



# Introduction to MPI Programming – Part 1





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### Outline

- Introduction what is MPI and why MPI
- MPI program structure
- Point-to-point communication





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### Memory system models

- Different ways of sharing data among processors
  - Distributed Memory
  - Shared Memory
  - Other memory models
    - Hybrid model
    - PGAS (Partitioned Global Address Space)









# Message Passing

- Context: distributed memory parallel computers
  - Each processor has its own memory space and cannot access the memory of other processors
  - Any data to be shared must be explicitly transferred from one to another









### Distributed memory model

- Each process has its own address space
  - Data is local to each process
- Data sharing achieved via explicit message passing (through network)



• Example: MPI (Message Passing Interface)









### Shared memory model

- All threads can access the global address space
- Data sharing achieved via writing to/reading from the same memory location



• Example: OpenMP









# Message Passing Interface

- MPI defines a standard API for message passing
  - What's in the standard
    - The syntax and semantics of a core set of functions
  - What's not in the standard
    - Implementation details
    - Runtime details (how many processes the code with run with etc.)
- MPI provides C/C++ and Fortran bindings











# Why MPI?

- Standardized
  - With efforts to keep it evolving (MPI 3.0 draft came out in 2010)
- Portability
  - MPI implementations are available on almost all platforms
- Scalability
  - In the sense that it is not limited by the number of processors that can access the same memory space
- Popularity
  - Many packages are based on MPI











### When NOT to use MPI

- Not suitable for small-scale loop level parallelization
  - Shared memory parallelism and accelerator are better









### **MPI Functions**

- Point-to-point communication functions
  - Message transfer from one process to another
- Collective communication functions
  - Message transfer involving all processes in a communicator
- Environment management functions
  - Initialization and termination
  - Process group and topology











### **MPI Program Structure**

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```
program hello
...
include "mpif.h"
integer :: nprocs,myid,ierr
...
call mpi_init(ierr)
...
call mpi_comm_size(mpi_comm_world,nprocs,ierr)
call mpi_comm_rank(mpi_comm_world,myid,ierr)
Write(*,'("There are",I3," processes")') nprocs
write(*,'("Process",I3," says Hello World!")') myid
...
call mpi_finalize(ierr)
...
```





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11





### **MPI Program Structure**

program hello								
 include "mpif.h"								
call mpi_init(ierr)								
call mpi_cc [lyan1@qb563 ex]\$ mpirun -np 4 ./a.out								
call mpi_cc There are 4 processes.								
Write(*,'("	There are 4 processes.							
write(*,'("	There are 4 processes.							
	There are 4 processes.							
call mpi_fi	Process 3 says Hello World!							
	Process 1 says Hello World!							
	Process 0 says Hello World!							
9	Process 2 says Hello World!							

#### Header file

Initialization

Computation and communication

**Termination** 





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#### C vs. Fortran

- Header file
  - C:mpi.h
  - Fortran:mpif.h
- Function names
  - C:MPI\_Xxx\_Yyyy
  - Fortran: mpi\_xxx\_yyyy (doesn't really matter)
- Error handles
  - C returns the error value, while Fortran passes it as an argument
    - C:int err = MPI\_Xxx(arg1,arg2,...,argN)
    - Fortran:call mpi\_xxx(arg1,arg2,...,argN,ierr)









# Initialization and Termination

- Initialization
  - Must be called before any other MPI calls
  - C: MPI\_Init()
  - Fortran: MPI\_INIT(ierr)
- Termination
  - Clean up data structures, terminate incomplete calls etc.
  - C:MPI\_Finalize()
  - Fortran: MPI\_FINALIZE(ierr)









# Communicators (1)

- A communicator is an identifier associated with a group of processes
  - Can be regarded as an ordered list of processes
  - Each process has a unique rank, which starts from 0 (root)
  - It is the context of MPI communicators and operations
    - When a function is called to send data to all processes, MPI needs to understand what "all" means









# Communicators (2)

- MPI\_COMM\_WORLD: the default communicator that contains all processes running the MPI program
- There can be many communicators
- A process can belong to multiple communicators
  - The rank is usually different











# **Getting Communicator Information**

- Get the rank of a communicator
  - C: MPI\_Comm\_Rank(MPI\_Comm comm, int
     \*rank)
  - Fortran: MPI\_COMM\_RANK ( COMM , RANK , ERR )
- Get the size in a communicator
  - C: MPI\_Comm\_Size(MPI\_Comm comm, int
     \*size)
  - Fortran: MPI\_COMM\_SIZE(COMM,SIZE,ERR)











### Compiling and Running MPI Programs

- Not a part of the standard
  - Could vary from platform to platform
  - Or even from implementation to implementation on the same platform
- On LONI Linux systems:
  - Compile
    - C:mpicc -o <executable name> <source file>
    - Fortran: mpif90 -o <executable name> <source file>
  - Run
    - mpirun -machinefile \$PBS\_NODEFILE -np <number of procs> <executable name> <input parameters>











### Exercise 1a: Process Color

- Write a MPI program where
  - Processes with odd rank print to screen "Process x has the color green"
  - Processes with even rank print to screen "Process x has the color red"









#### Exercise 1b: Laplace Solver version 0

- Goal: Distribute the work load among processes in 1-d manner
  - Find out the size of sub-matrix for each process
  - Let each process report which part of the domain it will work on, e.g. "Process x will process column (row) x through column (row) y."
    - Row-wise (C) or column-wise (Fortran)









### Point-to-point Communication

- Communication between a pair of processes, so two functions calls are required
  - The sending process calls the MPI\_SEND function
    - C:int MPI\_Send(void \*buf, int count, MPI\_Datatype dtype, int dest, int tag, MPI\_Comm comm);
    - Fortran: MPI\_SEND(BUF, COUNT, DTYPE, DEST, TAG, COMM, IERR)
  - The receiving process calls the MPI\_RECV function
    - C: int MPI\_Recv(void \*buf, int count, MPI\_Datatype dtype, int source, int tag, MPI\_Comm comm, MPI\_Status \*status);
    - Fortran: MPI\_RECV(BUF, COUNT, DTYPE, SOURCE, TAG, COMM, STATUS, IERR)
- The function arguments characterize the message being transferred









### **MPI** Message

- A MPI message consists of two parts
  - Message body
    - Buffer: starting location in memory for outgoing data (send) or incoming data (receive)
    - Data type: type of data to be sent or received
    - Count: number of items of type datatype to be sent or received
  - Message envelope
    - Destination (source): rank of the destination (source) of the message
    - Tag: what MPI uses to match messages between processes
    - Communicator
- The status argument contains information on the message that is received













### Example: Gathering Array Data



• Goal: gather some array data from each process and place it in the memory of the root process









### Example: Gathering Array Data

```
integer,allocatable :: array(:)
! Initialize MPI
call mpi_init(ierr)
call mpi_comm_size(mpi_comm_world,nprocs,ierr)
call mpi_comm_rank(mpi_comm_world,myid,ierr)
! Initialize the array
allocate(array(2*nprocs))
array(1)=2*myid
array(2)=2*myid+1
! Send data to the root process
if (myid.eq.0) then
  do i=1,nprocs-1
    call mpi_recv(array(2*i+1),2,mpi_integer,i,i,status,ierr)
  enddo
  write(*,*) "The content of the array:"
  write(*,*) array
else
  call mpi_send(array,2,mpi_integer,0,myid,ierr)
endif
```







### Example: Gathering Array Data

	 integer,a: <b>! Initial</b> : call mpi_: call mpi_: call mpi_: <b>! Initial</b> :	llocatable :: ize MPI init(ierr) comm_size(mpi comm_rank(mpi ize the array	array(:) comm_world, comm_world,	<pre>nprocs,ierr) myid,ierr)</pre>		
[lyan1@ The cor	oqb563 ex]\$ tent of th	mpirun -np 4 e array:	ł ./a.out			_
	0	1 7	2	3	4	5
CENTER FOR COMPUTATION & TECHNOLOGY	do i=1, call n enddo write(* write(* else call mp endif	nprocs-1 mpi_recv(arra ,*) "The cont ,*) array i_send(array,	y(2*i+1),2,n ent of the a 2,mpi_intege	mpi_integer,i array:" er,0,myid,ier:	,i,status,ie r)	rr)





# **Blocking Operations**

- MPI\_SEND and MPI\_RECV are blocking operations
  - They will not return from the function call until the communication is completed
  - When a blocking send returns, the send buffer can be safely overwritten
  - When a blocking receive returns, the data has been received and is ready to use









# Deadlock (1)

- Deadlock occurs when both processes awaits the other to make progress
  - // Exchange data between two processes
    If (process 0)
     Receive data from process 1
     Send data to process 1
    If (process 1)
     Receive data from process 0
     Send data to process 0



This is a guaranteed deadlock because both receives will be waiting for data, but no send can be called until the receive returns







# Deadlock (2)

• How about this one?

// Exchange data between two processes
If (process 0)
 Receive data from process 1
 Send data to process 1
If (process 1)
 Send data to process 0
 Receive data from process 0





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# Deadlock (2)

• How about this one?

```
// Exchange data between two processes
If (process 0)
    Receive data from process 1
    Send data to process 1
If (process 1)
    Send data to process 0
    Receive data from process 0
```



No deadlock will occur – process 0 will receive the data first, then send the data to process 1; However, there will be performance penalty because we turn concurrent operations into sequential.







# Deadlock (3)

• And this one?

// Exchange data between two processes
If (process 0)
 Send data to process 1
 Receive data from process 1
If (process 1)
 Send data to process 0
 Receive data from process 0





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# Deadlock (3)

• And this one?

// Exchange data between two processes
If (process 0)
 Send data to process 1
 Receive data from process 1
If (process 1)
 Send data to process 0
 Receive data from process 0



It depends. If one of the sends returns, then we are OKAY - most MPI implementations buffer the message, so a send could return even before the matching receive is posted. However, if this is not the case or the message is too large to be buffered, deadlock will occur.







# Non-blocking Operations (1)

- Non-blocking operations separate the initialization of a send or receive from its completion
- Two calls are required to complete a send or receive
  - Initialization
    - Send: MPI\_ISEND
    - Receive: MPI\_IRECV



Completion: MPI\_WAIT





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# Non-blocking Operations (2)

- MPI\_ISEND
  - C: int MPI\_Isend(void \*buf, int count, MPI\_Datatype
     dtype, int dest, int tag, MPI\_Comm comm, MPI\_Request
     \*request);
  - Fortran:MPI\_ISEND(BUF, COUNT, DTYPE, DEST, TAG, COMM, REQ, IERR)
- MPI\_IRECV
  - C: int MPI\_Irecv(void \*buf, int count, MPI\_Datatype
     dtype, int source, int tag, MPI\_Comm comm,
     MPI\_Request \*request);
  - Fortran:MPI\_IRECV(BUF, COUNT, DTYPE, SOURCE, TAG, COMM, REQUEST, IERR)
- MPI\_WAIT



• Fortran: MPI\_WAIT(**REQUEST**, STATUS, IERR )

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# Example: Exchange Data with Nonblocking calls

integer reqids,reqidr
integer status(mpi\_status\_size)

- if (myid.eq.0) then
  - call mpi\_isend(to\_p1,n,mpi\_integer,1,100,mpi\_comm\_world,reqids,ierr)
  - call mpi\_irecv(from\_p1,n,mpi\_integer,1,101,mpi\_comm\_world,reqidr,ierr)
- elseif (myid.eq.1) then
  - call mpi\_isend(to\_p0,n,mpi\_integer,0,101,mpi\_comm\_world,reqids,ierr)

call mpi\_irecv(from\_p0,n,mpi\_integer,0,100,mpi\_comm\_world,reqidr,ierr)
endif

```
call mpi_wait(status,reqids,ierr)
call mpi_wait(status,reqidr,ierr)
```











# Blocking vs. Non-blocking

- Blocking operations are data corruption proof, but
  - Possible deadlock
  - Performance penalty
- Non-blocking operations allow overlap of completion and computation
  - The process can work on other things between the initialization and completion
  - Should be used whenever possible











# Exercise 2a: Find Global Maximum

- Goal: Find the maximum in an array
  - Each process handle part of the array
  - Every process needs to know the maximum at the end of program
- Hints
  - This can be done in two steps
    - Step 1: each process send the local maximum to the root process to find the global maximum
    - Step 2: the root process send the global maximum to all other processes









#### Exercise 2b: Laplace Solver Version 1

- Goal: develop a working MPI Laplace solver
  - Distribute the workload in a one-dimensional manner
  - Initialize the sub-matrix at each process and set the boundary values
  - At the end of each iteration
    - Exchange boundary data with neighbors
    - Find the global convergence error and distribute to all processes



