 nodes on `m2.utech.edu` ending with the third a final subjob having two nodes on `m1.utech.edu` again. Note that inter-subjob messaging is *always* communicated over TCP, even if the two separate subjobs are the same machine.

3.4.1 Using mpirun By Supplying Your Own RSL Script

You would use `mpirun` supplying your own RSL script if you were submitting to a set of machines that could not run or access the same executable file (e.g., machines that are not binary compatible and/or do not share a file system). In this situation, we must currently use something called a *Resource Specification Language* (RSL) request to specify the executable filename for each machine. This technique is very flexible, but rather complex; work is currently underway to simplify the manner in which these issues are addressed.

The easiest way to learn how to write your own RSL request is to study the one generated for you by `mpirun`. Consider the example where we wanted to run an application on a cluster of workstations. Recall our `machines` file looked like this:

```
"m1.utech.edu" 10
"m2.utech.edu" 5
```

To view the RSL request generated in this situation, *without* actually launching the program, we type the following `mpirun` command:

```
% mpirun -dumprsl -np 12 myapp 123 456
```

which produces the following output:

```
+
( &(resourceManagerContact="m1.utech.edu")
  (count=10)
  (jobtype=mpi)
  (label="subjob 0")
  (environment=(GLOBUS_DUROC_SUBJOB_INDEX 0))
  (arguments=" 123 456")
  (directory=/homes/karonis/MPI/mpich.yukon/mpich/lib/IRIX64/globus)
  (executable=/homes/karonis/MPI/mpich.yukon/mpich/lib/IRIX64/globus/myapp)
)
( &(resourceManagerContact="m2.utech.edu")
  (count=2)
  (jobtype=mpi)
  (label="subjob 1")
  (environment=(GLOBUS_DUROC_SUBJOB_INDEX 1))
  (arguments=" 123 456")
  (directory=/homes/karonis/MPI/mpich.yukon/mpich/lib/IRIX64/globus)
  (executable=/homes/karonis/MPI/mpich.yukon/mpich/lib/IRIX64/globus/myapp)
)
```

Note that `(jobtype=mpi)` may appear only in those subjobs whose machines have vendor-supplied implementations of MPI. Additional environment variables may be added as in the example below:

```
+
( &(resourceManagerContact="m1.utech.edu")
  (count=10)
```

```

(jobtype=mpi)
(label="subjob 0")
(environment=(GLOBUS_DUROC_SUBJOB_INDEX 0)
              (MY_ENV 246))
(arguments=" 123 456")
(directory=/homes/karonis/MPI/mpich.yukon/mpich/lib/IRIX64/globus)
(executable=/homes/karonis/MPI/mpich.yukon/mpich/lib/IRIX64/globus/myapp)
)
( &(resourceManagerContact="m2.utech.edu")
  (count=2)
  (jobtype=mpi)
  (label="subjob 1")
  (environment=(GLOBUS_DUROC_SUBJOB_INDEX 1))
  (arguments=" 123 456")
  (directory=/homes/karonis/MPI/mpich.yukon/mpich/lib/IRIX64/globus)
  (executable=/homes/karonis/MPI/mpich.yukon/mpich/lib/IRIX64/globus/myapp)
)

```

After editing your own RSL file you may submit that directly to `mpirun` as follows:

```
% mpirun -globusrsl <myrslrequestfile>
```

Note that when supplying your own RSL it should be the *only* argument you specify to `mpirun`.

RSL is a flexible language capable of doing much more than has been presented here. For example, it can be used to stage executables and to set environment variables on remote computers before starting execution. A full description of the language can be found at <http://www.globus.org>.

3.5 MPMD Programs

It is possible to run a parallel program with different executables with several of the devices including the `ch_p4`, `ch_mpl`, and `globus2` devices. It is currently not possible to do this with the `ch_shmem` or `ch_p4mpd` devices. This style of parallel programming is often called MPMD for “multiple program multiple data”. In many cases, it is easy to convert a MPMD program into a single program that uses the rank of the process to invoke a different routine; doing so makes it easier to start parallel programs and often to debug them. If converting a MPMD program to a SPMD (single program multiple data, not to be confused with single *instruction* multiple data, or SIMD) is not feasible, then you can run MPMD programs using MPICH. However, you will not be able to use `mpirun` to start the programs; instead, you will need to follow the instructions for each device.

3.6 Debugging

Debugging parallel programs is notoriously difficult. Parallel programs are subject not only to the usual kinds of bugs but also to new kinds having to do with timing and synchronization errors. Often, the program “hangs,” for example when a process is waiting for a message to arrive that is never sent or is sent with the wrong tag. Parallel bugs often

disappear precisely when you add code to try to identify the bug, which is particularly frustrating. In this section we discuss several approaches to parallel debugging.

3.6.1 The printf Approach

Just as in sequential debugging, you often wish to trace interesting events in the program by printing trace messages. Usually you wish to identify a message by the rank of the process emitting it. This can be done explicitly by putting the rank in the trace message.

3.6.2 Error handlers

The MPI Standard specifies a mechanism for installing one's own error handler, and specifies the behavior of two predefined ones, `MPI_ERRORS_RETURN` and `MPI_ERRORS_ARE_FATAL`. As part of the MPE library, we include two other error handlers to facilitate the use of command-line debuggers such as `dbx` in debugging MPI programs.

```
MPE_Errors_call_dbx_in_xterm
MPE_Signals_call_debugger
```

These error handlers are located in the MPE directory. A configure option (`-mpedbg`) includes these error handlers into the regular MPI libraries, and allows the command-line argument `-mpedbg` to make `MPE_Errors_call_dbx_in_xterm` the default error handler (instead of `MPI_ERRORS_ARE_FATAL`).

3.6.3 Starting jobs with a debugger

The `-dbg=<name of debugger>` option to `mpirun` causes processes to be run under the control of the chosen debugger. For example, enter

```
mpirun -dbg=gdb or mpirun -dbg=gdb a.out
```

invokes the `mpirun_dbg.gdb` script located in the `'mpich/bin'` directory. This script captures the correct arguments, invokes the `gdb` debugger, and starts the first process under `gdb` where possible. There are five debugger scripts; `ddd`, `gdb`, `xxgdb`, `ddd`, and `totalview`. These may need to be edited depending on your system. There is another debugger script for `dbx`, but this one will always need to be edited as the debugger commands for `dbx` varies between versions. You can also use this option to call another debugger; for example, `-dbg=mydebug`. All you need to do is write a script file, `'mpirun_dbg.mydebug'`, which follows the format of the included debugger scripts, and place it in the `'mpich/bin'` directory. More information on using the Totalview debugger with MPICH can be found in Section 3.6.6.

3.6.4 Starting the debugger when an error occurs

It is often convenient to have a debugger start when a program detects an error. If MPICH was configured with the option `--enable-mpedbg`, then adding the command-line option

`-mpedbg` to the program will cause MPICH to attempt to start a debugger (usually `dbx` or `gdb`) when an error that generates a signal (such as `SIGSEGV`) occurs. For example,

```
mpirun -np 4 a.out -mpedbg
```

If you are not sure if your MPICH provides this service, you can use `-mpiversion` to see if MPICH was built with the `--enable-mpedbg` option. This feature may not be available with all devices.

3.6.5 Attaching a debugger to a running program

On workstation clusters, you can often attach a debugger to a running process. For example, the debugger `dbx` often accepts a process id (pid) which you can get by using the `ps` command. The form may be either

```
dbx a.out 1234
```

or

```
dbx -pid 1234 a.out
```

where 1234 is the process id².

To do this with `gdb`, start `gdb` and at its prompt do

```
file a.out
attach 1234
```

One can also attach the TotalView debugger to a running program (See Section 3.6.6 below).

3.6.6 Debugging MPI programs with TotalView

TotalView© [1] is a powerful, commercial-grade, portable debugger for parallel and multithreaded programs, available from Etnus (<http://www.etnus.com/>). TotalView understands multiple MPI implementations, including MPICH. By “understand” is meant that if you have TotalView installed on your system, it is easy to start your MPICH program under the control of TotalView, even if you are running on multiple machines, manage your processes both collectively and individually through TotalView’s convenient GUI, and even examine internal MPICH data structures to look at message queues [3]. The general operation model of TotalView will be familiar to users of command-line-based debuggers such as `gdb` or `dbx`.

²The unnecessary changes in the command syntax for programs such as `dbx` contributes to the poor state of software.

Starting an MPICH program under TotalView control. To start a parallel program under TotalView control, simply add ‘-dbg=totalview’ to your `mpirun` arguments:

```
mpirun -dbg=totalview -np 4 cpi
```

TotalView will come up and you can start the program by typing ‘G’. A window will come up asking whether you want to stop processes as they execute `MPI_Init`. You may find it more convenient to say “no” and instead to set your own breakpoint after `MPI_Init` (see Section 3.6.6). This way when the process stops it will be on a line in your program instead of somewhere inside `MPI_Init`.

Attaching to a running program. TotalView can attach to a running MPI program, which is particularly useful if you suspect that your code has deadlocked. To do this start TotalView with no arguments, and then press ‘N’ in the root window. This will bring up a list of the processes that you can attach to. When you dive through the initial MPICH process in this window TotalView will also acquire all of the other MPICH processes (even if they are not local). See the TotalView manual for more details of this process.

Debugging with TotalView. You can set breakpoints by clicking in the left margin on a line number. Most of the TotalView GUI is self-explanatory. You select things with the left mouse button, bring up an action menu with the middle button, and “dive” into functions, variables, structures, processes, etc., with the right button. Pressing `ctrl-?` in any TotalView window brings up help relevant to that window. In the initial TotalView window it brings up general help. The full documentation (*The TotalView User’s Guide*) is available from the Etnus web site.

You switch from viewing one process to the next with the arrow buttons at the top-right corner of the main window, or by explicitly selecting (left button) a process in the root window to re-focus an existing window onto that process, or by diving (right button) through a process in the root window to open a new window for the selected process. All the keyboard shortcuts for commands are listed in the menu that is attached to the middle button. The commands are mostly the familiar ones. The special one for MPI is the ‘m’ command, which displays message queues associated with the process.

Note also that if you use the MPI-2 function `MPI_Comm_set_name` on a communicator, TotalView will display this name whenever showing information about the communicator, making it easier to understand which communicator is which.

3.7 Log and tracefile tools

The MPE libraries that are distributed with MPICH contain several libraries for either logging information about the execution of each MPI call to a file for later analysis or for tracing each call as it occurs. These libraries may be accessed by simply providing a command line argument to the compilation scripts; further, this needs only be done when linking the program. For example, to create a log file of a program such as `cpi`, the following steps are needed:

```
mpicc -c cpi.c
mpicc -o cpi -mpilog cpi.o
```

The log file will be written to a file with the name ‘cpi.clog’ or ‘cpi.slog’, depending on the value of the environment variable `MPE_LOG_FORMAT` (clog is the default). The logfile can be graphically displayed using the Jumpshot program, described in Section 3.7.1.

Another tool of interest is FPMPI [?]; this tool reports a summary of all MPI calls, providing information about message lengths, communication patterns.

3.7.1 Jumpshot

Jumpshot is a program for displaying logfiles produced using the MPE logging libraries. Jumpshot is described in more detail in the *MPE Installation and User’s manual* [?]. Jumpshot is a Java program and requires a functioning Java environment. If you are lucky enough to have one, then you can view a logfile such as ‘cpi.clog’ in the example above by using

```
jumpshot cpi.clog
```

3.8 Execution tracing

Execution tracing is easily accomplished using the `-mpitrace` command line argument while linking:

```
mpicc -c cpi.c
mpicc -o cpi -mpitrace cpi.o
```

3.9 Performance measurements

The ‘mpich/examples/perftest’ directory contains a sophisticated tool for measuring latency and bandwidth for MPICH on your system. Simply change to the `mpich/examples/perftest` directory and do

```
make
mpirun -np 2 mptest -gnuplot > out.gpl
```

The file ‘out.gpl’ will then contain the necessary `gnuplot` commands. The file ‘mptest.gpl’ will contain the data. To view the data with `gnuplot`, use:

```
gnuplot out.gpl
```

or use

```
load 'out.gpl'
```

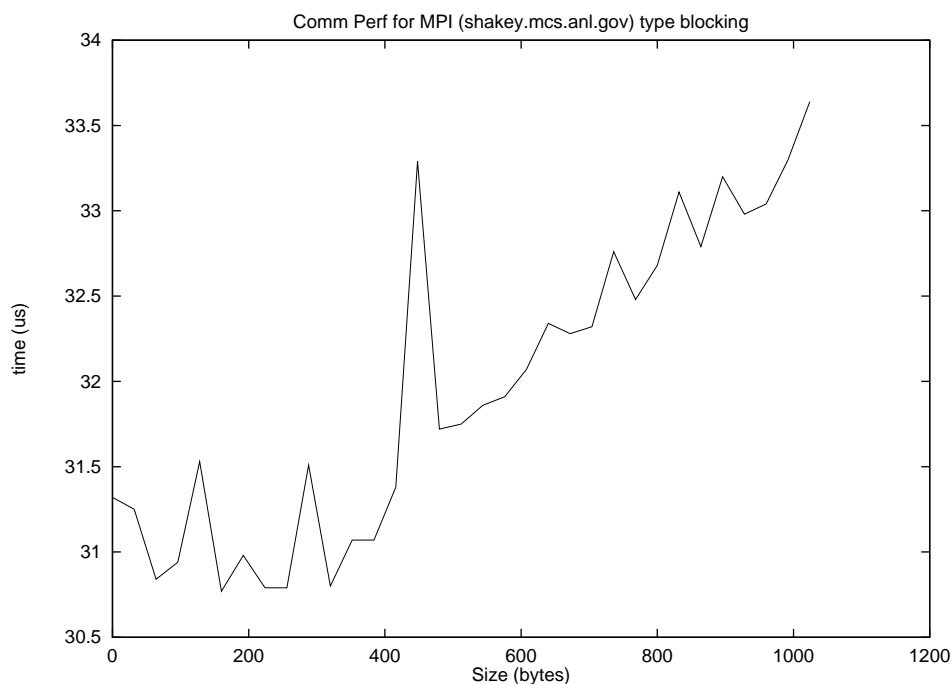


Figure 1: Sample output from `mpptest`

from within `gnuplot`. Depending on your environment and version of `gnuplot`, you may need to start `gnuplot` first and issue the command `set terminal x11` before executing `'load 'out.gpl'`. You can use

```
gnuplot
set term postscript eps
set output "foo.eps"
load 'out.gpl'
```

to create an Encapsulated Postscript graph such as the one in Figure 1.

The programs `mpptest` and `goptest` have a wide variety of capabilities; the option `-help` will list them. For example, `mpptest` can automatically pick message lengths to discover any sudden changes in behavior and can investigate the ability to overlap communication with computation. These programs are written using MPI, and may be used with *any* MPI implementation, not just MPICH. (See the `configure` file in the `'examples/perftest'` directory.) More information is available at <http://www.mcs.anl.gov/mpi/mpptest>.

Benchmarking can be very tricky. Some common benchmarking errors are discussed at <http://www.mcs.anl.gov/mpi/mpptest/hownot.html>. The paper [12] discusses these issues at more length.

4 Details

This section covers details that you should not normally need. Read this section if you need more detailed control over configuring, building, installing, or operating MPICH.

4.1 Configure options

The configure script documents itself in the following way. If you type

```
configure -usage
```

you will get a complete list of arguments and their meanings; these are also shown in Appendix D. The most important options are

- `--prefix=dir` The installation prefix. `configure` understands all of the usual GNU installation directory arguments, including `--libdir` and `--mandir`. We recommend that all users specify an installation directory with `--prefix`.
- `--with-device=devname` Set the MPICH device to use. `devname` must be the name of one of the directories in the ‘mpid’ directory, such as `ch_p4`, `ch_shmem`, `globus2`, or `ch_p4mpd`.
- `--enable-g` Add the `-g` option to the compile scripts. This is a prerequisite for using most debuggers, including `dbx`, and `gdb`.
- `--enable-debug` Turn on support for the Totalview© Debugger. This allows Totalview to display information on message queues.
- `--enable-sharedlib` Build both static and shared libraries for MPICH. This supports only a few systems, including those using `gcc` (e.g., most Linux Beowulf systems).
- `-automountfix=program` This is sometimes necessary for systems with automounter problems (see Section ??).

In addition, `configure` makes use of environment variables such as `MAKE`, `CC`, `FC`, `CFLAGS`, and `FFLAGS`.

Normally, you should use `configure` with as few arguments as you can. If you leave all arguments off, `configure` will usually guess the correct architecture (`arch`) unless you are in a cross-compiling environment, and will usually choose an appropriate device (`device`) as well. Where TCP/IP is an appropriate mechanism for communication, the TCP device (`ch_p4`) will be chosen by default.

4.1.1 MPI and PMPI routines

The MPI standard requires that each routine be available with both the MPI and PMPI prefix; for example, `MPI_Send` and `PMPI_Send`. MPICH attempts to use *weak symbols* to provide this feature; this reduces the size of the MPICH library. You can force MPICH to

make separate libraries for the MPI and PMPI versions by adding the configure option `--disable-weak-symbols`:

```
configure --disable-weak-symbols ...
```

Some MPI routines are implemented in terms of other MPI routines. For example, in MPICH, `MPI_Bcast` is implemented using `MPI_Send`. When weak symbols are used, even the PMPI versions of the routines are implemented using the MPI (not PMPI) versions. If you want the PMPI routines to only use the PMPI routines, use `--disable-weak-symbols` when configuring MPICH. Note that this behavior may change in later releases.

4.1.2 Configuring MPICH for use with threads

The MPICH implementation of MPI is currently not threadsafe. It may, however, be possible to use MPICH in a threaded application as long as all MPICH calls are made by a single thread. An example of this is OpenMP used for loop parallelism, combined with MPI.

4.1.3 Signals

In general, users should avoid using signals with MPI programs. The manual page for `MPI_Init` describes the signals that are used by the MPI implementation; these should not be changed by the user.

Because Unix does not chain signals, there is the possibility that several packages will attempt to use the same signal, causing the program to fail. For example, by default, the `ch_p4` device uses `SIGUSR1`; some thread packages also use `SIGUSR1`.

In a few cases, you can change the signal *before* calling `MPI_Init`. In those cases, your signal handler will be called after the MPICH implementation acts on the signal. For example, if you want to change the behavior of `SIGSEGV` to print a message, you can establish such a signal handler before calling `MPI_Init`. With devices such as the `ch_p4` device that handle `SIGSEGV`, this will cause your signal handler to be called after MPICH processes it.

4.2 Computational Grids with the globus2 device

Before configuring for the `globus2` device, a version of Globus must already be installed³. You will need to know the directory where Globus is installed (e.g., `/usr/local/globus`). Set the environment `GLOBUS_INSTALL_PATH` to that directory, for example,

```
setenv GLOBUS_INSTALL_PATH /usr/local/globus
```

When configuring for the `globus2` device, you may specify one of the Globus *flavors* (e.g., `mpi`, `debug` or `nodebug`, `threads`, `32-` or `64-bit`, etc.). To see the complete list of *all* Globus flavors (not all may be installed on your machine) use

```
$GLOBUS_INSTALL_PATH/bin/globus-development-path -help
```

³See <http://www.globus.org> for instructions regarding acquiring and installing Globus.

The flavors that are available to you (i.e., installed on your machine) are enumerated as directories in `$GLOBUS_INSTALL_PATH/development`. For example, Globus installation on a Solaris workstation might have the following flavors:

```
sparc-sun-solaris2.7_nothreads_standard_debug/  
sparc-sun-solaris2.7_pthreads_standard_debug/  
sparc-sun-solaris2.7_solaristhreads_standard_debug/
```

There are two ways to configure for the `globus2` device. Each method selects one of the Globus flavor directories in `$GLOBUS_INSTALL_PATH/development`. The first method is to specify the flavor directory *explicitly*, for example (all on one line):

```
configure --with-device=globus2:-dir=$GLOBUS_INSTALL_PATH/development/sparc-sun-solaris2.7_nothreads_standard_debug
```

Optionally, you may specify the flavor directory *implicitly*,

```
configure --with-device=globus2:-flavor=nothreads,debug
```

Finally, you may simply choose the default flavor (returned by `$GLOBUS_INSTALL_PATH/bin/globus-development-path`)

```
configure --with-device=globus2
```

You must specify `-mpi` to enable vendor-supplied MPI communication for intra-machine messaging. In other words, when configuring on machines that provide vendor implementations of the MPI standard, you must specify `-mpi` for optimal performance. Failing to specify `-mpi` will result in TCP intra-machine communication.

Selecting `-debug` can be helpful during debugging, but can slow down performance. `-nodebug` should be used for debugged production code.

In general, `-nothreads` should be used (the Globus2 device is not multithreaded). You should select a threaded flavor only if you intend to link your MPI application with other modules that require a threaded version of Globus (e.g., you have written a library that uses Nexus which requires threaded handlers). You should *not* select a threaded version of Globus simply because your MPI application is multithreaded.

When Globus was installed, a special ‘Makefile’ was automatically generated just for MPICH. The MPICH configure uses that file when configuring for the `globus2` device. That special ‘Makefile’ contains virtually all the information MPICH configure needs (include directory paths, special libraries, the names of C and Fortran compiler and linkers, etc.).

4.3 Alternate C Compilers

More and more systems, particularly clusters, come with multiple compilers. In many cases, you can build MPICH using just one of these compilers and then allow users to use

their favorite compiler when building their code. In many cases, no extra steps are needed. Users can simply use the command-line argument `-cc` or the environment variable `MPICH_CC` to specify a different compiler.

Unfortunately, this won't always work. For example, in some cases, different libraries may be needed by different compilers when linking programs. Some compilers may need different options to support ANSI/ISO C. In particular, support for the longer datatypes such as `long long` and `long double` may rely on runtime library support routines that are specific to each compiler. To handle all of these cases, For C and C++, you can create a file with the name `'mpicc-<compilename>.conf'`, e.g., `'mpicc-gcc.conf'` or `'mpicc-pgcc.conf'`, that contains any specifications that are needed by the `mpicc` command. `mpiCC` uses `'mpiCC-<name>.conf'`. Currently, these files must be created by hand, starting from the file `'mpichlib.conf'` that is created in the `'mpich/util'` directory by the `MPICH configure`. The simplest way to create each file is to use `configure`:

```
setenv CC cc
configure --prefix=/usr/local/mpich-1.2.3
make
make install
cp util/mpichlib.conf /usr/local/mpich-1.2.3/etc/mpicc-cc.conf
setenv CC gcc
configure --prefix=/usr/local/mpich-1.2.3
make
cp util/mpichlib.conf /usr/local/mpich-1.2.3/etc/mpicc-gcc.conf
```

In this example, the default value of `sysconffdir`, `$prefix/etc`, is used.

For example, if `MPICH` was built with `cc` as the compiler but a user wanted to use `gcc` instead, either the commands

```
MPICH_CC=gcc
mpicc ...
```

or

```
mpicc -config=gcc ...
```

would cause `mpicc` to load `'mpich-gcc.conf'` and use the appropriate definitions.

4.4 Fortran Compilers

`MPICH` provides support for both Fortran 77 and Fortran 90. Because `MPICH` is implemented in C, using `MPICH` from Fortran can sometimes require special options. This section discusses some of the issues. Note that `configure` tries to determine the options needed to support Fortran. You need the information in this section only if you have problems. Section 4.4.6 discusses how to support multiple Fortran compilers (e.g., `g77` and `pgf77`) with a single `MPICH` installation.

4.4.1 What if there is no Fortran compiler?

The `configure` program should discover that there is no Fortran compiler. You can force `configure` to not build the Fortran parts of the code with the option `--disable-f77`. In this case, only the C programs will be built and tested.

4.4.2 Fortran 90

During configuration, a number of Fortran 90-specific arguments can be specified. See the output of `configure -help`. In particular, when using the NAG Fortran 90 compiler, you should specify `-f90nag`.

4.4.3 Fortran 77 and Fortran 90

Selecting Fortran 90 with Fortran 77 should be done only when the two compilers are compatible, supporting the same datatypes and calling conventions. In particular, if the Fortran 90 compiler supports an 8-byte integer type, the Fortran 77 compiler must support `integer*8` (this is needed by the MPI-IO routines for the value of `MPI_OFFSET_KIND`). In addition, both compilers must support the same functions for accessing the command line, and the code for those commands must reside in the same library. If the two Fortran compilers are not compatible, you should either select the Fortran 90 compiler as both the Fortran 77 and Fortran 90 compiler (relying on the upward compatibility of Fortran), or build two separate configurations of `MPICH`. For example,

```
setenv FC f90
setenv F90 f90
configure
```

will use `f90` for both Fortran 77 and Fortran 90 programs. In many systems, this will work well. If there are reasons to have separate Fortran 90 and Fortran 77 builds, then execute the following commands (where `MPICH` is to be installed into the directory `'/usr/local'`):

```
setenv FC f77
configure --disable-f90 -prefix=/usr/local/mpich-1.2.3/f77-nof90
make
make install

setenv FC f90
setenv F90 f90
configure -prefix=/usr/local/mpich-1.2.3/f90
make
make install
```

This sequence of commands will build and install two versions of `MPICH`. An alternative approach that installs only a single version of `MPICH` is described in Section 4.4.6.

4.4.4 Fortran 90 Modules

If `configure` finds a Fortran 90 compiler, by default MPICH will try to create a Fortran 90 module for MPI. In fact, it will create two versions of an `mpi` module: one that includes only the MPI routines that do not take “choice” arguments and one that does include choice argument. A choice argument is an argument that can take any datatype; typically, these are the buffers in MPI communication routines such as `MPI_Send` and `MPI_Recv`.

The two different modules can be accessed with the `-nochoice` and `-choice` option to `mpif90`. The choice version of the module supports a limited set of datatypes (numeric scalars and numeric one- and two-dimensional arrays). This is an experimental feature; please send mail to `mpi-bugs@mcs.anl.gov` if you have any trouble. Neither of these modules offer full “extended Fortran support” as defined in the MPI-2 standard.

The reason for having two versions of the MPI module is that it is very difficult to provide a completely correct module that includes all of the functions with choice arguments. As it is, on many systems, the size of the Fortran 90 module to handle the routines with choice arguments will be larger than the entire C version of the MPI library. If you are uninterested in the Fortran 90 MPI module, or you wish to keep the installed version of MPICH small, you can turn off the creation of the Fortran 90 MPI module with the configure option `--disable-f90modules`.

4.4.5 Configuring with the Absoft Fortran Compiler

The Absoft compiler can be told to generate external symbols that are uppercase, lowercase, and lowercase with a trailing underscore (the most common case for other Unix Fortran compilers), or use mixed case (an extension of Fortran, which is only monospace). Each of these choices requires a *separate* MPICH configure and build step. MPICH has been tested in the mode where monospace names are generated; this case is supported because only this case supports common (and necessary for MPICH) extensions such as `getarg` and `iargc`. By default, MPICH forces the Absoft compiler to use lowercase; this matches most Unix Fortran compilers. MPICH will find the appropriate versions of `getarg` and `iargc` for this case. Because the examples and the test suite assume that the Fortran compiler is case-insensitive; the Fortran library produced by MPICH will only work with source code that uses monospace (either upper or lower) for all MPI calls.

In addition, you may need to use `-N90` if you use `character` data, because the MPICH Fortran interface expects the calling convention used by virtually all Unix Fortran systems (Cray UNICOS is handled separately). If you are building shared libraries, you will also need to set the environment variable `FC_SHARED_OPT` to `none`.

Early versions of the Absoft compiler could not handle multiple `-I` options. If you have trouble with this, you should get an update from Absoft.

4.4.6 Configuring for Multiple Fortran Compilers

In some environments, there are several different Fortran compilers, all of which define Fortran datatypes of the same size, and which can be used with the same C libraries. These

compilers may make different choices for Fortran name mappings (e.g., the external format of the names given to the linker) and use different approaches to access the command line. This section describes how to configure MPICH to support multiple Fortran compilers. However, if any of these steps fails, the best approach is to build a separate MPICH installation for each Fortran compiler.

The first step is to configure MPICH with the `--with-flibname` option. For example, if one of the compilers is `g77`, use

```
setenv FC g77
./configure --with-flibname=mpich-g77 ... other options ...
```

After you build, test, *and install* this version of MPICH, you can configure support for additional Fortran compilers as follows:

1. Change directory to `'src/fortran'`
2. Execute

```
setenv FC pgf77
./configure --with-mpichconfig --with-flibname=mpich-pgf77
make
make install-alt
```

To use a particular Fortran compiler, either select it on the `mpif77` command line with the `-config=name` option (e.g., `-config=pgf77`) or by selecting a particular `mpif77` command (e.g., `mpif77-pgf77`).

4.5 C++

The C++ support in MPICH has been provided by Indiana University (formerly the University of Notre Dame), and uses its own configure process (it also supports other MPI implementations). This version supports only the MPI-1 functions, and does not include support for the MPI-2 functions such as I/O or the functions for manipulating `MPI_Info`. Questions, comments, suggestions, and requests for additional information should be sent to `mpi2cpp-devel@osl.iu.edu`. Bug reports should also be sent to `mpi-bugs@mcs.anl.gov`. More information is available at the web site <http://www.osl.iu.edu/research/mpi2c++/>.

4.6 Using Shared Libraries

Shared libraries can help reduce the size of an executable. This is particularly valuable on clusters of workstations, where the executable must normally be copied over a network to each machine that is to execute the parallel program. However, there are some practical problems in using shared libraries; this section discusses some of them and how to solve most of those problems. Currently, shared libraries are not supported from C++.

In order to build shared libraries for MPICH, you must have configured and built MPICH with the `--enable-sharedlib` option. Because each Unix system and in fact each compiler uses a different and often incompatible set of options for creating shared objects and libraries, MPICH may not be able to determine the correct options. Currently, MPICH understands Solaris, GNU `gcc` (on most platforms, including Linux and Solaris), and IRIX. Information on building shared libraries on other platforms should be sent to `mpi-bugs@mcs.anl.gov`.

Once the shared libraries are built, you must tell the MPICH compilation and linking commands to use shared libraries (the reason that shared libraries are not the default will become clear below). You can do this either with the command line option `-shlib` or by setting the environment variable `MPICH_USE_SHLIB` to `yes`. For example,

```
mpicc -o cpi -shlib cpi.c
```

or

```
setenv MPICH_USE_SHLIB yes
mpicc -o cpi cpi.c
```

Using the environment variable `MPICH_USE_SHLIB` allows you to control whether shared libraries are used without changing the compilation commands; this can be very useful for projects that use makefiles.

Running a program built with shared libraries can be tricky. If you have trouble, particular if programs will either not start or start and issue error messages about missing libraries, see Section 6.4.

4.7 File System Issues

Most users do not need to worry about file systems. However, there are two issues: using NFS (the Network File System) with MPI-IO and using NFS with some automounters. These issues are covered in the next two sections.

4.7.1 NFS and MPI-IO

To use MPI-IO multihost on NFS file systems, NFS should be version 3, and the shared NFS directory must be mounted with the “no attribute caching” (`noac`) option set (the directory cannot be automounted). If NFS is not mounted in this manner, the following error could occur:

```
MPI_Barrier: Internal MPI error: No such file or directory
File locking messages
```

In order to reconfigure NFS to handle MPI-IO properly, the following sequence of steps are needed (root permission required):

1. confirm you are running NFS version 3

```
rpcinfo -p 'hostname' | grep nfs
```

for example, there should be a '3' in the second column

```
fire >rpcinfo -p fire | grep nfs
100003      3      udp      2049      nfs
```

2. edit '/etc/fstab' for each NFS directory read/written by MPI-IO on each machine used for multihost MPI-IO. The following is an example of a correct fstab entry for /epm1:

```
root >grep epm1 /etc/fstab
gershwin:/epm1 /rmt/gershwin/epm1 nfs bg,intr,noac 0 0
```

If the “noac” option is not present, add it and then remount this directory on each of the machines that will be used to share MPI-IO files.

```
root >umount /rmt/gershwin/epm1
root >mount /rmt/gershwin/epm1
```

3. confirm that the directory is mounted noac

```
root >grep gershwin /etc/mnttab
gershwin:/epm1 /rmt/gershwin/epm1 nfs
noac,acregmin=0,acregmax=0,acdirmin=0,acdirmax=0 0 0 899911504
```

Turning off of attribute caching may reduce performance of MPI-IO applications as well as other applications using this directory. The load on the machine where the NFS directory is hosted will increase.

4.8 Building MPICH

Once `configure` has determined the features of your system, all you have to do now is

```
make
```

This will clean all the directories of previous object files (if any), compile both profiling and non-profiling versions of the source code, including Romio and the C++ interface, build all necessary libraries, and link both a sample Fortran program and a sample C program as a test that everything is working. If anything goes wrong, check Section 6 to see if there is anything said there about your problem. If not, follow the directions in Section 6.2 for submitting a bug report. To simplify checking for problems, it is a good idea to use

```
make |& tee make.log
```

Specific (non-default) targets can also be made. See the ‘**Makefile**’ to see what they are.

After running this **make**, the size of the distribution will be about 45 Megabytes (depending on the particular machine it is being compiled for and the selected options), before building any of the examples or the extensive test library. The ‘**Makefile**’s are built for the various example subdirectories, but the example programs themselves have to be made “by hand”.

4.9 Installing MPICH for Others to Use

This step is optional. However, if you are installing MPICH, you should make sure that you specified the directory into which MPICH is to be installed when you configure MPICH by using the **-prefix** option. For example, if you plan to install MPICH into ‘**/usr/local/mpich-1.2.3/**’, then you should configure with the option **-prefix=/usr/local/mpich-1.2.3/**. If there is any possibility at all that you will build MPICH for several systems and/or devices, you should include that information in the prefix. For example, by using

```
-prefix=/usr/local/mpich-1.2.3/solaris/ch_p4,
```

you can later add

```
-prefix=/usr/local/mpich-1.2.3/solaris/ch_p4smp'
```

for a version that is built with the configure option **-comm=shared** (suitable for clusters of symmetric multiprocessors, hence the “smp” in the directory name). Once you have tested all parts of the MPI distribution (including the tools, particularly **upshot** and/or **nupshot**), you may install MPICH into a publically available directory, and disseminate information for other users, so that everyone can use the shared installation. To install the libraries and include files in a publicly available place, change to the top-level MPICH directory, and do

```
make install
```

The **man** pages will have been copied with the installation, so you might want to add ‘**/usr/local/mpich-1.2.3/man**’ to the default system **MANPATH**. The **man** pages can be conveniently browsed with the **mpiman** command, found in the **mpich/bin** directory.

It is possible to specify the directory into which MPICH should be installed after building MPICH by setting the value of **PREFIX** when executing the installation step:

```
make install PREFIX=/usr/local/mpich-1.2.3
```

However, some features, particularly the ability of Totalview to show MPICH message queues, will work only if MPICH is configured with the **prefix** set to the installation directory.

A good way to handle multiple releases of MPICH is to install them into directories whose names include the version number and then set a link from **mpi** to that directory. For example, if the current version is 1.2.3, the installation commands to install into ‘**/usr/local**’ are

```
make install PREFIX=/usr/local/mpi-1.2.3
rm /usr/local/mpi
ln -s /usr/local/mpi-1.2.3 /usr/local/mpi
```

The script ‘bin/mpiinstall’ provides more control over the installation of MPICH (in fact, `make install` just runs this script). For example, you can change the protection of the files when they are installed with the options `-mode=nnnn` (for regular files) and `-xmode=nnnn` (for executables and directories). You can set the directory into which the man pages will be placed with `-manpath=<path>`. The option `-help` shows the full set of options for `mpiinstall`.

Installing `nupshot` can sometimes be troublesome. You can use the switch `-nonupshot` to `mpiinstall` to not install `nupshot`; alternately, you can use the switch `-cpnupshot` to install the copy in ‘mpich/profiling/nupshot’. Normally, `mpiinstall` builds a new version of `nupshot` to insure that all of the paths are correct (`nupshot` needs to find files where it is installed). If you need to “manually” build `nupshot` for installation, the `-cpnupshot` switch will allow you to install that version.

You can test the installation by using the `configure` in ‘mpich/examples/test’. For example, if you have installed MPICH into ‘/usr/local/mpich-1.2.3/’ for architecture `solaris` and device `ch_p4`, execute

```
cd examples/test
./configure -mpichpath=/usr/local/mpich-1.2.3/solaris/ch_p4/bin
make testing
```

The test codes in the ‘mpich/examples/test’ directory may be used with *any* implementation of MPI, not just the MPICH implementation.

4.9.1 User commands

The commands `mpirun`, `mpicc`, `mpif77`, `mpiCC`, `mpif90`, and `mpiman` should be in the user’s search path. Note that if several architectures and/or MPICH devices are supported, it is important that the correct directory be added to the user’s path. These are installed into the ‘bin’ directory. If multiple architectures or devices are being used, be sure that the installation path distinguishes these. For example, if both the `ch_p4` and `ch_shmem` devices are built on a Solaris system, set the installation prefix to include these names:

```
-prefix='/usr/local/mpich-1.2.3/solaris/ch_p4', and
-prefix='/usr/local/mpich-1.2.3/solaris/ch_shmem'
```

respectively. Alternately, set the prefix to ‘/usr/local/mpich-1.2.3/’ and set the `execprefix` to

```
‘/usr/local/mpich-1.2.3/solaris/ch_p4’ and
‘/usr/local/mpich-1.2.3/solaris/ch_shmem’
```

respectively.

4.9.2 Installing documentation

The MPICH implementation comes with several kinds of documentation. Installers are encouraged to provide site-specific information, such as the location of the installation (particularly if it is not in `/usr/local/mpich-1.2.3/`).

4.9.3 Man pages

A complete set of Unix man pages for the MPICH implementation are in `mpich/man`. `man/man1` contains the commands for compiling, linking, and running MPI programs; `man/man3` contains the MPI routines; `man/man4` contains the MPE routines, and `man/man5` contains the MPID routines (these are for the low-level part of the MPICH implementation, are not of interest to users). The command `mpich/bin/mpiman` is a script that can present the manual pages for MPICH in various forms, using either the terminal-style `man`, `xman`, or one of several HTML browsers.

4.9.4 Examples

Users often prefer working from example `Makefile`'s and programs. The directory that is installed in the `examples` directory contains a C and Fortran version of the `pi` program, along with a `Makefile.in`. Other examples there include a simple parallel I/O program and an MPI program written using the C++ bindings for the MPI functions. Users may be interested in some of the examples that are in the source tree, also in the `examples` directory.

4.10 Computational Grids: the globus2 device

The `globus2` device⁴ supports the execution of MPI programs on “computational grids” that may include parallel computers and workstations, and that may span multiple sites. In such grid environments, different sites may support different security mechanisms and different process creation mechanisms. The `globus2` device hides these low-level details from you, allowing you to start programs with `mpirun` as on MPPs and workstation clusters. The `globus2` device also provides other convenient features, such as remote access to files and executable staging. These features are provided by using services supplied by the Globus toolkit: see <http://www.globus.org> for details.

The `globus2` device requires that special servers be running on the computers where processes are to be created. In our discussion of how to use the `globus2` device, we assume that we are using the `globus2` device on a collection of machines on which various Globus servers are installed and running. Such a collection is often referred to as a *computational grid*, for example, NASA's Information Power Grid (IPG). If possible, we recommend that you use the `globus2` device in this environment. If you wish to use the `globus2` device in other situations, please send email to developers@globus.org. Details of how to run MPI programs using the `globus2` device on Globus-enabled computational grids are in Appendix 3.3.

⁴`globus2` replaces the `globus` device distributed in previous releases of MPICH.

4.11 Thorough Testing

The `examples/test` directory contains subdirectories of small programs that systematically test a large subset of the MPI functions. The command

```
make testing
```

in the `MPICH` directory will cause these programs to be compiled, linked, executed, and their output to be compared with the expected output. Linking all these test programs takes up considerable space, so you might want to do

```
make clean
```

in the test directory afterwards. The individual parts of MPI (point-to-point, collective, topology, etc.) can be tested separately by

```
make testing
```

in the separate subdirectories for `examples/test`.

If you have a problem, first check the troubleshooting guides and the lists of known problems. If you still need help, send detailed information to `mpi-bugs@mcs.anl.gov`.

4.12 Internationalization

`MPICH` has support for providing error messages in different languages. This makes use of the X/Open message catalog services, which are a standard way of providing multi-language support. This multi-language support is often called NLS, for National Language Support. `MPICH` comes with error messages in US English; additional languages will be provided as we get the translations (if you wish to provide one, please send mail to `mpi-bugs@mcs.anl.gov`). More precisely, `MPICH` uses an English version that uses the ISO Latin-1 character set (ISO8859-1). We expect to provide other versions that also use the Latin-1 character set, subject to getting translations of the messages.

To create a new message catalog, copy the file `mpich.En_US.msg` to `mpich.mylanguage.msg` and translate the entries. The value of `mylanguage` should match the ones used for your system; for example, `mpich.De_DE.msg` for German. Many systems put their NLS files in `/usr/lib/nls/msg`; you can also check the value of the environment variable `NLSPATH` on your system. Note that some systems provide the routines and programs to support NLS, but do not make use of it and do not provide a initial `NLSPATH` value.

For emacs users, check the Emacs info under “European Display”. The commands

```
M-x standard-display-european
```

```
M-x iso-accents-mode
```

can be used to input most European languages. You can also load `iso-transl` and use `C-x 8` to compose characters (this sets the high bit in the character). `MPICH` currently does not

support languages that require multi-byte character sets (such as Japanese). However, the only changes needed are in the file `'src/env/errmsg.c'`; if you are interested in developing a multi-byte character set version, please let us know.

By default, MPICH uses the value of `'NLSPATH'` to find the message catalogs. If this fails, it tries `'MPICHNLSPATH'`, and if that fails, it uses English language versions that are coded into the library.

The catalogs are not, however, installed into these directories. Instead, you will find them in the library directory for a particular architecture; for example, `'mpich/rs6000/lib'`.

5 Documentation

This distribution of MPICH comes with complete `man` pages for the MPI routines, commands for compiling and linking MPI programs, and the MPE extensions. The command `mpiman` in `'mpich/bin'` is a good interface to the `man` pages.⁵ The `'mpich/www'` directory contains HTML versions of the `man` pages for MPI and MPE. The `'mpich/doc'` directory contains this *Installation and User's Guide*. All documentation is also available on the web at www.mcs.anl.gov/mpi/mpich/docs.html.

Information about MPI is available from a variety of sources. Some of these, particularly WWW pages, include pointers to other resources.

- The Standard itself:
 - As a Technical report [4].
 - As Postscript and HTML at www.mpi-forum.org, for both MPI-1 and MPI-2.
 - As a journal article in the Fall 1994 issue of the Journal of Supercomputing Applications [14] for MPI-1 and as a journal article in the International Journal of High Performance Computing Applications in 1998.
- MPI Forum discussions
 - The MPI Forum email discussions and both current and earlier versions of the Standard are available from www.netlib.org. MPI-2 discussions are available at www.mpi-forum.org.
- Books:
 - *Using MPI: Portable Parallel Programming with the Message-Passing Interface, Second Edition*, by Gropp, Lusk, and Skjellum [10].
 - *Using MPI-2: Advanced Features of the Message-Passing Interface*, by Gropp, Lusk, and Thakur [11].
 - *MPI—The Complete Reference: Volume 1, The MPI Core*, by Snir, et al. [16].
 - *MPI—The Complete Reference: Volume 2, The MPI-2 Extensions*, by Gropp, et al. [5].

⁵The `mpiman` command is created by the configure process.

- *Parallel Programming with MPI*, by Peter S. Pacheco [15].
- Newsgroup:
 - `comp.parallel.mpi`
- Mailing lists:
 - `mpi-comments@mpi-forum.org`: The MPI Forum discussion list.
 - `mpi-impl@mcs.anl.gov`: The implementors' discussion list.
 - `mpi-bugs@mcs.anl.gov` is the address to which you should report problems with `mpich`.
- Implementations available from the web:
 - MPICH is available from `http://www.mcs.anl.gov/mpi/mpich` or by anonymous ftp from `ftp.mcs.anl.gov` in the directory 'pub/mpi/mpich', file 'mpich.tar.gz'.
 - Links to other implementations are available at `www.mcs.anl.gov/mpi/implementations.html`.
- Test code repository:
 - `ftp://ftp.mcs.anl.gov/pub/mpi/mpi-test`

6 In Case of Trouble

This section describes some of the problems that you may run into and some solutions, along with information on submitting bug reports.

6.1 Things to try first

If something goes wrong, the first thing to do is to check the output of `configure` or `make` for some obvious problem, such as an improperly specified compiler or inadequate disk space. Next, check the web page `www.mcs.anl.gov/mpi/mpich/buglist-tbl.html` for any recent patches that may fix your problem. After that, check Section 6.3 for common problems. If you still can't find a solution to your problem, submit a bug report and we will try to help you.

6.2 Submitting bug reports

Send any problem that you can not solve by checking this section to `mpi-bugs@mcs.anl.gov`.

Please include:

- The version of MPICH (e.g., 1.2.3)

- The output of running your program with the `-mpiversion` argument (e.g., `mpirun -np 1 a.out -mpiversion`)
- The output of

```
uname -a
```

for your system. If you are on an SGI system, also

```
hinv
```

- The files `'config.log'` and `'src/fortran/config.log'`.
- If the problem is with a script such as `configure` or `mpirun`, run the script with the `-echo` argument (e.g., `mpirun -echo -np 4 a.out`).

If you have more than one problem, please send them in separate messages; this simplifies our handling of problem reports.

The rest of this section contains some information on trouble-shooting MPICH. Some of these describe problems that are peculiar to some environments and give suggested work-arounds. Each section is organized in question and answer format, with questions that relate to more than one environment (workstation, operating system, etc.) first, followed by questions that are specific to a particular environment. Problems with workstation clusters are collected together as well. To make it easier to find solutions, the most common problems are described first.

6.3 The Most Common Problems

This section describes some of the most common problems encountered when building and using MPICH. See also Section A which covers frequently asked questions, including some additional problems.

Missing symbols when linking. The most common source of missing symbols is a failure of the MPICH configure step to determine how to pass command line arguments to Fortran. Check the output of the configure step for any error messages or warnings about building the Fortran libraries. If you do not require Fortran, reconfigure MPICH using the configure option `--disable-f77` and remake MPICH. If you need Fortran and cannot figure out how to make MPICH work with Fortran, send a bug report to `mpi-bugs@mcs.anl.gov`.

Another common problem with programs that mix Fortran and C is missing libraries. The MPICH configure attempts to determine the libraries that are necessary when linking C with Fortran, but may miss some. There are additional suggestions for this problem in Section 6.9.

SIGSEGV. Any message that mentions SIGSEGV is referring to a “segmentation violation” during program execution. This is usually due to an error in the user’s program, such as an array overwrite or use of an uninitialized variable in referencing storage.

6.4 Troubleshooting Shared Libraries

Shared libraries provide a useful and powerful tool for speeding linking and execution of programs. However, the implementation of shared libraries in most operating systems leaves much to be desired. This section covers some of the problems and workarounds for them.

Some (most?) systems *do not remember* where the shared library was found when the executable was linked!⁶ Instead, they depend on finding the shared library in either a default location (such as `/lib`) or in a directory specified by an environment variable such as `LD_LIBRARY_PATH` or by a command line argument such as `-R` or `-rpath` (more on this below). The MPICH configure tests for this and will report whether an executable built with shared libraries remembers the location of the libraries. It also attempts to use a compiler command line argument to force the executable to remember the location of the shared library.

If you need to set an environment variable to indicate where the MPICH shared libraries are, you need to ensure that both the process that you run `mpirun` from and any processes that `mpirun` starts gets the environment variable. The easiest way to do this is to set the environment variable within your `.cshrc` (for `csh` or `tcsh` users) or `.profile` (for `sh` and `ksh` users) file.

However, setting the environment variable within your startup scripts can cause problems if you use several different systems. For example, you may have a single `.cshrc` file that you use with both an SGI (IRIX) and Solaris system. You do not want to set the `LD_LIBRARY_PATH` to point the SGI at the Solaris version of the MPICH shared libraries⁷. Instead, you would like to set the environment variable before running `mpirun`:

```
setenv LD_LIBRARY_PATH ${LD_LIBRARY_PATH}:/usr/local/mpich/lib/shared
mpirun -np 4 cpi
```

Unfortunately, this won't always work. Depending on the method that `mpirun` and MPICH use to start the processes, the environment variable may not be sent to the new process. This will cause the program to fail with a message like

```
ld.so.1: /home/me/cpi: fatal: libmpich.so.1.0: open failed: No such
file or directory
Killed
```

Some devices support starting new processes with the current environment; check the documentation for `mpirun` (Section 3.2).

An alternative to using `LD_LIBRARY_PATH` and the secure server is to add an option to the link command that provides the path to use in searching for shared libraries. Unfortunately,

⁶There is a reason for this unfortunate choice that has to do with distributing executables linked with shared libraries. However, the capability to save the location should be provided, at least as an option, by the linker.

⁷You can make `.cshrc` check for the kind of system that you are running on and pick the paths appropriately. This isn't as flexible as the approach of setting the environment variable from the running shell.

the option that you would like is “append this directory to the search path” (such as you get with `-L`). Instead, many compilers provide only “replace the search path with this path.”⁸ For example, some compilers allow `-Rpath:path:...:path` to specify a replacement path. Thus, if both `MPICH` and the user provide library search paths with `-R`, one of the search paths will be lost. Eventually, `mpicc` and friends can check for `-R` options and create a unified version, but they currently do not do this. You can, however, provide a complete search path yourself if your compiler supports an option such as `-R`.

The preceeding may sound like a lot of effort to go to, and in some ways it is. For large clusters, however, the effort will be worth it: programs will start faster and more reliably, because there is less network and file system traffic.

6.5 Other Problems

The following items describe miscellaneous problems that we have encountered. These may help solve less common problems.

6.6 Problems configuring

6.6.1 General

1. **Q:** Configure reports that floating point is not commutative! How do I fix it?
A: Check your compiler’s documentation. On RS/6000’s, the `-qnomaf` (no multiply-add floating point) option. On some other systems, intermediate results may be stored in 80-bit registers (Intel CPUs do this); this can also lead to inconsistent rounding. You may be able to force the compiler to round to 64 bits.

6.6.2 Linux

1. **Q:** The configure step issues the message:

```
checking that the compiler f77 runs... no
Fortran compiler returned non-zero return code
Output from test was
f2ctmp_conftest.f:
  MAIN main:
```

A: This is probably caused by a problem in the Fortran compiler in older versions of Linux. The `f77` command in Linux was often a shell script that uses the `f2c` program to convert the Fortran program to C, and then compile it with the C compiler. In many versions of Linux, this script has an error that causes a non-zero return code even when the compilation is successful.

To fix this problem, you need a corrected `f77` script. If you can edit the script yourself, change the last 3 lines from

⁸Even though the linker may provide the “append to search path” form.

```
case $cOPT in 2) $CC $G -o $OUTF $OFILES -lf2c -lm;; esac
rc=$?
exit $rc
```

to

```
case $cOPT in 2) $CC $G -o $OUTF $OFILES -lf2c -lm;; esac
rc=$?
trap 0
exit $rc
```

2. **Q:** The link test fails on Linux with messages like

```
overtake.o(.text+0x59): undefined reference to 'MPI_COMM_WORLD'
overtake.o(.text+0x81): undefined reference to 'MPI_COMM_WORLD'
...
```

A: This is probably caused by a problem in the Fortran compiler in Linux. In some early versions of Linux, the `f77` command in Linux is often a shell script that uses the `f2c` program to convert the Fortran program to C, and then compile it with the C compiler. In many versions of Linux, this script has an error that causes a non-zero return code even when the compilation is successful.

To fix this problem, you need a corrected `f77` script. If you can edit the script yourself, change the last 3 lines from

```
case $cOPT in 2) $CC $G -o $OUTF $OFILES -lf2c -lm;; esac
rc=$?
exit $rc
```

to

```
case $cOPT in 2) $CC $G -o $OUTF $OFILES -lf2c -lm;; esac
rc=$?
trap 0
exit $rc
```

3. **Q:** During the configure step, messages like

```
/homes/me/mpich/configure: 134956160: Permission denied
```

sometimes appear. What is wrong?

A: This is a bug in the Linux (actually GNU) `sh` shell. The shell is attempting to create a file with the name `/tmp/t<processid>-sh` (e.g., `/tmp/t11203-sh`), but was unable to. This may happen if `/tmp` is full; however, it can also happen when the shell created the same file for another user days before. (You can see this happen by running `configure` under `strace -f`). The only fix is to have your systems administrator clean old files out of `/tmp`.

6.7 Problems building MPICH

6.7.1 General

1. **Q:** When running make on MPICH, I get this error:

```
ar: write error: No such file or directory
*** Error code 1
```

I've looked, and all the files are accessible and have the proper permissions.

A: Check the amount of space in `/tmp`. This error is sometimes generated when there is insufficient space in `/tmp` to copy the archive (this is a step that `ar` takes when updating a library). The command `df /tmp` will show you how much space is available. Try to insure that at least twice the size of the library is available.

2. **Q:** When running make on MPICH, I get errors when executing `ranlib`.

A: Many systems implement `ranlib` with the `ar` command, and they use the `/tmp` directory by default because it “seems” obvious that using `/tmp` would be faster (`/tmp` is often a local disk). Unfortunately, some systems have ridiculously small `/tmp` partitions, making any use of `/tmp` very risky. In some cases, the `ar` commands used by MPICH will succeed because they use the `l` option—this forces `ar` to use the local directory instead of `/tmp`. The `ranlib` command, on the other hand, may use `/tmp` and cannot be fixed.

In some cases, you will find that the `ranlib` command is unnecessary. In these cases, you can reconfigure with `-noranlib`. If you must use `ranlib`, either reduce the space used in `/tmp` or increase the size of the `/tmp` partition (your system administrator will need to do this). There should be at least 20–30 MBytes free in `/tmp`.

3. **Q:** When doing the link test, the link fails and does not seem to find any of the MPI routines:

```
/homes/me/mpich/IRIX32/ch_p4/bin/mpicc -o overtake overtake.o test.o
ld: WARNING 126: The archive /homes/me/mpich/IRIX32/ch_p4/lib/libmpi.a
defines no global symbols. Ignoring.
ld: WARNING 84: /usr/lib/libsun.a is not used for resolving any symbol.
ld: ERROR 33: Unresolved data symbol "MPI_COMM_WORLD" -- 1st referenced by overtake.o.
ld: ERROR 33: Unresolved text symbol "MPI_Send" -- 1st referenced by overtake.o.
...
```

A: Check that the `ar` and `ranlib` programs are compatible. One site installed the Gnu `ranlib` in such a way that it would be used with the vendors `ar` program, with which it was incompatible. Use the `-noranlib` option to `configure` if this is the case.

6.7.2 Workstation Networks

1. **Q:** When building MPICH, the make fails with errors like this:


```

making p4 in directory lib
make libp4.a
cc -Aa -g -I../include -I../..../include -c p4_globals.c
cc: "/usr/include/netinet/in.h", line 69: error 1000: Unexpected symbol: "u_long".
cc: "/usr/include/netinet/in.h", line 127: error 1000: Unexpected symbol: "u_short".

```

etc.

A: The problem here is that the system (not MPICH) include files are incompatible with your choice of compiler. Check to see if `cc` is aliased (in C shell, do `alias cc`). If it is, either unalias it or set the environment variable `CC` to the full path for the compiler. To get the full path, do

```

unalias cc
setenv CC `which cc`

```

and then reconfigure.

6.7.3 Cray T3D

1. **Q:** When linking I get

```

mppldr-133 cf77: CAUTION
Unsatisfied external references have been encountered.

```

```

Unsatisfied external references
Entry name      Modules referencing entry

GETARG (equivalenced to $USX1)
                MPIR_GETARG

```

A: You may have specified the Fortran compiler with the `F77` environment variable or the `-fc` argument to configure. The MPICH Fortran implementation of MPI uses a common Fortran extension, `GETARG`, to get the command line arguments. Most Fortran runtime systems support this, but Cray uses `call pxfgetarg(i,s,len(s),ierr)` instead. You can change the file `'src/env/farg.f'` manually to call the correct routine (but note that configure builds a new `'farg.f'` from `'farg.f.in'` each time that it is run).

MPICH now attempts to determine the correct names of the routines to access the command line. If you find that MPICH fails to determine the names correctly, please send a bug report to mpi-bugs@mcs.anl.gov.

6.7.4 SGI

1. **Q:** The build on an SGI Power Challenge fails with

```

Signal: SIGSEGV in Back End Driver phase.
> ### Error:
> ### Signal SIGSEGV in phase Back End Driver -- processing aborted

```

```
> f77 ERROR: /usr/lib64/cmplrs/be died due to signal 4
> f77 ERROR: core dumped
> *** Error code 2 (bu21)
> *** Error code 1 (bu21)
> *** Error code 1 (bu21)
```

A: Our information is that setting the environment variable `SGI_CC` to `-ansi` will fix this problem.

2. **Q:** The build on an SGI with architecture `IRIXN32` fails with

```
cc: Warning: -c should not be used with ucode -O3 -o32 \
on a single file; use -j instead to get inter-module optimization.
```

A: Amazingly, the standard `-c` option is *not valid* for the SGI compilers when both `-O3` and `-n32` are specified. This is a “feature” of the SGI compiler, and there is no way to work around this for `MPICH` (other than a massive and non-portable rewrite of all the ‘Makefile’s). Your only option is to not use the `-O3` option.

6.7.5 Linux

1. **Q:** The link test failed on Linux with

```
...
cc -o overtake overtake.o test.o -L/usr/local/mpich/LINUX/ch_p4/lib
-lmpi
overtake.o(.text+0x71): undefined reference to 'MPI_COMM_WORLD'
overtake.o(.text+0x82): undefined reference to 'MPIR_I_DOUBLE'
overtake.o(.text+0xe1): undefined reference to 'MPI_COMM_WORLD'
...
```

A: We have been informed that there is a error in the `f77` script in some versions of Linux which causes this problem. Try either getting a patch for the `f77` script or reconfiguring with `-nof77`.

2. **Q:** The build fails for the `ch_p4` device when using the Compaq C compiler.

A: There is an incompatibility with the system include files (not the `MPICH` include files). If you can modify ‘`/usr/include/rpc/xdr.h`’, add the following near the top of that file:

```
#if defined(__DECC) || defined(__DECCXX)
typedef long int int64_t;
#endif
```

6.7.6 Compaq ULTRIX and Tru64

1. **Q:** When trying to build, the `make` aborts early during the cleaning phase:

```
amon:MPICH/mpich>make clean
/bin/rm -f *.o *~ nupshot
*** Error code 1
```

A: This is a bug in the shell support on some Compaq ULTRIX systems. You may be able to work around this with

```
setenv PROG_ENV SYSTEM_FIVE
```

Configuring with `-make=s5make` may also work.

6.8 Problems in testing

The MPICH test suite, in `examples/test`, performs a fairly complete test of an MPI implementation. If there is an error, it usually indicates a problem with the implementation of MPI; if you encounter such a problem, please report it to `mpi-bugs@mcs.anl.gov`. However, there are a few exceptions that are described here.

6.8.1 General

1. **Q:** The test `pt2pt/structf` fails with

```
0 - Error in MPI_ADDRESS : Invalid argument: Address of location
given to MPI_ADDRESS does not fit in Fortran integer
[0] Aborting program!
```

A: This is not an error; it is a gap in the MPI-1 definition that is fixed in MPI-2 (with the routine `MPI_Get_address`. This indicates that Fortran integers are not large enough to hold an address. This does indicate that MPI programs written in Fortran should not use the `MPI_Address` function on this system.

2. **Q:** The test `env/timers` fails with

```
Timer around sleep(1) did not give 1 second; gave 0.399949
```

A: The low-level software that MPICH uses probably makes use of the `SIGALRM` signal, thus denying it to the user's program. This is not an error (the standard permits systems to make use of any signals) though it is unfortunate.

6.9 Problems compiling or linking Fortran programs

6.9.1 General

1. **Q:** When linking the test program, the following message is generated:

```
f77 -g -o secondf secondf.o -L/usr/local/mpich/lib/sun4/ch_p4 -lmpich
invalid option -L/usr/local/mpich/lib/sun4/ch_p4
ld: -lmpich: No such file or directory
```

A: This f77 program does not accept the `-L` command to set the library search path. Some systems provide a shell script for f77 that is very limited in its abilities. To work around this, use the full library path instead of the `-L` option:

```
f77 -g -o secondf secondf.o /usr/local/mpich/lib/sun4/ch_p4/libmpich.a
```

As of the MPICH 1.2.0 release, the MPICH configure attempts to find the correct option for indicating library paths to the Fortran compiler. If you find that the MPICH configure has made an error, please submit a bug report to mpi-bugs@mcs.anl.gov.

2. **Q:** When linking Fortran programs, I get undefined symbols such as

```
f77 -c secondf.f
secondf.f:
  MAIN main:
f77 -o secondf secondf.o -L/home/mpich/lib/solaris/ch_shmem -lmpich
Undefined                               first referenced
  symbol                                in file
getdomainname
/home/mpich/lib/solaris/ch_shmem/libmpi .a(shmempriv.o)
ld: fatal: Symbol referencing errors. No output written to secondf
```

There is no problem with C programs.

A: This means that your C compiler is providing libraries for you that your Fortran compiler is not providing. Find the option for the C compiler and for the Fortran compilers that indicate which library files are being used (alternately, you may find an option such as `-dryrun` that shows what commands are being used by the compiler). Build a simple C and Fortran program and compare the libraries used (usually on the `ld` command line). Try the ones that are present for the C compiler and missing for the Fortran compiler. MPICH attempts to determine these libraries for you. If you have trouble, send a bug report and include the contents of the file `'mpich/src/fortran/config.log'`.

3. **Q:** When trying to compile Fortran code with a Fortran 90 or Fortran 95 compiler, I get error messages like

```
Error: foo.f, line 30: Inconsistent datatype for argument 1 in MPI_SEND
```

A: The Fortran language requires that in two calls to the same subroutine, the types of arguments must be the same. That is, if you called `MPI_SEND` with a `REAL` buffer as the first argument and then called it with an `INTEGER` buffer as the first argument, a Fortran compiler can consider this an error. Few Fortran 77 compilers would complain about this; more Fortran 90 and Fortran 95 compilers check for this. There are two solutions. One is to use the MPI module (in the “choice” version: use the `-choicemod` option for `mpif90`); the other is to use an option to tell the Fortran 90 compiler to allow argument mismatches. For example, the argument `-mismatch` will cause the NAG Fortran compilers to allow mismatched arguments. Using the MPI module is the preferred approach.

Fortran 77 users may sometimes see a similar message, particularly with later versions of `g77`. The option `-Wno-globals` will suppress these warning messages.

6.10 Problems Linking C Programs

6.10.1 General

1. **Q:** When linking programs, I get messages about `__builtin_saveregs` being undefined.

A: You may have a system on which C and Fortran compilers are incompatible (for example, using `gcc` and the vendor’s Fortran compiler). If you do not plan to use Fortran, the easiest fix is to rebuild with the `-nof77` option to configure.

You should also look into making your C compiler compatible with your Fortran compiler. Because this is very dependent on the specific system and compilers that you are using, you may need to find a local expert who can help you.

The easiest but ugliest possibility is use `f2c` to convert Fortran to C, then use the C compiler to compile everything. If you take this route, remember that *every* Fortran routine has to be compiled using `f2c` and the C compiler.

Alternatively, you can use various options (check the man pages for your compilers) to see what libraries that add when they link. Add those libraries to the link line for the *other* compiler. If you find a workable set of libraries, edit the appropriate scripts (e.g., `mpicc`) to include the necessary libraries. `MPICH` attempts to find all the libraries that you need but is not always successful.

6.10.2 HPUX

1. **Q:** When linking on HPUX, I get an error like this:

```
cc -o pgm pgm.o -L/usr/local/mpich/lib/hpux/ch_p4 -lmpich -lm
/bin/ld: Unsatisfied symbols:
sigrelse (code)
sigset (code)
sighold (code)
*** Error code 1
```

A: You need to add the link option `-lv3`. The `ch_p4` device uses the System V signals on the HP; these are provided in the 'V3' library.

6.10.3 LINUX

1. **Q:** When linking a Fortran program, I get

Linking:

```
foo.o(.data+0x0): undefined reference to 'pmpi_wtime_'
```

A: This is a bug in the `pgf77` compiler (which is itself a workaround for a bug in the LINUX `ld` command). You can fix it by either adding `-lmpich` to the link line or modifying the '`mpif.h`' to remove the `external pmpi_wtime, pmpi_wtick` statement.

The MPICH configure attempts to determine if `pmpi_wtime` and `pmpi_wtick` can be declared in '`mpif.h`' and removes them if there is a problem. If this happens and you use `pmpi_wtime` or `pmpi_wtick` in your program, you will need to declare them as functions returning double precision values.

6.11 Problems starting programs

6.11.1 General

1. **Q:** When trying to start a program with

```
mpirun -np 2 cpi
```

either I get an error message or the program hangs.

A: On some systems such as IBM SPs, there are many mutually exclusive ways to run parallel programs; each site can pick the approach(es) that it allows. The script `mpirun` tries one of the more common methods, but may make the wrong choice. Use the `-v` or `-t` option to `mpirun` to see how it is trying to run the program, and then compare this with the site-specific instructions for using your system. You may need to adapt the code in `mpirun` to meet your needs. See also the next question.

2. **Q:** When trying to run a program with, e.g., `mpirun -np 4 cpi`, I get

```
usage : mpirun [options] <executable> [<dstnodes>] [-- <args>]
```

or

```
mpirun [options] <schema>
```

A: You have a command named `mpirun` for a different implementation of MPI in your path ahead of the MPICH version. Execute the command

```
which mpirun
```

to see which command named `mpirun` was actually found. The fix is to either change the order of directories in your path to put the MPICH version of `mpirun` first, or to define an alias for `mpirun` that uses an absolute path. For example, in the `cs` shell, you might do

```
alias mpirun /usr/local/mpich/bin/mpirun
```

to set `mpirun` to the MPICH version.

3. **Q:** When I try to run a program on more than one processor, I get an error message

```
mpirun -np 2 cpi
/home/me/cpi: error in loading shared libraries: libcxa.so.1: cannot open
shared object file: No such file or directory
```

There is no trouble running on one processor.

A: This means that some shared library used by the system cannot be found on remote processors. There are two possibilities:

- (a) The shared libraries are not installed on the remote processor. To fix this, have your system administrators install the libraries.
- (b) The shared libraries are not in the default path. This can happen if the path is in an environment variable that is set in your current shell but that is not part of your default or remote shell environment. The fix in this case is harder, because you must communicate the location of the shared library to the executable (a major deficiency in most Unix shared-library designs is that executables do not, by default, remember where a shared library was found when linking). The simplest fix may be to have your system administrators place the necessary shared libraries into one of the directories that is searched by default. If this is not possible, then you will need to help the compiler and linker out.

Many linkers provide a way to specify the search path for shared libraries. The trick is to (a) pass this command to the linker program and (b) specify all of the libraries that are needed.

For example, on Linux systems, the linker command to specify the shared library search path is `-rpath path`, e.g., `-rpath /usr/lib:/usr/local/lib`. To pass this command to the linker through the Intel C compiler `icc`, the command `-Qoption,link,-rpath,path` is used. By default, the Linux linker looks in `/usr/lib` and the directories specified by the environment variable `LD_LIBRARY_PATH`. Thus, to force the linker to include the path to the shared library, you can use

```
mpicc -o cpi cpi -Qoption,link,-rpath,$LD_LIBRARY_PATH:/usr/lib
```

If this works, then consider editing the value of `LDFLAGS` in the compiler scripts (e.g., `mpicc`) to include this option.

Unfortunately, each compiler has a different way of passing these arguments to the linker, and each linker has a different set of arguments for specifying the shared library search path. You will need to check the documentation for your system to find this options.

4. **Q:** When attempting to run `cpilog` I get the following message:

```
ld.so.1: cpilog: fatal: libX11.so.4: can't open file: errno 2
```

A: The X11 version that `configure` found isn't properly installed. This is a common problem with Sun/Solaris systems. One possibility is that your Solaris machines are running slightly different versions. You can try forcing static linking (`-Bstatic` on Solaris).

Alternately, consider adding these lines to your `‘.login’` (assuming C shell):

```
setenv OPENWINHOME /usr/openwin
setenv LD_LIBRARY_PATH /opt/SUNWspro/lib:/usr/openwin/lib
```

(you may want to check with your system administrator first to make sure that the paths are correct for your system). Make sure that you add them *before* any line like

```
if ($?USER == 0 || $?prompt == 0) exit
```

5. **Q:** My program fails when it tries to write to a file.

A: If you opened the file *before* calling `MPI_INIT`, the behavior of MPI (not just the MPICH implementation of MPI) is undefined. In the `ch_p4` device, only process zero (in `MPI_COMM_WORLD`) will have the file open; the other processes will not have opened the file. Move the operations that open files and interact with the outside world to after `MPI_INIT` (and before `MPI_FINALIZE`).

6. **Q:** Programs seem to take forever to start.

A: This can be caused by any of several problems. On systems with dynamically-linked executables, this can be caused by problems with the file system suddenly getting requests from many processors for the dynamically-linked parts of the executable (this has been measured as a problem with some DFS implementations). You can try statically linking your application.

6.11.2 IBM RS6000

1. **Q:** When trying to run on an IBM RS6000 with the `ch_p4` device, I got

```
% mpirun -np 2 cpi
Could not load program /home/me/mpich/examples/basic/cpi
Could not load library libC.a[shr.o]
Error was: No such file or directory
```

A: This means that MPICH was built with the `x1C` compiler but that some of the machines in your `‘util/machines/machines.rs6000’` file do not have `x1C` installed. Either install `x1C` or rebuild MPICH to use another compiler (either `x1c` or `gcc`; `gcc` has the advantage of never having any licensing restrictions).

6.11.3 IBM SP

1. **Q:** When starting my program on an IBM SP, I get this:

```
$ mpirun -np 2 hello
ERROR: 0031-124  Couldn't allocate nodes for parallel execution.  Exiting ...
ERROR: 0031-603  Resource Manager allocation for task: 0, node:
me1.myuniv
.edu, rc = JM_PARTIONCREATIONFAILURE
ERROR: 0031-635  Non-zero status -1 returned from pm_mgr_init
```

A: This means that either `mpirun` is trying to start jobs on your SP in a way different than your installation supports or that there has been a failure in the IBM software that manages the parallel jobs (all of these error messages are from the IBM `poe` command that `mpirun` uses to start the MPI job). Contact your system administrator for help in fixing this situation. Your system administrator can use

```
dsh -av "ps aux | egrep -i 'poe|pmd|jmd'"
```

from the control workstation to search for stray IBM POE jobs that can cause this behavior. The files `/tmp/jmd_err` on the individual nodes may also contain useful diagnostic information.

2. **Q:** When trying to run on an IBM SP, I get the message from `mpirun`:

```
ERROR: 0031-214  pmd: chdir </a/user/gamma/home/mpich/examples/basic>
ERROR: 0031-214  pmd: chdir </a/user/gamma/home/mpich/examples/basic>
```

A: These are messages from the IBM system, not from `mpirun`. They may be caused by an incompatibility between POE, the automounter (especially the AMD automounter) and the shell, especially if you are using a shell other than `ksh`. There is no good solution; IBM often recommends changing your shell to `ksh`!

3. **Q:** When trying to run on an IBM SP, I get this message:

```
ERROR: 0031-124  Less than 2 nodes available from pool 0
```

A: This means that the IBM POE/MPL system could not allocate the requested nodes when you tried to run your program; most likely, someone else was using the system. You can try to use the environment variables `MP_RETRY` and `MP_RETRYCOUNT` to cause the job to wait until the nodes become available. Use `man poe` to get more information.

4. **Q:** When running on an IBM SP, my job generates the message

```
Message number 0031-254 not found in Message Catalog.
```

and then dies.

A: If your user name is eight characters long, you may be experiencing a bug in the IBM POE environment. The only fix at the time this was written was to use an account whose user name was seven characters or less. Ask your IBM representative about PMR 4017X (poe with userids of length eight fails) and the associated APAR IX56566.

6.12 Programs fail at startup

6.12.1 General

1. **Q:** With some systems, you might see

```
/lib/dld.sl: Bind-on-reference call failed
/lib/dld.sl: Invalid argument
```

(This example is from HP-UX), or

```
ld.so: libc.so.2: not found
```

(This example is from SunOS 4.1; similar things happen on other systems).

A: The problem here is that your program is using shared libraries, and the libraries are not available on some of the machines that you are running on. To fix this, relink your program without the shared libraries. To do this, add the appropriate command-line options to the link step. For example, for the HP system that produced the errors above, the fix is to use `-Wl,-Bimmediate` to the link step. For Solaris, the appropriate option is `-Bstatic`.

6.12.2 Workstation Networks

1. **Q:** I can run programs using a small number of processes, but once I ask for more than 4–8 processes, I do not get output from all of my processes, and the programs never finish.

A: We have seen this problem with installations using AFS. The remote shell program, `rsh`, supplied with some AFS systems limits the number of jobs that can use standard output. This seems to prevent some of the processes from exiting as well, causing the job to hang. There are four possible fixes:

- (a) Use a different `rsh` command. You can probably do this by putting the directory containing the non-AFS version first in your `PATH`. This option may not be available to you, depending on your system. At one site, the non-AFS version was in `'/bin/rsh'`.
- (b) Use the secure server (`serv_p4`). See the discussion in the Users Guide.
- (c) Redirect all standard output to a file. The MPE routine `MPE_IO_Stdout_to_file` may be used to do this.

- (d) Get a fixed `rsh` command. The likely source of the problem is an incorrect usage of the `select` system call in the `rsh` command. If the code is doing something like

```
int mask;
mask |= 1 << fd;
select( fd+1, &mask, ... );
```

instead of

```
fd_set mask;
FD_SET(fd, &mask);
select( fd+1, &mask, ... );
```

then the code is incorrect (the `select` call changed to allow more than 32 file descriptors many years ago, and the `rsh` program (or programmer!) hasn't changed with the times).

A fourth possibility is to get an AFS version of `rsh` that fixes this bug. As we are not running AFS ourselves, we do not know whether such a fix is available.

2. **Q:** Not all processes start.

A: This can happen when using the `ch_p4` device and a system that has extremely small limits on the number of remote shells you can have. Some systems using “Kerberos” (a network security package) allow only three or four remote shells; on these systems, the size of `MPI_COMM_WORLD` will be limited to the same number (plus one if you are using the local host).

The only way around this is to try the secure server; this is documented in the `MPICH` installation guide. Note that you will have to start the servers “by hand” since the `chp4_serve` script uses remote shell to start the servers.

6.13 Programs fail after starting

6.13.1 General

1. **Q:** I use `MPI_Allreduce`, and I get different answers depending on the number of processes I'm using.

A: The MPI collective routines may make use of associativity to achieve better parallelism. For example, an

```
MPI_Allreduce( &in, &out, MPI_DOUBLE, 1, ... );
```

might compute

$$(((((((a + b) + c) + d) + e) + f) + g) + h)$$

or it might compute

$$((a + b) + (c + d)) + ((e + f) + (g + h)),$$

where a, b, \dots are the values of `in` on each of eight processes. These expressions are equivalent for integers, reals, and other familiar objects from mathematics but are *not*

equivalent for datatypes, such as floating point, used in computers. The association that MPI uses will depend on the number of processes, thus, you may not get exactly the same result when you use different numbers of processes. Note that you are not getting a wrong result, just a different one (most programs assume the arithmetic operations are associative). All processes do get the same result from a single call to `MPI_Allreduce`.

2. **Q:** My Fortran program fails with a BUS error.

A: The C compiler that MPICH was built with and the Fortran compiler that you are using have different alignment rules for variables of type like `DOUBLE PRECISION`. For example, the GNU C compiler `gcc` may assume that all doubles are aligned on eight-byte boundaries, but the Fortran language requires only that `DOUBLE PRECISION` align with `INTEGERs`, which may be four-byte aligned.

There is no good fix. Consider rebuilding MPICH with a C compiler that supports weaker data alignment rules. Some Fortran compilers will allow you to force eight-byte alignment for `DOUBLE PRECISION` (for example, `-dalign` or `-f` on some Sun Fortran compilers); note though that this may break some correct Fortran programs that exploit Fortran's storage association rules.

Some versions of `gcc` may support `-munaligned-doubles`; MPICH should be rebuilt with this option if you are using `gcc`, version 2.7 or later. MPICH attempts to detect and use this option where available.

3. **Q:** I'm using `fork` to create a new process, or I'm creating a new thread, and my code fails.

A: The MPICH implementation is not thread safe and does not support either `fork` or the creation of new processes. Note that the MPI specification is thread safe, but implementations are not required to be thread safe. At this writing, few implementations are thread-safe, primarily because this reduces the performance of the MPI implementation (you at least need to check to see if you need a thread lock, actually getting and releasing the lock is even more expensive).

The MPICH implementation supports the `MPI_Init_thread` call; with this call, new in MPI-2, you can find out what level of thread support the MPI implementation supports. As of version 1.2.0 of MPICH, only `MPI_THREAD_SINGLE` is supported. We believe that version 1.2.0 and later support `MPI_THREAD_FUNNELED`, and some users have used MPICH in this mode (particularly with OpenMP), but we have not rigorously tested MPICH for this mode. Future versions of MPICH will support `MPI_THREAD_MULTIPLE`.

Q: C++ programs execute global destructors (or constructors) more times than expected. For example:

```
class Z {
public:
    Z() { cerr << "*Z" << endl; }
    ~Z() { cerr << "+Z" << endl; }
};

Z z;
```

```

int main(int argc, char **argv) {
    MPI_Init(&argc, &argv);
    MPI_Finalize();
}

```

when running with the `ch_p4` device on two processes executes the destructor twice for each process.

A: The number of processes running before `MPI_Init` or after `MPI_Finalize` is not defined by the MPI standard; you can not rely on any specific behavior. In the `ch_p4` case, a new process is `forked` to handle connection requests; it terminates with the end of the program.

You can use the threaded listener with the `ch_p4` device, or use the `ch_p4mpd` device instead. Note, however, that this code is not portable because it relies on behavior that the MPI standard does not specify.

6.13.2 HPUX

1. **Q:** My Fortran programs seem to fail with `SIGSEGV` when running on HP workstations.

A: Try compiling and linking the Fortran programs with the option `+T`. This *may* be necessary to make the Fortran environment correctly handle interrupts used by `MPICH` to create connections to other processes.

6.13.3 LINUX

1. **Q:** Processes fail with messages like

```
p0_1835: p4_error: Found a dead connection while looking for messages: 1
```

A: What is happening is that the TCP implementation on this platform is deciding that the connection has “failed” when it really hasn’t. The current `MPICH` implementation assumes that the TCP implementation will not close connections and has no code to reanimate failed connections. Future versions of `MPICH` will work around this problem.

In addition, some users have found that the single processor Linux kernel is more stable than the SMP kernel.

6.14 Trouble with Input and Output

6.14.1 General

1. **Q:** I want output from `printf` to appear immediately.

A: This is really a feature of your C and/or Fortran runtime system. For C, consider

```
setbuf( stdout, (char *)0 );
```

or

```
setvbuf( stdout, NULL, _IONBF, 0 );
```

6.14.2 Workstation Networks

1. **Q:** I want standard output (`stdout`) from each process to go to a different file.

A: MPICH has no built-in way to do this. In fact, it prides itself on gathering the `stdouts` for you. You can do one of the following:

- (a) Use Unix built-in commands for redirecting `stdout` from inside your program (`dup2`, etc.). The MPE routine `MPE_IO_Stdout_to_file`, in `'mpe/src/mpe_io.c'`, shows one way to do this. Note that in Fortran, the approach of using `dup2` will work only if the Fortran `PRINT` writes to `stdout`. This is common but by no means universal.
- (b) Write explicitly to files instead of to `stdout` (use `fprintf` instead of `printf`, etc.). You can create the file name from the process's rank. This is the most portable way.

6.14.3 HP-UX

1. **Q:** When trying to run `upshot` under HP-UX, I get error messages like

```
set: Variable name must begin with a letter.
```

or

```
upshot: syntax error at line 35: '(' unexpected
```

A: Your version of HP-UX limits the shell names to very short strings. `Upshot` is a program that is executed by the `wish` shell, and for some reason HP-UX is both refusing to execute in this shell and then trying to execute the `upshot` program using your current shell (e.g., `'sh'` or `'csh'`), instead of issuing a sensible error message about the command name being too long. There are two possible fixes:

- (a) Add a link with a much shorter name, for example

```
ln -s /usr/local/tk3.6/bin/wish /usr/local/bin/wish
```

Then edit the `upshot` script to use this shorter name instead. This may require root access, depending on where you put the link.

- (b) Create a regular shell program containing the lines

```
#!/bin/sh
/usr/local/tk3.6/bin/wish -f /usr/local/mpi/bin/upshot
```

(with the appropriate names for both the `'wish'` and `'upshot'` executables).

Also, file a bug report with HP. At the very least, the error message here is wrong; also, there is no reason to restrict general shell choices (as opposed to login shells).

6.15 Special debugging arguments

These are currently undocumented, and some require configure options to have been specified (like `-mpipktsize` and `-chmemdebug`). The option `-mpiversion` is useful for finding out how your installation of MPICH was configured and exactly what version it is. The options include:

- mpedbg** If an error occurs, start `xterms` attached to the process that generated the error. Requires the MPICH be configured with `-mpedbg` and works on only some workstations systems.
- mpiversion** Print out the version and configuration arguments for the MPICH implementation being used.

These arguments are provided to the program, not to `mpirun`. For example,

```
mpirun -np 2 a.out -mpiversion
```

Acknowledgments

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Appendices

A Frequently Asked Questions

- Introduction
- Installing MPICH
- Using MPICH
- Permission Denied

- Notes on getting MPICH running on RH 7.2
- poll: protocol failure during circuit creation
- Using SSH
- SIGSEGV
- Compiler Switches
- MPMD (Multiple Program Multiple Data) Programs
- Reporting problems and support
- Algorithms used in MPICH
- Jumpshot and X11

A.1 Introduction

MPICH is a freely available, portable implementation of MPI, the Standard for message-passing libraries.

A.2 Installing MPICH

Building and installing MPICH often requires only

```
./configure --prefix=/home/me/mpich
make
make install
```

where the value of the `--prefix` argument to `configure` is the directory in which MPICH should be installed. See the *Installation Guide* for more detailed instructions.

A.3 Using MPICH

Information on using MPICH can be found in the *Installation and Users' Guide*. There are versions of this guide for each MPICH device; make sure that you read the one that applies to your device. You can find out which device is installed by running a simple MPI program with the option `-mpiversion`:

```
mpirun -np 1 a.out -mpiversion
```


A.4 Permission Denied

Question:

When I use mpirun, I get the message `Permission denied, connection reset by peer, or poll: protocol failure in circuit setup` when trying to run MPICH.

Answer:

If you see something like this

```
% mpirun -np 2 cpi
Permission denied.
```

(or `connection reset by peer` or `poll: protocol failure in circuit setup`) when using the `ch_p4` device, it probably means that you do not have permission to use rsh to start processes. The script `tstmachines` can be used to test this. For example, if the architecture type (the `-arch` argument to `configure`) is `sun4`, then try

```
tstmachines sun4
```

If this fails, then you may need a `.rhosts` or `/etc/hosts.equiv` file (you may need to see your system administrator) or you may need to use the p4 server. Another possible problem is the choice of the remote shell program; some systems have several. Check with your systems administrator about which version of rsh or remsh you should be using.

If your system allows a `.rhosts` file, do the following:

- Create a file `.rhosts` in your home directory
- Change the protection on it to user read/write only: `chmod og-rwx .rhosts`.
- Add one line to the `.rhosts` file for each processor that you want to use. The format is

```
host username
```

For example, if your username is `doe` and you want to use machines `a.our.org` and `b.our.org`, your `.rhosts` file should contain

```
a.our.org doe
b.our.org doe
```

Note the use of fully qualified host names (some systems require this).

On networks where the use of `.rhosts` files is not allowed, (such as the one in MCS at Argonne), you should use the p4 server to run on machines that are not trusted by the machine that you are initiating the job from.

Finally, you may need to use a non-standard rsh command within MPICH. MPICH must be reconfigured with `-rsh=command_name`, and perhaps also with `-rshno1` if the remote shell command does not support the `-1` argument. Systems using Kerberos and/or AFS may need this.

A.5 Notes on getting MPICH running on RH 7.2

Introduction The purpose of this document is to describe the steps necessary to allow MPICH processes to be started and to communicate with one another. The installation that we are focusing on is a RedHat 7.2 installation with medium security (the default). While other distributions will certainly vary, this is a good example of the sorts of problems that one might run across.

There are three methods for starting MPICH processes that are typically used on clusters today. These are rsh, ssh, and mpd.

We will first describe getting the rsh service working. We will include rlogin in this process because it is helpful for testing. Next we will describe getting ssh working and enabling the ssh-agent to allow for logins without password typing. Finally we will discuss issues related to process communication in MPICH and firewalls.

Enabling rsh By default the rsh server is not installed, and it is necessary for use of the rsh service in starting MPICH processes. The rsh server, in.rshd, is part of the rsh-server RPM. This RPM is located on the first disc of the RedHat 7.2 distribution. The rlogin server, in.rlogind, is also included in this package.

The xinetd server controls the availability of the rsh and rlogin services. This server is installed by default, but by default rsh and rlogin services are disabled. To enable these services, you must edit the files ‘/etc/xinetd.d/rsh’ and ‘/etc/xinetd.d/rlogin’. Here is the rsh file as it looks by default:

```
# default: on
# description: The rshd server is the server for the rcmd(3) routine and, \
#               consequently, for the rsh(1) program. The server provides \
#               remote execution facilities with authentication based on \
#               privileged port numbers from trusted hosts.
service shell
{
    socket_type          = stream
    wait                 = no
    user                 = root
    log_on_success        += USERID
    log_on_failure        += USERID
    server                = /usr/sbin/in.rshd
    disable               = yes
}
```

You must enable the service by changing “disable = yes” to “disable = no”. The same must be done to the rlogin config file to enable that service.

At this point the xinetd daemon must be restarted to register these changes:

```
/etc/rc.d/init.d/xinetd restart
```

At this point you should receive a "Permission denied." if you attempt a command such as "rsh localhost hostname" as a non-root user (or as root for that matter).

To allow users to rsh without passwords you need to edit '/etc/hosts.equiv', the system-wide host file for rsh and rlogin. This file should hold hostnames of machines that you would like users to be able to start MPICH processes from. For example, simply adding:

```
localhost.localdomain
```

Should allow users to perform the command "rsh localhost hostname" successfully. Likewise adding other hostnames will allow users on those hosts to rsh to this host.

However, there is another catch! By default (with medium security) packet filtering is enabled as well, and this will prevent users from remote hosts from connecting to this machine using the rsh or rlogin services. This packet filter, or firewall, is administered using the ipchains package (which is installed by default).

The firewall configuration is written out by a program called lokkit at installation time (I think). The configuration is stored in '/etc/sysconfig/ipchains' and by default looks like this:

```
# Firewall configuration written by lokkit
# Manual customization of this file is not recommended.
# Note: ifup-post will punch the current nameservers through the
#       firewall; such entries will *not* be listed here.
:input ACCEPT
:forward ACCEPT
:output ACCEPT
-A input -s 0/0 67:68 -d 0/0 67:68 -p udp -i eth0 -j ACCEPT
-A input -s 0/0 67:68 -d 0/0 67:68 -p udp -i eth1 -j ACCEPT
-A input -s 0/0 -d 0/0 -i lo -j ACCEPT
-A input -p tcp -s 0/0 -d 0/0 0:1023 -y -j REJECT
-A input -p tcp -s 0/0 -d 0/0 2049 -y -j REJECT
-A input -p udp -s 0/0 -d 0/0 0:1023 -j REJECT
-A input -p udp -s 0/0 -d 0/0 2049 -j REJECT
-A input -p tcp -s 0/0 -d 0/0 6000:6009 -y -j REJECT
-A input -p tcp -s 0/0 -d 0/0 7100 -y -j REJECT
```

While an in-depth discussion of ipchains rules is outside the context of this document, it's worth talking about how this works a bit. First, the rules are applied in order from top of the list to the bottom of the list. The argument to -j says what to do if a packet matches; it's usually either ACCEPT (let the packet in), or REJECT (toss it out). If a packet makes it through the entire list then the default policy is applied. In this case the default policy is ACCEPT.

The following line tells the packet filter to allow all localhost (-i lo) traffic to pass unmolested:

```
-A input -s 0/0 -d 0/0 -i lo -j ACCEPT
```

This line blocks all new TCP connections going to ports 0-1023, which is the range of most services, including rsh/rlogin:

```
-A input -p tcp -s 0/0 -d 0/0 0:1023 -y -j REJECT
```

We're going to modify this file to allow rsh and rlogin traffic.

```
# Firewall configuration written by lokkit
# Manual customization of this file is not recommended.
# Note: ifup-post will punch the current nameservers through the
#       firewall; such entries will *not* be listed here.
:input ACCEPT
:forward ACCEPT
:output ACCEPT
-A input -s 0/0 67:68 -d 0/0 67:68 -p udp -i eth0 -j ACCEPT
-A input -s 0/0 67:68 -d 0/0 67:68 -p udp -i eth1 -j ACCEPT
-A input -s 0/0 -d 0/0 -i lo -j ACCEPT
#
# New rules for rlogin/rsh traffic, incoming or outgoing
#
-A input -p tcp -s 0/0 -d 0/0 513 -b -j ACCEPT
-A input -p tcp -s 0/0 -d 0/0 514 -b -j ACCEPT
#
# End of new rules
#
-A input -p tcp -s 0/0 -d 0/0 0:1023 -y -j REJECT
-A input -p tcp -s 0/0 -d 0/0 2049 -y -j REJECT
-A input -p udp -s 0/0 -d 0/0 0:1023 -j REJECT
-A input -p udp -s 0/0 -d 0/0 2049 -j REJECT
-A input -p tcp -s 0/0 -d 0/0 6000:6009 -y -j REJECT
-A input -p tcp -s 0/0 -d 0/0 7100 -y -j REJECT
```

At this point users on remote systems with accounts on this system should be able to rsh/rlogin to this machine without using a password.

Enabling ssh Enabling ssh is somewhat easier.

First the ssh server, sshd, must be installed. This is part of the openssh-server RPM. This RPM is located on the first disc of the RedHat 7.2 distribution.

Once the server is installed, it must be started:

```
/etc/rc.d/init.d/sshd start
```

The service will be automatically started on reboot.

At this point ssh on the localhost should work, although a password will still be required. However, our firewall rules will be preventing connections from other machines.

We again modify `/etc/sysconfig/ipchains`, this time to allow ssh traffic in and out. See the above section for a discussion of what we are doing here.

```
# Firewall configuration written by lokkit
# Manual customization of this file is not recommended.
# Note: ifup-post will punch the current nameservers through the
#       firewall; such entries will *not* be listed here.
:input ACCEPT
:forward ACCEPT
:output ACCEPT
-A input -s 0/0 67:68 -d 0/0 67:68 -p udp -i eth0 -j ACCEPT
-A input -s 0/0 67:68 -d 0/0 67:68 -p udp -i eth1 -j ACCEPT
-A input -s 0/0 -d 0/0 -i lo -j ACCEPT
#
# New rules for ssh traffic, incoming or outgoing
#
-A input -p tcp -s 0/0 -d 0/0 22 -b -j ACCEPT
#
# End of new rules
#
-A input -p tcp -s 0/0 -d 0/0 0:1023 -y -j REJECT
-A input -p tcp -s 0/0 -d 0/0 2049 -y -j REJECT
-A input -p udp -s 0/0 -d 0/0 0:1023 -j REJECT
-A input -p udp -s 0/0 -d 0/0 2049 -j REJECT
-A input -p tcp -s 0/0 -d 0/0 6000:6009 -y -j REJECT
-A input -p tcp -s 0/0 -d 0/0 7100 -y -j REJECT
```

At this point users on remote systems should be able to ssh into the machine, but they will still need a password.

Users should set up a private/public authentication key pair in order for ssh to operate without passwords. This process is documented in the installation guide, but a summary of the steps for RH7.2 will be included here.

First run the `"ssh-keygen -t rsa"` application to create the private/public key pair. By default this will create the files `~/.ssh/id_rsa` and `~/.ssh/id_rsa.pub`. Use a password.

Next place the public key (`~/.ssh/id_rsa.pub`) in the file `~/.ssh/authorized_keys`. If more than one machine is going to be used, then this key must be put in the `~/.ssh/authorized_keys` file on each machine. The permissions on the `.ssh` directory should be set to 700; otherwise the `sshd` may choose to not accept the keys.

This will allow you to connect using rsa keys rather than simple UNIX passwords. The next step is to enable an SSH agent so that you do not need to repeatedly type your password.

The agent is started with "`ssh-agent <cmd>`". Typically `<cmd>` is `$SHELL`, so that your default shell is started. The agent will then handle authentication on your behalf any time you attempt to use ssh from this shell. To give the ssh-agent your password, type "`ssh-add`". This will query you for the passphrase that accompanies your rsa key.

Once you have completed this, you will be able to ssh to other systems on which your key is authorized without typing a password.

Interprocess communication MPICH processes use the standard UNIX mechanisms for allocating ports for intercommunication. Using this mechanism processes are given ports in the range of 1024–65535.

Unfortunately for us, the default firewall configuration blocks some port ranges that our MPICH processes might be given to use for communication. This leads to a situation where MPICH applications will occasionally fail to communicate (when they happen to get the wrong port value).

We're going to modify the ipchains configuration file to remove lines disabling ranges of ports that our processes might use for intercommunication.

The two default rules of interest are the following:

```
-A input -p tcp -s 0/0 -d 0/0 6000:6009 -y -j REJECT
-A input -p tcp -s 0/0 -d 0/0 7100 -y -j REJECT
```

The first blocks incoming TCP connections to ports 6000-6009 (often used by X), while the second blocks incoming TCP connections to port 7100 (often used by the X font server).

We simply remove these rules:

```
# Firewall configuration written by lokkit
# Manual customization of this file is not recommended.
# Note: ifup-post will punch the current nameservers through the
#       firewall; such entries will *not* be listed here.
:input ACCEPT
:forward ACCEPT
:output ACCEPT
-A input -s 0/0 67:68 -d 0/0 67:68 -p udp -i eth0 -j ACCEPT
-A input -s 0/0 67:68 -d 0/0 67:68 -p udp -i eth1 -j ACCEPT
-A input -s 0/0 -d 0/0 -i lo -j ACCEPT
-A input -p tcp -s 0/0 -d 0/0 0:1023 -y -j REJECT
-A input -p tcp -s 0/0 -d 0/0 2049 -y -j REJECT
-A input -p udp -s 0/0 -d 0/0 0:1023 -j REJECT
-A input -p udp -s 0/0 -d 0/0 2049 -j REJECT
#
# Removed these rules to eliminate chance of MPICH comm. failure
#
# -A input -p tcp -s 0/0 -d 0/0 6000:6009 -y -j REJECT
```

```
# -A input -p tcp -s 0/0 -d 0/0 7100 -y -j REJECT
#
# End of removed rules
#
```

This modification, in conjunction with one to allow process startup, should prepare your system for MPICH jobs.

A.6 poll: protocol failure during circuit creation

You may see this message if you attempt to run too many MPI programs in a short period of time. For example, in Linux and when using the `ch_p4` device (without the secure server or `ssh`), MPICH uses `rsh` to start the MPI processes. Depending on the particular Linux distribution and version, there may be a limit of as few as 40 processes per minute. When running the MPICH test suite or starting short parallel jobs from a script, it is possible to exceed this limit.

To fix this, you can do one of the following:

1. Wait a few seconds between running parallel jobs. You may need to wait up to a minute.
2. Modify `/etc/inetd.conf` to allow more processes per minute for `rsh`. For example, change

```
shell stream tcp nowait root /etc/tcpd2 in.rshd
```

to

```
shell stream tcp nowait.200 root /etc/tcpd2 in.rshd
```

3. Use the `ch_p4mpd` device or the secure server option of the `ch_p4` device instead. Neither of these relies on `inetd`.

A.7 Using SSH

The secure shell (`ssh`) may be used with the `ch_p4` device, but requires careful setup. See *configuring with ssh* in the *Installation and User's* manual.

Make sure that `ssh` is set up to *not* require a password. The command

```
ssh -n 'hostname' date
```

should return the date without any prompts for passwords. See the installation manual if you have problems.

A.8 SIGSEGV

If your program fails with

```
p4_error: interrupt SIGSEGV
```

the problem is probably not with MPI. Instead, check for program bugs including

1. Array overwrites or accesses beyond array bounds. Be particularly careful of `a[size]` in C, where `a` is declared as `int a[size]`.
2. Invalid pointers, including null pointers.
3. Missing or mismatched parameters to subroutines or functions. Fortran users should check that all MPI calls include the integer error return parameter and that any status variable is dimensioned as an array of size `MPI_STATUS_SIZE`.

A.9 Compiler Switches

Normally, you should let `configure` determine compiler switches. However, you can use the configure options `-cflags=...` and `-fflags=...` to specify special flags. See also *compiler switches*.

A.10 MPMD (Multiple Program Multiple Data) Programs

MPICH, depending on the device, supports MPMD programs. However, the `mpirun` script currently does not support MPMD programs. For the `ch_p4` device, the user must create a procgroup file and invoke the program that will have rank zero in `MPI_COMM_WORLD` with the command-line option `-p4pg filename`. See the *Installation and User's Guide* for more information.

A.11 Reporting problems and support

1. First, check *the list of known bugs and patches* for the problem you are seeing.
2. Also check the troubleshooting guides in the *Installation* and *Users* guides.
3. If that doesn't help, send mail to `mpi-bugs@mcs.anl.gov`.

A.12 Algorithms used in MPICH

1. Does MPICH use IP Multicast for `MPI_Bcast`?

No. In principle, MPICH could use multicast, but in practice this would be very difficult. To start with, IP multicast is unreliable; additional code to make it reliable needs to be added. In fact, there is an effort to provide a reliable multicast, built on top of the unreliable multicast. The second problem is that not all systems allow

user programs (or *any* program) to perform an IP multicast. In fact, that is the case for the systems that we have been developing on. Thus, we will always need the point-to-point version. There is a fairly easy way to replace any collective routine in MPI, but no-one has offered us a multicast-based `MPI_Bcast` yet...

A.13 Jumpshot and X11

Jumpshot relies on the Swing toolkit to perform its graphics operations. We have noticed some problems with Swing and X11 window system support on some PC's running Windows. In particular, the combination of SecureCRT and Exceed or SecureCRT and Xfree86 often have problems (sometimes serious, including hangs and crashes). For now, the only workaround is to use OpenSSH and Xfree86 if possible when using a Windows PC.

B History of MPICH

MPICH was developed during the MPI standards process to provide feedback to the MPI Forum on implementation and usability issues. With the release of the MPI standard, MPICH was designed to provide an implementation of the MPI standard that could replace the proprietary message-passing systems on the massively parallel computers of the day, such as the Intel Paragon, IBM SP, and TMC CM5. MPICH used an early version of the abstract device interface (ADI), based on the Chameleon [?] portability system, to provide a light-weight implementation layer. To enable development on desktop systems, a device layered on top of the P4 [2] system was used. Over time, other devices were developed; as systems have vanished (e.g., the TMC CM5 and the Ncube), these devices have been dropped from our distribution. Because MPICH used P4 for workstation networks, MPICH has supported both MIMD programming and heterogeneous clusters from the very beginning.

Because MPICH was designed to enable ports to other systems, many parallel computer vendors and research groups have used MPICH as the basis for their implementation. Many users are now familiar only with the version of MPICH that uses the `ch_p4` device for workstation and Beowulf clusters. However, MPICH continues to support other systems and continues to serve as a platform for research into MPI implementations.

C File Manifest

This section briefly describes the files and directories at the top level of the MPICH source tree.

COPYRIGHT Copyright statement. This code is free but not public domain. It is copyrighted by the University of Chicago and Mississippi State University.

Makefile.in Template for the 'Makefile', which will be produced when you run `configure`.

MPI-2-C++ The C++ system from Notre Dame. It includes the C++ bindings for the MPI-1 functions.

README Basic information and instructions for configuring.

aclocal.m4 Used for building ‘**configure**’ from ‘**configure.in**’; not needed for most installations. The file ‘**aclocal_tcl.m4**’ is included by ‘**aclocal.m4**’.

ccbugs Directory for programs that test the C compiler during configuration, to make sure that it will be able to compile the system.

configure The script that you run to create Makefiles throughout the system.

configure.in Input to **autoconf** that produces **configure**.

doc Assorted tools for producing documentation, together with this manual.

examples Directory containing further directories of example MPI programs. Of particular note are **basic**, with a few small examples to try first, **test**, with a test suite for exercising MPICH, and **perftest**, containing benchmarking code.

include The include files, both user and system.

bin Contains the programs and executable scripts, such as **mpicc** and **mpirun**, used to build and run MPI programs.

man Man pages for MPI, MPE, and internal routines.

mpe The source code for the MPE extensions for logging and X graphics. The **contrib** directory contains examples. Best are the **mandel** and **mastermind** subdirectories. The **profiling** subdirectory contains the profiling subsystem, including a system for automatically generating the “wrappers” for the MPI profiling interface. MPE also includes the performance visualization programs, such as **jumpshot** (see Section 3.7.1).

mpid The source code for the various “devices” that customize MPICH for a particular machine, operating system, and environment.

romio The ROMIO parallel I/O system, which includes an implementation of most of the MPI-2 parallel I/O standard.

src The source code for the portable part of MPICH. There are subdirectories for the various parts of the MPI specification.

util Utility programs and files.

www HTML versions of the **man** pages.

D Configure Usage

The command **configure -help** will print out

```
Configuring with args --help
Configuring MPICH Version 1.2.3 (pre-alpha) of : 2001/08/20 18:08:23
Usage: ./configure [--with-arch=ARCH_TYPE] [--with-comm=COMM_TYPE]
        [--with-device=DEVICE]
```

```

[--with-mpe] [--without-mpe]
[--disable-f77] [--disable-f90] [--with-f90nag] [--with-f95nag]
[--disable-f90modules]
[--disable-gencat] [--disable-doc]
[--enable-cxx ] [--disable-cxx]
[--enable-mpedbg] [--disable-mpedbg]
[--enable-devdebug] [--disable-devdebug]
[--enable-debug] [--disable-debug]
[--enable-traceback] [--disable-traceback]
[--enable-long-long] [--disable-long-long]
[--enable-long-double] [--disable-long-double]
[-prefix=INSTALL_DIR]

[-c++=[C++_COMPILER] ] [noc++]
[-opt=OPTFLAGS]
[-cc=C_COMPILER] [-fc=FORTRAN_COMPILER]
[-clinker=C_LINKER] [-flinker=FORTRAN_LINKER]
[-c++linker=CC_LINKER]
[-cflags=CFLAGS] [-fflags=FFLAGS] [-c++flags=CCFLAGS]
[-optcc=C_OPTFLAGS] [-optf77=F77_OPTFLAGS]
[-f90=F90_COMPILER] [-f90flags=F90_FLAGS]
[-f90inc=INCLUDE_DIRECTORY_SPEC_FORMAT_FOR_F90]
[-f90linker=F90_LINKER]
[-f90libpath=LIBRARY_PATH_SPEC_FORMAT_FOR_F90]
[-lib=LIBRARY] [-mpilibname=MPILIBNAME]
[-mpe_opts=MPE_OPTS]
[-make=MAKEPGM ]
[-memdebug] [-ptrdebug] [-tracing] [-dlast]
[-listener_sig=SIGNAL_NAME]
[-cross]
[-adi_collective]
[-automountfix=AUTOMOUNTFIX]
[-noranlib] [-ar_nolocal]
[-rsh=RSHCOMMAND] [-rshnol]
[-noromio] [-file_system=FILE_SYSTEM]
[-p4_opts=P4_OPTS]

```

where

ARCH_TYPE	= the type of machine that MPI is to be configured for
COMM_TYPE	= communications layer or option to be used
DEVICE	= communications device to be used
INSTALL_DIR	= directory where MPI will be installed (optional)
MPE_OPTS	= options to pass to the mpe configure
P4_OPTS	= options to pass to the P4 configure (device=ch_p4)
C++_COMPILER	= default is to use xlc, g++, or CC (optional)
OPTFLAGS	= optimization flags to give the compilers (e.g. -g)
CFLAGS	= flags to give C compiler
FFLAGS	= flags to give Fortran compiler
MAKEPGM	= version of make to use
LENGTH	= Length of message at which ADI switches from short to long message protocol
AUTOMOUNTFIX	= Command to fix automounters
RSHCOMMAND	= Command to use for remote shell
MPILIBNAME	= Name to use instead of mpich in the name of the MPI library. If set, libMPILIBNAME will be used instead

or libmpich. This can be used on systems with several different MPI implementations.

FILE_SYSTEM = name of the file system ROMIO is to use. Currently supported values are nfs, ufs, pfs (Intel), piofs (IBM), hfs (HP), sfs (NEC), and xfs (SGI).

SIGNAL_NAME = name of the signal for the P4 (device=ch_p4) device to use to indicate that a new connection is needed. By default, it is SIGUSR1.

All arguments are optional, but if 'arch', 'comm', or 'prefix' arguments are provided, there must be only one. 'arch' must be specified before 'comm' if they both appear.

Packages that may be included with MPICH

```
--with-device=name      - Use the named device for communication. Known
                           names include ch_p4, ch_mpl, ch_shmem, and globus2.
                           If not specified, a default is chosen. Special
                           options for the device are specified after the
                           device name, separated by a colon. E.g.,
                           --with-device=globus2:-flavor=mpi,nothreads
--with-romio[=OPTIONS] - Use ROMIO to provide MPI-I/O from MPI-2 (default).
                           The options include -file_system=FSTYPE, where
                           fstype can be any combination of nfs, ufs,
                           pfs (intel), piofs (IBM), hfs (HP), sfs (NEC), and
                           xfs (SGI), combined with '+'. If romio is not
                           included, the Fortran 90 modules cannot be built.
--with-mpe              - Build the MPE environment (default)
--with-f90nag           - Choose the NAG f90 compiler for Fortran
                           (preliminary version intended for use *instead*
                           of a Fortran 77 compiler)
--with-f95nag           - Choose the NAG f95 compiler for Fortran
--with-cross=file       - Use the file for cross compilation. The file
                           should contain assignments of the form
                           CROSS_SIZEOF_INT=4
                           for each cross compilation variable. The command
                           egrep 'CROSS_[A-Z]*=' configure | sed 's/=.*/g'
                           will list each variable.
```

You can use --without-<featurename> to turn off a feature (except for device).

Options for device ch_lfshmem:

```
--with-device=ch_lfshmem[:~usesysv]
```

The option '~usesysv' applies to the ch_shmem device, and causes the device to attempt and use System V shared memory and semaphore routines, rather than what would be chosen by default (often mmap or a system-specific method).

Options for device ch_meiko:

```
--with-device=ch_meiko
```

Options for device ch_mpl:

```
--with-device=ch_mpl
```

Options for device ch_p4:

./configure: ./mpid/ch_p4/setup_ch_p4: Permission denied

Options for device ch_p4mpd:

--with-device=ch_p4mpd[:**-listener_sig**=SIGNALNAME] [**-dlast**] [**-socksize**=BYTES]

The option '**-listener_sig**' applies to the ch_p4mpd device, and changes the signal that is used to signal that a new connection should be made. By default, SIGUSR1 is used.

The option '**-dlast**' causes the p4 device to remember the last few debugging messages, printing them out only when the job terminates abnormally.

The option '**-socksize**' changes the size of the socket buffers used.

Options for device ch_shmem:

--with-device=ch_shmem[:**-usesysv**]

The option '**-usesysv**' applies to the ch_shmem device, and causes the device to attempt and use System V shared memory and semaphore routines, rather than what would be chosen by default (often mmap or a system-specific method).

Options for device globus:

--with-device=globus[:**-globusdir**=GLOBUSDIR]

'**-globusdir**=GLOBUS' allows one to specify the location of an installed version of Globus. You can acquire Globus from <http://www.globus.org>."

Options for device globus2:

GLOBUS_INSTALL_PATH must be set

Features that may be included with MPICH

--enable-c++	- Build C++ interfaces to the MPI-1 routines (default)
--enable-f77	- Build Fortran 77 interfaces to the MPI routines (default)
--enable-weak-symbols	- Use weak symbols for MPI/PMPI routines. This uses weak symbols, if available, for the profiling interface (default)
--enable-debug	- Enable support for debuggers to access message queues
--enable-traceback	- Enable printing of a call stack when MPI and the user's program is built with certain compilers (currently only some versions of gcc are supported).
--enable-mpedbg	- Enable the -mpedbg command-line argument (e.g., errors can start an xterm running a debugger). Only works with some workstation systems.
--enable-sharedlib	- Attempt to build shared libraries. Static
--enable-sharedlib=dir	libraries are always built. If a directory is specified, the shared libraries will be placed in that directory. This can be used to place the shared libraries in a uniform location in local disks on a cluster.
--enable-f90modules	- Build Fortran 90 module support (default if a Fortran 90 or 95 compiler is found). If ROMIO is not built, no Fortran 90 modules will be built.

The following are intended for MPI implementors and debugging of configure

<code>--enable-strict</code>	- Try and build MPICH using strict options in Gnu gcc
<code>--enable-echo</code>	- Cause configure to echo what it does
<code>--enable-devdebug</code>	- Enable debugging code in the ADI.

You can use `--disable-<featurename>` to turn off a feature.

Notes on configure usage:

The suggestions for GNU configure usage suggest that configure not be used to build different tools, only controlling some basics of the features enabled or the packages included. Our use of configure does not follow these rules because configure is too useful but we need the flexibility that allows the user to produce variations of MPICH.

More notes on command-line parameters:

You can select a different C and Fortran compiler by using the `'-cc'` and `'-fc'` switches. The environment variables `'CC'` and `'FC'` can also provide values for these but their settings may be overridden by the configure script. Using `'-cc=$CC -fc=$FC'` will force configure to use those compilers.

The option `'-opt'` allows you to specify optimization options for the compilers (both C and Fortran). For example, `'-opt=-O'` chooses optimized code generation on many systems. `'-optcc'` and `'-optf77'` allow you to specify options for just the C or Fortran compilers. Use `-cflags` and `-fflags` for options not related to optimization.

Note that the `'-opt'` options are not passed to the `'mpicc'`, `'mpif77'`, `'mpiCC'`, and `'mpif90'` scripts. The `'-opt'` options are used only in building MPICH.

The option `'-lib'` allows you to specify the location of a library that may be needed by a particular device. Most devices do NOT need this option; check the installation instructions for those that might.

The option `'-make'` may be used to select an alternate make program. For example, to make use of VPATH builds (building in one directory with the source in a different directory), `-make=gnumake` may be required.

The option `'--disable-short-longs'` may be used to suppress support for the C types `'long long'` (a common extension) and `'long double'` (ANSI/ISO C) when they are the same size as `'long'` and `'double'` respectively. Some systems allow these long C types, but generate a warning message when they are used; this option may be used to suppress these messages (and support for these types). `'--disable-long-long'` disables just `'long long'`; `'--disable-long-double'` disables just `'long double'`.

The option `'-ar_nolocal'` prevents the library archive command from attempting to use the local directory for temporary space. This option should be used when (a) there isn't much space (less than 20 MB) available in the partition where MPICH resides and (b) there is enough space in `/tmp` (or wherever `ar` places temporary files by default).

The option `'-noranlib'` causes the `'ranlib'` step (needed on some systems to build an object library) to be skipped. This is particularly useful on systems where `'ranlib'` is optional (allowed but not needed; because it is allowed, configure chooses to use it just in case) but can fail (some `'ranlib'`'s are implemented as scripts using `'ar'`; if they don't use the local directory, they can fail (destroying the library in the process) if the temporary directory (usually `'/tmp'`) does not have enough space. This has occurred on some OSF systems.

The option `'-rsh'` allows you to select an alternative remote shell command (by default, configure will use `'rsh'` or `'remsh'` from your `'PATH'`). If your remote shell command does not support the `'-l'` option (some AFS versions of `'rsh'` have this bug), also give the option `'-rshnol'`. These options are useful only when building a network version of MPICH (e.g., `'--with-device=ch_p4'`).

Special Tuning Options:

There are a number of options for tuning the behavior of the ADI (Abstract Device Interface) which is the low-level message-passing interface. These should NOT be used unless you are sure you know what you are doing.

The option `'-pkt_size=LENGTH'` allows you to choose the message length at which the ADI (Abstract Device Interface) switches from its short to long message format. `LENGTH` must be positive.

The option `'-adi_collective'` allows the ADI to provide some collective operations in addition to the basic point-to-point operations. Currently, most systems do not support this option (it is ignored) and on the others it has not been extensively tested.

Options for Experts:

The option `'-memdebug'` enables extensive internal memory debugging code. This should be used only if you are trying to find a memory problem (it can be used to help find memory problems in user code as well). Running programs with the option `'-mpidb memdump'` will produce a summary, when `'MPI_Finalize'` is called, of all unfreed memory allocated by MPI. For example, a user-created datatype that was not later freed would be reported.

The option `'-tracing'` enables tracing of internal calls. This should be used only for debugging the MPICH implementation itself.

The option `'-dlast'` enables tracing of the most recent operations performed by the device. These can be output when a signal (like `SIGINT`), error, or call to a special routine occurs. There is a performance penalty for this option, but it can be very useful for implementors attempting to debug problems.

Sample Configure Usage:

To make for running on Sun's running Solaris with `ch_p4` as the device,

and with the installation directory equal to the current directory:

```
./configure --with-device=ch_p4 --with-arch=solaris
make
```

Known devices are

- ch_nx (native Intel NX calls),
- ch_mpl (native IBM EUI or MPL calls),
- ch_p4 (p4)
- ch_p4mpd (p4 with the MPD startup system)
- globus2 (Globus: globus_io/vMPI)
- ch_meiko (for Meiko CS2, using NX compatibility library),
- ch_shmem (for shared memory systems, such as SMPs),
- ch_lfshmem (for shared memory systems, such as SMPs; uses
lock-free message buffers),
- ch_cenju3 (native NEC Cenju-3 calls)

The following devices were supported with ADI-1, but are currently unsupported. Please contact us if you are interested in helping us support these devices:

- meiko (for Meiko CS2, using elan tport library), and
- nx (for Intel Paragon),
- t3d (for the Cray T3D, using Cray shmem library).
- ch_nc (native nCUBE calls, requires -arch=ncube),
- ch_cmmd (native TMC CM-5 CMMD calls)

These are no longer distributed with the MPICH distribution.

Known architectures include (case is important)

- alpha (Compaq alpha)
- CRAY (CRAY XMP, YMP, C90, J90, T90)
- cray_t3d (CRAY T3D)
- EWS_UX_V (NEC EWS4800/360AD Series workstation. Untested.)
- freebsd (PC clones running FreeBSD)
- hpux (HP UX)
- intelnx (Intel i860 or Intel Delta)
- IRIX (synonym for sgi)
- IRIX32 (IRIX with 32bit objects -32)
- IRIXN32 (IRIX with -n32)
- IRIX64 (IRIX with 64bit objects)
- ksr (Kendall Square KSR1 and KSR2)
- LINUX (PC clones running LINUX)
- LINUX_ALPHA (Linux on Alpha processors)
- meiko (Meiko CS2)
- netbsd (PC clones running NetBSD)
- paragon (Intel Paragon)
- rs6000 (AIX for IBM RS6000)
- sgi (Silicon Graphics IRIX 4.x, 5.x or 6.x)
- sgi5 (Silicon Graphics IRIX 5.x on R4400's, for the MESHINE)
- solaris (Solaris)
- solaris86 (Solaris on Intel platforms)
- sppux (SPP UX)
- sun4 (SUN OS 4.x)
- SX_4_float0
(NEC SX-4; Floating point format float0)


```

                Conforms IEEE 754 standard.
C:      sizeof (int)      = 4; sizeof (float) = 4
FORTRAN: sizeof (INTEGER) = 4; sizeof (REAL)  = 4)
SX_4_float1
(NEC SX-4; Floating point format float1
        IBM floating point format.
C:      sizeof (int)      = 4; sizeof (float) = 4
FORTRAN: sizeof (INTEGER) = 4; sizeof (REAL)  = 4)
SX_4_float2
(NEC SX-4; Floating point format float2
        CRAY floating point format.
C:      sizeof (int)      = 4; sizeof (float) = 8
FORTRAN: sizeof (INTEGER) = 8; sizeof (REAL)  = 8)
!!! WARNING !!! This version will not run
                  together with FORTRAN routines.
                  sizeof (INTEGER) != sizeof (int)
SX_4_float2_int64
(NEC SX-4; Floating point format float2 and
        64-bit int's)
C:      sizeof (int)      = 8; sizeof (float) = 8
FORTRAN: sizeof (INTEGER) = 8; sizeof (REAL)  = 8)
tflops    (Intel TFLOPS)
UXPM      (UXP/M. Untested.)
uxpv      (uxp/v. Untested.)

```

Others may be recognized.

Special notes:

For SGI (--with-arch=IRIX) multiprocessors running the ch_p4 device, use -comm=ch_p4 to disable the use of the shared-memory p4 communication device, and -comm=shared to enable the shared-memory p4 communication device. The default is to enable the shared-memory communication device.

E Mpirun Usage

The options for mpirun as shown by mpirun -help, are (note that not all options are supported by all devices). Depending on the specific device, the output of mpirun -help may differ; the following is for the globus2 device.

```
mpirun [mpirun_options...] <progname> [options...]
```

```

mpirun_options:
  -arch <architecture>
        specify the architecture (must have matching machines.<arch>
        file in /usr/local/mpich/bin/machines) if using the execer
  -h      This help
  -machine <machine name>
        use startup procedure for <machine name>

```

`mpirun [mpirun_options...] <progname> [options...]`

`mpirun_options:`

- `-arch <architecture>`
 - specify the architecture (must have matching machines.<arch> file in /usr/local/mpich/bin/machines) if using the execer
- `-h` This help
- `-machine <machine name>`
 - use startup procedure for <machine name>
 - Currently supported:
 - paragon
 - p4
 - sp1
 - ibmbsp
 - anlsp
 - sgi_mp
 - ipsc860
 - inteldelta
 - cray_t3d
 - execer
 - smp
 - symm_ptx
- `-machinefile <machine-file name>`
 - Take the list of possible machines to run on from the file <machine-file name>. This is a list of all available machines; use `-np <np>` to request a specific number of machines.
- `-np <np>`
 - specify the number of processors to run on
- `-nodes <nodes>`
 - specify the number of nodes to run on (for SMP systems, currently only ch_mpl device supports this)
- `-nolocal`
 - don't run on the local machine (only works for ch_p4 jobs)
- `-all-cpus, -allcpus`
 - Use all available CPUs on all the nodes.
- `-all-local`
 - Run all processes on the master node.
- `-exclude <list>`
 - Exclude nodes in a colon delimited list.
- `-map <list>`
 - Use the colon delimited list to specify which rank runs on which nodes.
- `-stdin filename`
 - Use filename as the standard input for the program. This is needed for programs that must be run as batch jobs, such as some IBM SP systems and Intel Paragons using NQS (see `-paragontype` below).
 - use
 - `-stdin /dev/null`
 - if there is no input and you intend to run the program in the background. An alternate is to redirect standard input from /dev/null, as in
 - `mpirun -np 4 a.out < /dev/null`
- `-t` Testing - do not actually run, just print what would be

executed

-v Verbose - throw in some comments

-dbg The option '-dbg' may be used to select a debugger. For example, -dbg=gdb invokes the mpirun_dbg.gdb script located in the 'mpich/bin' directory. This script captures the correct arguments, invokes the gdb debugger, and starts the first process under gdb where possible. There are 4 debugger scripts; gdb, xgdb, ddd, totalview. These may need to be edited depending on your system. There is another debugger script for dbx, but this one will always need to be edited as the debugger commands for dbx varies between versions. You can also use this option to call another debugger; for example, -dbg=mydebug. All you need to do is write a script file, 'mpirun_dbg.mydebug', which follows the format of the included debugger scripts, and place it in the mpich/bin directory.

-ksq Keep the send queue. This is useful if you expect later to attach totalview to the running (or deadlocked) job, and want to see the send queues. (Normally they are not maintained in a way which is visible to the debugger).

Options for the globus2 device:

With the exception of -h, these are the only mpirun options supported by the globus device.

-machinefile <machine-file name>
Take the list of possible machines to run on from the file <machine-file name>

-np <np>
specify the number of processors to run on

-dumprsl - display the RSL string that would have been used to submit the job. using this option does not run the job.

-globusrsl <globus-rsl-file name>
<globus-rsl-file name> must contain a Globus RSL string. When using this option all other mpirun options are ignored.

Special Options for Batch Environments:

-mvhome Move the executable to the home directory. This is needed when all file systems are not cross-mounted. Currently only used by anlspdx

-mvback files
Move the indicated files back to the current directory. Needed only when using -mvhome; has no effect otherwise.

-maxtime min
Maximum job run time in minutes. Currently used only by anlspdx. Default value is \$max_time minutes.

-nopoll Do not use a polling-mode communication.

Available only on IBM SPs.

Special Options for IBM SP2:

`-cac name`

CAC for ANL scheduler. Currently used only by `anlspix`.

If not provided will choose some valid CAC.

On exit, `mpirun` returns a status of zero unless `mpirun` detected a problem, in which case it returns a non-zero status.

When using the `ch_p4` device, multiple architectures may be handled by giving multiple `-arch` and `-np` arguments. For example, to run a program on 2 sun4s and 3 rs6000s, with the local machine being a sun4, use

```
mpirun -arch sun4 -np 2 -arch rs6000 -np 3 program
```

This assumes that `program` will run on both architectures. If different executables are needed, the string `'%a'` will be replaced with the arch name. For example, if the programs are `program.sun4` and `program.rs6000`, then the command is

```
mpirun -arch sun4 -np 2 -arch rs6000 -np 3 program.%a
```

If instead the executables are in different directories; for example, `‘/tmp/me/sun4’` and `‘/tmp/me/rs6000’`, then the command is

```
mpirun -arch sun4 -np 2 -arch rs6000 -np 3 /tmp/me/%a/program
```

It is important to specify the architecture with `-arch` *before* specifying the number of processors. Also, the *first* `arch` command must refer to the processor on which the job will be started. Specifically, if `-nolocal` is *not* specified, then the first `-arch` must refer to the processor from which `mpirun` is running.

F Deprecated Features

During the development of MPICH, various features were developed for the installation and use of MPICH. Some of these have been superseded by newer features that are described above. This section archives the documentation on the deprecated features.

F.1 Getting Tcl, Tk, and wish

These software packages are available by anonymous ftp from `ftp.scrip.tics.com/pub/tcl/tcl\protect\unhbox\voidb@x\kern.06em\vbox{\hrulewidth.3em}old`. They are needed only for the `upshot` and `nupshot` programs; you do not need them in order to

install MPICH. If you cannot find them at `ftp.scriptics.com`, copies of Tcl and Tk are available at `ftp://ftp.mcs.anl.gov/mpi/tcltk`.

You should get `tcl7.3.tar.Z` and `tk3.6.tar.Z` (and patch `tk3.6p1.patch`). Later versions of both Tcl and Tk are incompatible with these and do not work with `nupshot`. The `upshot` program has been modified to work with either Tk 3.6 or Tk 4.0. `Upshot` may work with later versions, but we are no longer tracking the changes to Tcl/Tk.

It is necessary that the `wish` program be accessible to users; the other parts of Tcl and Tk do not need to be installed (but make sure that everything that `wish` itself needs is installed).

To build Tcl and Tk, we recommend the following approach:

1. Fetch the compressed tar files and the patch file into an empty directory, preferably in a local (not NFS) file system such as `‘/tmp’` (but make sure that you have enough space in that file system; xxx should be adequate).

2. Unpack the tar files:

```
gunzip -c tcl7.3.tar.Z | tar xf -
gunzip -c tk3.6.tar.Z | tar xf -
```

3. Apply the patch to Tk:

```
cd tk3.6
patch -p 1 < ../tk3.6p1patch
cd ..
```

(Note that the instructions say to use `patch -p`; newer versions of patch require an argument and the correct value in this case is one; other versions of patch will want `-p1` (no space between p and the one).)

4. Configure Tcl. Pick an installation directory that clearly indicates the Tcl and Tk versions. For example, to build Tcl to install into `‘/usr/local/tcl73tk36’`, use

```
cd tcl7.6
./configure -prefix=/usr/local/tcl73tk36
```

5. Build and install Tcl. Before you execute the `make install` step, make sure that the directory specified in the `-prefix` argument to `configure` exists.

```
mkdir /usr/local/tcl73tk36
make
make install
```

6. Configure, build, and install Tk. Use the same installation directory for Tk that you used for Tcl:

```
cd ../tcl7.6
./configure --prefix=/usr/local/tcl73tk36
make
make install
```

This will provide you with a Tcl and Tk installation that can be used with the Tcl and Tk tools provided with MPICH. If you have installed these into a non-standard location (such as the one used above), you can set the environment variable `TCL73TK36_DIR` to the location used as the prefix in the configure commands:

```
setenv TCL73TK36_DIR /usr/local/tcl73tk36
```

This will allow MPICH to find these versions of Tcl and Tk. We no longer use or recommend Tcl/Tk because of the lack of compatibility between versions of Tcl/Tk.

F.2 Obsolete Systems

To configure for the Intel Paragon, use

```
configure --with-device=ch_nx --with-arch=paragon
```

Two troubleshooting tips for the Paragon are

1. **Q:** I got the following messages when I tried to build on the Paragon:

```
PGC-W-0115-Duplicate standard type (init.c: 576)
PGC/Paragon Paragon Rel R5.0: compilation completed with warnings
PGC-W-0115-Duplicate standard type (init.c: 576)
PGC/Paragon Paragon Rel R5.0: compilation completed with warnings
```

A: This is because the compiler doesn't handle `long long` but doesn't reject it either. It causes no harm.

2. **Q:** I get errors compiling or running Fortran programs.

A:

Fortran programs will need to use a *absolute* path for the `'mpif.h'` include file, due to a bug in the `if77` compiler (it searches include directories in the wrong order).

A troubleshooting tip for the older Intel i860 system is

1. **Q:** The link test fails on an Intel i860 with

```
icc -o overtake overtake.o test.o -L/mpich/lib/intelnx/ -lmpi -lnode
/usr/ipsc/XDEV/i860/bin/ld860: Error: undefined symbol '_MPI_Keyval_create'
/usr/ipsc/XDEV/i860/bin/ld860: Fatal: no output file created
```

A: You are probably building MPICH on an old 386 running System V release 2. This version of Unix has very severe limitations on the length of filenames (more severe than we are willing to cater to). The specific problem here is that the name of the file ‘`mpich/src/context/keyval_create.c`’ is too long for this system, and was not properly archived. Your best bet is to build MPICH on a different, more modern system (for example, a Sun running SunOS or Solaris).

F.3 More detailed control over compiling and linking

Much of MPICH’s portability is handled through the careful construction of system-dependant Makefiles by the `configure` program. This is fine for installing MPICH, but what can you do when you are building a new application? For simple applications, the `mpicc` and `mpif77` commands may be the simplest way to build a new application. For more complex codes, we recommend taking a sample ‘`Makefile.in`’ file, for example, in ‘`mpich/examples/test/pt2pt`’. Modify those parts that are relevant, such as the `EXECS` and specific program targets. To create a ‘`Makefile`’, just execute

```
mpireconfig Makefile
```

(`mpireconfig` is in the same directory as `mpirun`). This generates a new ‘`Makefile`’ from ‘`Makefile.in`’, with the correct parameters for the MPICH that was installed.

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