Message Passing Interface (MPI)

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What is MPI?

A message-passing library specification for explicitly programming message-passing systems.

- ► Fully featured
 - point-to-point, collective, and one-sided communications
 - I/O routines and profiling interface
- ► Multiple language bindings
 - C, C++ and Fortran
- Available on many platforms
 - Production grade implementations on supercomputers
 - Open source implementations (e.g. Open MPI) on Linux and Windows
- ► Has been stable for many years:
 - MPI 2.0 was released in 1996, and it lasted 16 years
 - MPI 3.0 was released in Sept. 2012 (Manual 852 pages)

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Features of MPI

- Point-to-Point Communications
 - Structured buffers and derived data types, heterogeneity
 - many modes: blocking vs non-blocking vs synchronous, ready vs buffered
- Collective Communications
 - Both built-in and user-defined collective operations
 - Large number of data movement routines
 - Subgroups defined directly or by topology
 - Built-in support for grids and graphs
- One-Sided Communications
 - Allow remote memory access
- Others
 - Message security, thread safety
 - Profiling hooks, error control

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MPI for Beginners

One need not master all parts of MPI to use it. These six functions allow you to write many programs:

- ► MPI_Init starting MPI
- ► MPI_Finalize exiting MPI
- ▶ MPI_Comm_size the number of processes
- ▶ MPI_Comm_rank the id of *this* process
- ▶ MPI_Send sending a message
- ▶ MPI_Recv receiving a message

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"Hello world!" in MPI

C Version:

```
#include "mpi.h"
int main(int argc, char **argv) {
   MPI_Init(&argc, &argv);
   printf("Hello world!\n");
   MPI_Finalize();
   return 0;
}
```

► C++ Version:

```
#include "mpi++.h"
int main(int argc, char **argv) {
   MPI::Init(argc, argv);
   cout << "Hello world!" << endl;
   MPI::Finalize();
   return 0;
}</pre>
```

MPI_Init() and MPI_Finalize() must be included in every MPI program.

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"Hello world!" in MPI (Version 2)

```
#include "mpi.h"
int main(int argc, char **argv) {
   int rank, size;
   MPI_Init(&argc, &argv);
   MPI_Comm_rank(MPI_COMM_WORLD, &rank);
   MPI_Comm_size(MPI_COMM_WORLD, &size);
   printf("Hello world! I'm %d of %d\n", rank, size);
   MPI_Finalize();
   return 0;
}
```

Communication in MPI takes place with respect to communicators.

- ► MPI_COMM_WORLD is the default communicator, it contains all MPI processes. User can define other communicators.
- ▶ MPI_Init() and MPI_Finalize() must be called by *all* processes.

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Basic Send/Receive

- ▶ buf initial address of send/receive buffer
- ▶ count, type number of and type of elements to be sent/received
- dest, src rank of destination and source processes; a special value for src is MPI_ANY_SOURCE
- tag message tag
- ▶ comm communicator
- ▶ st status of the receive command; its info includes MPI_SOURCE, MPI_TAG, MPI_ERROR, and message length

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A Simple Send/Receive Program

```
#include <mpi.h>
int main(int argc, char **argv) {
 int i, rank, size, dest, to, src, from;
 int count, tag, st_count, st_src, st_tag;
 double data[100];
 MPI_Status st;
 MPI_Init(&argc, &argv);
 MPI_Comm_rank(MPI_COMM_WORLD, &rank);
 MPI_Comm_size(MPI_COMM_WORLD, &size);
 printf("Process %d of %d is alive\n", rank, size);
 src = 0; dest = size - 1;
 if (rank == src) {
   to = dest;
   tag = 2001;
   count = 100;
   for (i = 0; i < 100; i++)
     data[i] = i;
   MPI_Send(data, count, MPI_DOUBLE, to, tag, MPI_COMM_WORLD);
 }
```

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A Simple Send/Receive Program (cont.)

```
else if (rank == dest) {
    from = MPI_ANY_SOURCE;
    tag = MPI_ANY_TAG;
    count = 100;
   MPI_Recv(data, count, MPI_DOUBLE, from, tag, MPI_COMM_WORLD, &st);
   MPI_Get_count(&st, MPI_DOUBLE, &st_count);
   st_src = status.MPI_SOURCE;
   st_tag = status.MPI_TAG;
   printf("Status info: source = %d, tag = %d, count = %d\n",
           st_src, st_tag, st_count);
   printf(" %d received: ", rank);
   for (i = 0; i < st_count; i++)
      printf("%lf ", data[i]);
    printf("\n");
 MPI_Finalize();
  return 0;
}
```

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Non-Blocking Send and Receive

Both routines create and allocate a *request object* and return a pointer to it in the req variable.

This object is useful for future testing and waiting for the finish of the non-blocking operation:

```
int MPI_Test(MPI_Request *reg, int *flag, MPI_Status *st)
int MPI_Wait(MPI_Request *reg, MPI_Status *st)
```

▶ If the value of flag is true, the operation has completed.

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Combined Send/Receive

The routine performs both a send and a receive. Advantages over pair of send and receive:

- avoids deadlock
- does not require data buffering

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Other Send Routines

Synchronous Sends — Won't complete until the corresponding receive has been posted:

```
int MPI_Ssend(...)
int MPI_Issend(...)
```

▶ Ready Sends — Assume the corresponding receive has been posted; otherwise they are errorous:

```
int MPI_Rsend(...)
int MPI_Irsend(...)
```

▶ Buffered Sends — If the corresponding receive has not been posted, the system must buffer the data and return; it's the user's responsibility to allocate the buffer:

```
int MPI_Bsend(...)
int MPI_Ibsend(...)
```

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Collective Communications

MPI Collective communications are coordinated among a group of processes, as specified by communicator.

- Message tags are not used
- ▶ All collective operations are blocking
- ► All processes in the communicator group must call the collective operation

Categories:

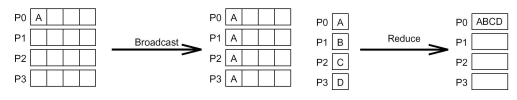
- ► One-to-All Broadcast
- All-to-One Reduction
- Scatter and Gather
- ► Prefix Scan
- ► All-to-All Broadcast & Reduction

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Basic Broadcast and Reduce



▶ The reduce routine take can take the following built-in collective operators:

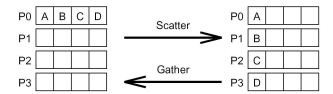
```
MPI_SUM MPI_MAX MPI_LAND MPI_BAND MPI_LXOR MPI_MAXLOC
MPI_PROD MPI_MIN MPI_LOR MPI_BOR MPI_BXOR MPI_MINLOC
```

▶ It can also take user-defined reduction functions.

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Scatter and Gather

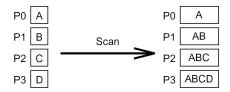


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Prefix Scan

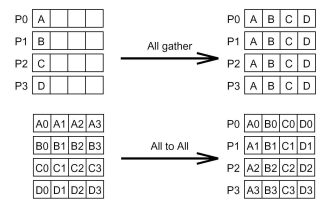


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All-Version and V-Version Routines

▶ All-versions deliver results to all participating processes.



MPI_ALLGATHER MPI_ALLREDUCE MPI_ALLTOALL

▶ V-versions allow the chunks to have different sizes.

MPI_ALLGATHERV MPI_GATHERV MPI_ALLTOALLV

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Example: Collective Communication

```
#include <mpi.h>
int main(int argc, char **argv) {
 int rank, size, myn, i, N;
 double *vector, *myvec, sum, mysum, total;
 MPI_Init(&argc, &argv);
 MPI_Comm_rank(MPI_COMM_WORLD, &rank);
 MPI_Comm_size(MPI_COMM_WORLD, &size);
 /* In the root process read the vector length, initialize
    the vector and determine the sub-vector sizes */
  if (rank == 0) {
   printf("Enter the vector length : ");
   scanf("%d", &N);
    vector = (double *)malloc(sizeof(double) * N);
   for (i = 0, sum = 0; i < N; i++)
     vector[i] = 1.0;
   myn = N / size;
 MPI_Bcast(&myn, 1, MPI_INT, 0, MPI_COMM_WORLD);
```

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Example: Collective Communication (cont.)

```
myvec = (double *)malloc(sizeof(double)*myn);
MPI_Scatter(vector, myn, MPI_DOUBLE, myvec, myn, MPI_DOUBLE,
            0, MPI_COMM_WORLD);
/* Find the local sum, then the global sum of the vectors */
for (i = 0, mysum = 0; i < myn; i++)
 mysum += myvec[i];
MPI_Allreduce(&mysum, &total, 1, MPI_DOUBLE, MPI_SUM, MPI_COMM_WORLD);
/* Multiply the local part of the vector by the global sum */
for (i = 0; i < myn; i++)
 myvec[i] *= total;
MPI_Gather(myvec, myn, MPI_DOUBLE, vector, myn, MPI_DOUBLE,
           O, MPI_COMM_WORLD);
if (rank == 0)
 for (i = 0; i < N; i++)
    printf("[%d] %f\n", rank, vector[i]);
MPI_Finalize();
return 0;
```

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One-Sided Communications

(A.k.a. Remote memory access or RMA.)

- ▶ Allow one process to specify all communication parameters, both for the sending side and for the receiving side.
- ► De-couple communication of data and syncrhonization between sender and receiver.
- ▶ The semantics is *very* complicated.

Sample Routines:

- ► MPI_Put remote write
- ▶ MPI_Get remote read
- ▶ MPI_Accumulate remote update
- MPI_Compare_and_swap remote atomic swap

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One-Sided Communications (cont.)

▶ One-sided communications can only access data in specific memory regions called "windows", which have to be explicitly created.

```
MPI_Win_create(...)
```

▶ One-sided communications can only occur during specific temporal intervals called "epochs", which are bracketed with synchronization calls.

```
MPI_Win_lock(...)
... // RMA operation here
MPI_Win_unlock(...)
```

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Example: One-Sided Communication

Process A:

```
int n;
MPI_WIN nwin;

MPI_Win_create(&n, sizeof(int), 1, MPI_INFO_NULL, MPI_COMM_WORLD, &nwin);
...
n = 1000; /* local update */
MPI_Barrier(MPI_COMM_WORLD);
...
```

Process B:

```
int n;
MPI_WIN nwin;

MPI_Win_create(MPI_BOTTOM, 0, 1, MPI_INFO_NULL, MPI_COMM_WORLD, &nwin);
...
MPI_Barrier(MPI_COMM_WORLD);
MPI_Win_lock(MPI_LOCK_EXCLUSIVE, 0, 0, nwin);
MPI_Get(&n, 1, MPI_INT, 0, 0, 1, MPI_INT, nwin); /* remote read */
MPI_Win_unlock(MPI_LOCK_EXCLUSIVE, 0, 0, nwin);
```

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File I/O

- ▶ Both routines are collective routines:
 - Input parameters (except info) must have the same values;
 - MPI_File_close implies a MPi_File_sync.
- ► However, it's possible to open a file to just one process by using MPI_COMM_SELF as comm's value.
- ► File access modes include the usual choices, e.g., read-only, read-write, append, etc.
- ► File Info is for advanced uses (i.e. passing optimization hints). For normal cases, just use MPI_INFO_NULL.

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Example: Reading From a File

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Example: Writing to a File

```
int main(int argc, char *argv[])
 int cnt=4, buf[4], i, rank;
 char fname[10];
 MPI_File fh;
 MPI_Status st;
 MPI_Init(&argc,&argv);
 MPI_Comm_rank(MPI_COMM_WORLD, &rank);
 for (i=0; i<cnt; i++) buf[i] = rank*100 + i;
  sprintf(fname, "%s.%d", "test", rank);
 MPI_File_open(MPI_COMM_SELF, fname, MPI_MODE_CREATE|MPI_MODE_RDWR,
               MPI_INFO_NULL, &fh);
 MPI_File_set_view(fh, 0, MPI_INT, MPI_INT, "native", MPI_INFO_NULL);
 MPI_File_write(fh, buf, cnt, MPI_INT, &st);
 MPI_File_close(&fh);
 MPI_Finalize();
 return 0;
```

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File View

This routine sets individual process's view of the data in the file.

- disp displacement. Specifies where within the file to start access. Useful for reading different sections to different processes.
- etype elementary datatype.
- ▶ ftype file type. For advanced uses. Specifies gaps between desirable data items. For simple cases, just use the same value as etype.
- ▶ datarep data representation. Normally just use "native".

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Open MPI

An open-source implementation of MPI.

- Comprehensive support many platforms (including GPUs)
- ► Flexible lots of user-level configuration/tuning controls
- ► High-Performance used by many TOP500 supercomputers

(The name 'Open MPI' can be shortend to OMPI, but not to OpenMPI.)

The steps to use Open MPI:

- Creating a hostfile
- Compiling a program
- Running a program

Optionally, one can also create an appfile.

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Open MPI: Creating a Hostfile

A hostfile contains a list of computer names.

Example:

```
linux> cat myhosts
# a line starting with # is a comment
african
chatham
chinstrap
```

- ► The MPI program must be accessible by the same pathname on all host computers.
- ► The environment variable OMPI_MCA_orte_default_hostfile can be set to point to a default hostfile.
- ▶ It's a good idea to create multiple host files for different number of hosts: e.g. mpihost2, mpihost4, mpihost8, etc.

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Open MPI: Compiling a Program

The command: mpicc

Example:

```
linux> mpicc -o ring ring.c
```

mpicc is just a gcc wrapper:

```
linux> mpicc --showme
gcc -I/usr/lib/openmpi/include -I/usr/lib/openmpi/include/openmpi
-pthread -L/usr/lib/openmpi/lib -lmpi -lopen-rte -lopen-pal -ldl
-Wl,--export-dynamic -lnsl -lutil -lm -ldl
```

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Open MPI: Running a Program

The command:

```
mpirun -host <hostnames> -n <#process> program>
mpirun -hostfile <hostfile> -n <#process> program>
```

Example:

```
linux> mpirun -n 4 ring
linux> mpirun -host african -n 4 ring
linux> mpirun -host african, chatham -n 4 ring
linux> mpirun -hostfile host2 -n 4 ring
linux> mpirun -hostfile host4 -n 4 ring
```

- ▶ If there is no host or hostfile specified, the program will run on the console computer.
- ▶ The number of processes do not need to match the number of hosts.
- ▶ If a default hostfile exists, the -hostfile switch can be omitted.

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Open MPI: Running a Program (cont.)

Open MPI also supports master/slave programs:

Example:

```
linux> mpirun -hostfile myhosts -n 1 master -n 4 slave
```

This command will run one copy of 'master' and four copies of 'slave'.

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Creating an Appfile

mpirun command-line parameters can be saved in an appfile; and a program can be executed from an appfile.

Example:

```
linux> cat myapp
# execute 4 copies of 'ring' on 2 hosts
-host african,chatham -n 4 ring
linux> mpirun -app myapp
```