Distributed Parallel Programming with MPI

Ramses van Zon

PHY1610 Winter 2024



Ramses van Zon

Distributed Parallel Programming with MPI

Issues with shared memory programming

- Parallel tasks are run by threads.
- All threads live on the same node and share the memory.
- Limited to the resources of a single node.
- Creation and deletion of threads can cause overhead.
- Can lead to bugs like race conditions.



Solution: distributed memory programming

- Parallel tasks are processes.
- Each process has only its own, private memory.
- Processes need not be on the same node.
- You can scale up the size of your system to as many resources as you have.
- Harder to create race condition bugs, but now you get new bugs like dead-lock.
- Must explicitly code in the communication between processes: Message Passing Interface aka MPI



MPI Intro



Message Passing Interface (MPI)

What is it?

- An open standard library interface for message passing, ratified by the MPI Forum
- Version: 1.0 (1994), 1.1 (1995), 1.2 (1997), 1.3 (2008)
- Version: 2.0 (1997), 2.1 (2008), 2.2 (2009)
- Version: 3.0 (2012), 3.1 (2015)
- Version: 4.0 (2021), 4.1 (2023)

MPI Implementations (Teach)

- OpenMPI www.open-mpi.org
 - \$ module load gcc/13 openmpi/5
 - \$ module load intel/2023u1 openmpi/5

Currently these give you OpenMPI version 5.0.0.

- MPICH www.mpich.org (MPICH, MVAPICH2, IntelMPI)
 - \$ module load gcc/13 intelmpi/2023u1
 - \$ module load intel/2023u1 intelmpi/2023u1

MPI is a Library for Message-Passing

Library:

- Not built in to compiler.
- Function calls that can be made from any compiler, many languages.
- Just link to it.
- Compiler wrappers: mpicc, mpif90, mpicxx
- Runtime wrappers: mpiexec/mpirun

```
#include <iostream>
#include <string>
#include <mpi.h>
using std::cout;
using std::to string;
int main(int argc, char **argv)
{
    int rank. size:
    MPI_Init(&argc, &argv);
    MPI Comm size(MPI COMM WORLD, &size):
    MPI Comm rank(MPI COMM WORLD, &rank):
    cout << "Hello from task " +</pre>
            to string(rank) + " of " +
            to_string(size) + "\n";
```

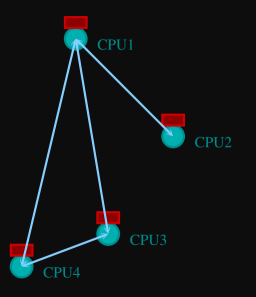
```
MPI_Finalize();
```



}

PHY1610 Winter 2024

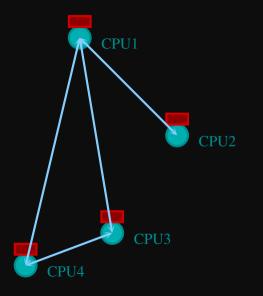
MPI is a Library for Message Passing



- Communication/coordination between tasks done by sending and receiving messages.
- Each message involves a function call from each of the programs.



MPI is a Library for Message Passing



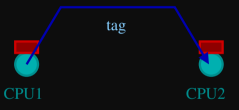
Three basic sets of functionality:

- Pairwise communications via messages;
- Collective operations via messages;
- Efficient routines for getting data from memory into messages and vice versa.



Messages

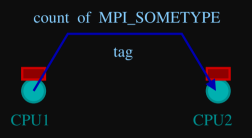
count of MPI_SOMETYPE



- Messages have a sender and a receiver.
- When you are sending a message, you don't need to specify the sender (it is the current processor).
- A sent message has to be actively received by the receiving process



Messages



- MPI messages are a string of length count all of some fixed MPI type.
- MPI types exist for characters, integers, floating point numbers, etc.
- An arbitrary non-negative integer tag is also included – helps keep things straight if lots of messages are sent.



Size of MPI Library

- Many, many functions (>200).
- Not nearly so many concepts.
- We'll get started with just 10-12, use more as needed.

MPI_Init()
MPI_Comm_size()
MPI_Comm_rank()
MPI_Ssend()
MPI_Recv()
MPI_Finalize()



Example: Hello World

```
#include <iostream>
#include <string>
#include <mpi.h>
using std::cout;
using std::to string;
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);
   cout<< "Hello from task" + to_string(rank) +</pre>
          " of " + to string(size) + " world\n":
   MPI_Finalize();
```



Example: Hello World

Compile with MPI

MPI provides compiler wrappers

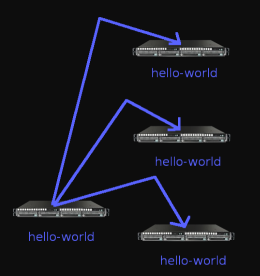
- mpicc
- mpicxx
- mpif90

that set all the -I, -L, -1, etc. options properly for the base compiler.

```
$ git clone /scinet/course/phy1610/mpi
$ cd mpi
$ module load gcc/13 openmpi/5 rarray
$ mpicxx -02 -std=c++17 -o mpi-hello-world mpi-hello-world.cc # or: 'make mpi-hello-world'
$ mpirun -n 16 ./mpi-hello-world
```



What mpirun Does



- Launches *n* processes, assigns each an MPI rank and starts the program.
- Usually, the processes run the same executable, therefore **each process runs the exact same code**.
- For multinode runs, has a list of nodes, and logs in (effectively) to each node, where it launches the program.



Ramses van Zon

Distributed Parallel Programming with MPI

Number of Processes

- Number of processes to use is almost always equal to the number of processors.
- But not necessarily. (memory-bound, hybrid, i/o bound)
- On a Teach debugjob, what happens when you run this?

\$ debugjob -n 16 \$ mpirun -n 16 ./mpi-hello-world Hello from task 5 of 16 Hello from task 1 of 16 Hello from task 0 of 16 Hello from task 4 of 16 Hello from task 6 of 16 Hello from task 13 of 16 Hello from task 10 of 16 Hello from task 12 of 16 Hello from task 7 of 16 Hello from task 9 of 16 Hello from task 14 of 16 Hello from task 11 of 16 Hello from task 2 of 16 Hello from task 3 of 16 Hello from task 15 of 16 Hello from task 8 of 16



mpirun runs any program

- mpirun will start its process-launching procedure for any program.
- Sets variables somehow that mpi programs recognize so that they know which process they are.

E.g., try this:

\$ hostname
teach31.scinet.local
\$ mpirun -n 4 hostname

teach31.scinet.local
teach31.scinet.local
teach31.scinet.local
teach31.scinet.local

\$ ls \$ mpirun -n 4 ls



Example: Hello World

\$ mpirun -n 4 ./mpi-hello-world Hello from task 2 of 4 world Hello from task 1 of 4 world Hello from task 0 of 4 world Hello from task 3 of 4 world

\$ mpirun --output TAG-DETAILED -n 4 ./mpi-hello-world [1,1][teach31:21851]<stdout>: Hello from task 1 of 4 [1,0][teach31:21850]<stdout>: Hello from task 0 of 4 [1,2][teach31:21852]<stdout>: Hello from task 2 of 4 [1,3][teach31:21853]<stdout>: Hello from task 3 of 4

The --output flag is specific for the OpenMPI implementation of MPI.



MPI Basics



MPI Basics

```
#include <iostream>
#include <string>
#include <mpi.h>
using std::cout;
using std::to_string;
```

```
int main(int argc, char **argv)
{
```

```
int rank, size;
```

```
MPI_Init(&argc, &argv);
```

```
MPI_Finalize();
```

Basic MPI Components

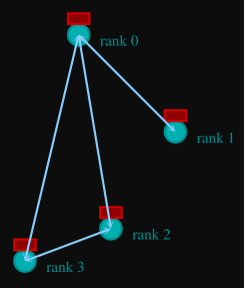
- #include <mpi.h> MPI library definitions
- MPI_Init(&argc,&argv)
 MPI Intialization, must come first
- MPI_Finalize()
 Finalizes MPI, must come last
- Formally, MPI routines return an error code. But in fact, MPI applications by default abort when there is an error.

Communicator Components

- A communicator is a handle to a group of processes that can communicate.
- MPI_Comm_rank(MPI_COMM_WORL
- MPI_Comm_size(MPI_COMM_WORI



Communicators



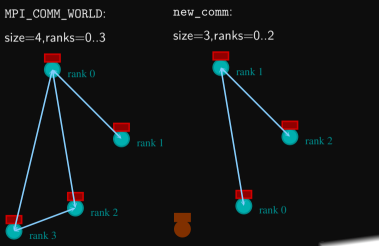
- MPI groups processes into communicators.
- Each communicator has some size number of tasks.
- Every task has a rank 0..size-1
- Every task in your program belongs to MPI_COMM_WORLD.

 $\begin{array}{l} \texttt{MPI_COMM_WORLD:} \\ \texttt{size} = \texttt{4}, \ \texttt{ranks} = \texttt{0..3} \end{array}$



Communicators

- One can create one's own communicators over the same tasks.
- May break the tasks up into subgroups.
- May just re-order them for some reason.





PHY1610 Winter 2024 21 / 40

MPI Basics - Communicator Components

• MPI_COMM_WORLD:

Global Communicator

- MPI_Comm_rank(MPI_COMM_WORLD,&rank)
 Get current tasks rank
- MPI_Comm_size(MPI_COMM_WORLD,&size)

Get communicator size



MPI = Rank and Size

Rank and Size are much more important in MPI than in OpenMP

- In OpenMP, the compiler assigns jobs to each thread; you do not need to know which one is which (usually).
- In MPI, all proceses run the same code.
- In MPI, processes determine amongst themselves which piece of puzzle to work on, based on their **rank**, then communicate with appropriate others.









Distributed Parallel Programming with MPI

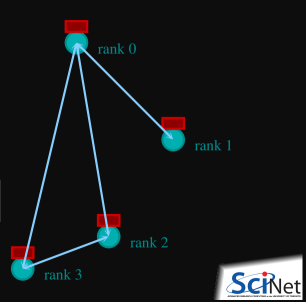
MPI = Communication

Explicit Communication between Tasks

- In OpenMP, threads can communicate using the memory.
- In MPI, a process which needs data of another process needs to communicate with that process by passing messages.

MPI_Ssend(...)

MPI_Recv(...)



Distributed Parallel Programming with MPI

MPI: Send & Receive

MPI_Ssend(sendptr, count, MPI_TYPE, destination,tag, Communicator);

MPI_Recv(recvptr, count, MPI_TYPE, source, tag, Communicator, MPI_status)

- sendptr/recvptr: pointer to message
- count: number of elements in message
- MPI_TYPE: one of MPI_DOUBLE, MPI_FLOAT, MPI_INT, MPI_CHAR, etc.
- destination/source: rank of sender/reciever
- tag: unique id for message pair
- Communicator: MPI_COMM_WORLD or user created
- status: receiver status (error, source, tag)

Note: MPI has a Fortran and C interface. We can use the C interface in C++ but will have to deal with pointers, i.e., we'll give arguments likes $\mathcal{O}(array[0])$ or array.data() instead of just array.



PHY1610 Winter 2024

25 / 40

MPI: Send & Receive

```
#include <iostream>
#include <string>
#include <mpi.h>
using std::cout:
using std::to_string;
int main(int argc, char **argv) {
    int rank. size:
    int tag = 1:
    double msgsent, msgrcvd;
    MPI Status rstatus:
    MPI Init(&argc, &argv):
    MPI Comm rank(MPI COMM WORLD, &rank):
    MPI Comm size(MPI COMM WORLD, &size):
    msgsent = 111.:
    msgrcvd = -999.:
    if (rank == 0) {
        MPI_Ssend(&msgsent, 1, MPI_DOUBLE, 1, tag, MPI_COMM_WORLD);
        cout << "Sent " + to_string(msgsent) + " from " + to_string(rank) + "\n";</pre>
    if (rank == 1) {
        MPI_Recv(&msgrcvd, 1, MPI_DOUBLE, 0, tag, MPI_COMM_WORLD, &rstatus);
        cout << "Received " + to string(msgrcvd) + " on " + to string(rank) + "\n":
    3
    MPI Finalize():
                                         Distributed Parallel Programming with MPI
           Ramses van Zon
```

MPI: Send & Receive

\$ make firstmessage \$ mpirun -n 2 ./firstmessage Send 111.000000 from 0 Received 111.000000 on 1



MPI Communication Patterns

Send a message to the right:





Specials

Special Source/Destination MPI_PROC_NULL

MPI_PROC_NULL basically ignores the relevant operation; can lead to cleaner code.

Special Source MPI_ANY_SOURCE

MPI_ANY_SOURCE is a wildcard; matches any source when receiving.

Special Status MPI_STATUS_IGNORE

Use MPI_STATUS_IGNORE if you do not want to capture the status in a receive.



Deadlocks



Deadlocks are a classic parallel bug

- In this explicit message passing model, it is possible to completely freeze the application.
- This can happen when a process is sending a message, but no process is or will ever be ready to receive it.
- This is called deadlock
- To see how that could happen, let's look at an example.



MPI: Send Right, Receive Left

```
#include <iostream>
#include <string>
#include <mpi.h>
using std::cout;
using std::to_string;
int main(int argc, char **argv)
    int
              rank, size, left, right, tag = 1;
    double
                msgsent, msgrcvd:
    MPI Init(&argc, &argv):
    MPI Comm rank(MPI COMM WORLD, &rank):
    MPI_Comm_size(MPI_COMM_WORLD, &size);
    left = rank - 1:
    if (left < 0) left = MPI PROC NULL:
    right = rank + 1:
    if (right >= size) right = MPI_PROC_NULL;
    msgsent = rank*rank:
    msgrcvd = -999.:
    MPI_Ssend(&msgsent, 1, MPI_DOUBLE, right, tag, MPI_COMM_WORLD);
    MPI Recv(&msgrcvd, 1, MPI DOUBLE, left, tag, MPI COMM WORLD, MPI STATUS IGNORE);
    cout << to_string(rank) + ": Sent " + to_string(msgsent)</pre>
          + " and got " + to_string(msgrcvd) + "\n";
    MPI Finalize():
```

MPI: Send Right, Receive Left

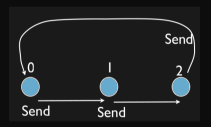
\$ make secondmessage \$ mpirun -n 3 ./secondmessage 2: Sent 4.000000 and got 1.000000 0: Sent 0.000000 and got -999.000000 1: Sent 1.000000 and got 0.000000 \$

\$ mpirun -n 6 ./secondmessage 4: Sent 16.000000 and got 9.000000 5: Sent 25.000000 and got 16.000000 0: Sent 0.000000 and got -999.000000 1: Sent 1.000000 and got 0.000000 2: Sent 4.000000 and got 1.000000 3: Sent 9.000000 and got 4.000000



MPI: Send Right, Receive Left with Periodic BCs

Periodic Boundary Conditions:





MPI: Send Right, Receive Left with Periodic BCs

```
...
left = rank - 1;
if (left < 0) left = size-1; // Periodic BC
right = rank + 1;
if (right >= size) right =0; // Periodic BC
msgsent = rank*rank;
msgrcvd = -999.;
...
```

\$ make thirdmessage
\$ mpirun -n 3 ./thirdmessage

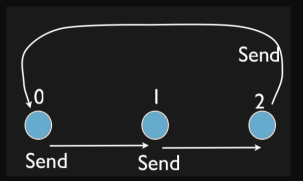
Program hangs!



Deadlock!

- A classic parallel bug.
- Occurs when a cycle of tasks are waiting for the others to finish.
- Whenever you see a closed cycle, you likely have (or risk) a deadlock.
- Here, all processes are waiting for the send to complete, but no one is receiving.

Sends and receives must be paired when sending

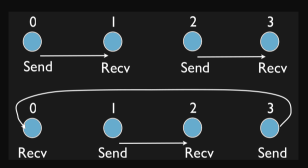




How do we fix the deadlock?

Without using new MPI routine, how do we fix the deadlock?

Even-odd solution



- First: evens send, odds receive
- Then: odds send, evens receive
- Will this work with an odd number of processes? How about 2? 1?



MPI: Send Right, Recv Left with Periodic BCs - fixed

```
if ((rank % 2) == 0) {
    MPI_Ssend(&msgsent, 1, MPI_DOUBLE, right, tag, MPI_COMM_WORLD);
    MPI_Recv(&msgrcvd, 1, MPI_DOUBLE, left, tag, MPI_COMM_WORLD, MPI_STATUS_IGNORE);
} else {
    MPI_Recv(&msgrcvd, 1, MPI_DOUBLE, left, tag, MPI_COMM_WORLD, MPI_STATUS_IGNORE);
    MPI_Ssend(&msgsent, 1, MPI_DOUBLE, right, tag, MPI_COMM_WORLD);
}
```

```
$ make fourthmessage
$ mpirun -n 5 ./fourthmessage
1: Sent 1.000000 and got 0.000000
2: Sent 4.000000 and got 1.000000
3: Sent 9.000000 and got 4.000000
4: Sent 16.000000 and got 9.000000
0: Sent 0.000000 and got 16.000000
```



MPI: Sendrecv

- A blocking send and receive built together.
- Lets them happen simultaneously.
- Can automatically pair send/recvs.
- Why 2 sets of tags/types/counts?



Send Right, Receive Left with Periodic BCs - Sendrecv

