





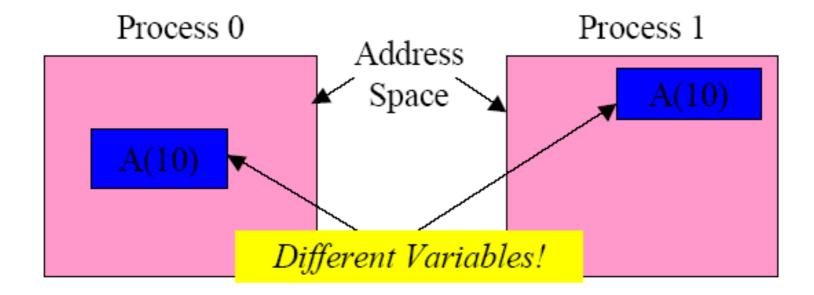
# Lo4: intro to MPI programming

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# Message passing paradigm

- Parallel programs consist of separate processes, each with its own address space
- Programmer manages memory by placing data in a particular process
- Data sent explicitly between processes
- Programmer manages memory motion
- Collective operations
  - On arbitrary set of processes
- Data distribution
  - Also managed by programmer

# Distributed memory (shared nothing approach)



#### What is MPI?

- A message-passing library specification
- extended message-passing model
- not a language or compiler specification
- not a specific implementation or product
- For parallel computers, clusters, and heterogeneous networks
- Full-featured
- Designed to provide access to advanced parallel hardware for end users, library writers, and tool developers
- Latest version of the standard MPI- 3.1

#### What is MPI?

#### A STANDARD...

- The actual implementation of the standard is demanded to the software developers of the different systems
- In all systems MPI has been implemented as a library of subroutines over the network drivers and primitives
- many different implementations
  - MPICH (the original one)
  - OpenMPI
  - IntelMPI

#### Some reason to use MPI

- International standard
- MPI evolves: MPI 1.0 was first introduced in 1994, most current version is MPI 3.3 (Nov. 2016)
- Available on almost all parallel systems (free MPICH, Open MPI used on many clusters), with interfaces for C/C++ and Fortran
- Supplies many communication variations and optimized functions for a wide range of needs
- Works both on distributed memory (DM) and shared memory (SM) hardware architectures
- Supports large program development and integration of multiple modules

### How to program with MPI?

- MPI is a library:
  - All operations are performed with subroutine calls
- Basic definitions are in
  - mpi.h for C/C++
  - mpif.h for Fortran 77 and 90
  - MPI module for Fortran 90 (optional)

# **Compiling MPI Programs**

#### NO STANDARD: left to the implementations:

- Generally:
  - You should specify the appropriate include directory (i.e. -I/mpidir/include)
  - You should specify the mpi library (i.e. -L/mpidir/lib -lmpi)
- Usually MPI compiler wrappers do this job for you. (i.e. Mpicc)
  - Check on your machine...

# Running MPI programs

The MPI Standard does not specify how to run an MPI program, just as the Fortran standard does not specify how to run a Fortran program.

In general, starting an MPI program is dependent on the implementation of MPI you are using, and might require various scripts, program arguments, and/or environment variables.

Many implementations provided mpirun -np 4 a.out to run an MPI program

Many parallel systems use a batch environment to share resources among users

The specific commands to run a program on a parallel system are defined by the environment installed on the parallel computer

# Writing MPI program..

 Modify your serial program to insert MPI routines to distribute data and loads to different processors.

#### **Basic Features of MPI routines**

#### Calls may be roughly divided into four classes:

- Calls used to initialize, manage, and terminate communications
- Calls used to communicate between pairs of processors.
   (Pair communication)
- Calls used to communicate among groups of processors.
   (Collective communication)
- Calls to create data types.

#### **Basic MPI routines**

- MPI INIT: initialize MPI
- MPI\_COMM\_SIZE: how many Processors?
- MPI\_COMM\_RANK: identify the Processor
- MPI SEND: send data
- MPI RECV: receive data
- MPI FINALIZE: close MPI

(Almost) All you need is to know this 6 calls

# Your First Program: Hello World!

```
PROGRAM hello
INCLUDE 'mpif.h'
INTEGER err
CALL MPI_INIT(err)
call MPI_COMM_RANK(MPI_COMM_WORLD,rank,ierr)
call MPI_COMM_SIZE(MPI_COMM_WORLD,size,ierr)
print *, 'I am ', rank, ' of ', size
CALL MPI_FINALIZE(err)
END
```

```
#include <stdio.h>
#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( "I am %d of %d\n", rank, size );
  MPI_Finalize();
}
```

#### Some initial notes

- All MPI programs begin with MPI\_Init and end with MPI\_Finalize
- MPI\_COMM\_WORLD is defined by mpi.h (in C) or mpif.h (in Fortran) and designates all processes in the MPI "job"
- Each statement executes independently in each process including the printf/print statements
  - I/O not part of MPI-1 (MPI-IO part of MPI-2)
  - print and write to standard output or error not part of MPI-1 or MPI-2 or MPI-3
- output order is undefined (may be interleaved by character, line, or blocks of characters),
- A consequence of the requirement that non-MPI statements execute independently

# **Initializing and Exiting MPI**

```
Initializing the MPI environment
C: int MPI Init(int *argc, char ***argv);
Fortran:
       INTEGER IERR
       CALL MPI INIT (IERR)
Finalizing MPI environment
       int MPI Finalize()
Fortran:
       INTEGER IERR
       CALL MPI FINALIZE (IERR)
This two subprograms should be called by all processes, and no other
MPI calls are allowed before mpi init and after mpi finalize
```

#### **MPI Communicator**

The Communicator is a variable identifying a group of processes that are allowed to communicate with each other.

There is a default communicator (automatically defined):

MPI\_COMM\_WORLD

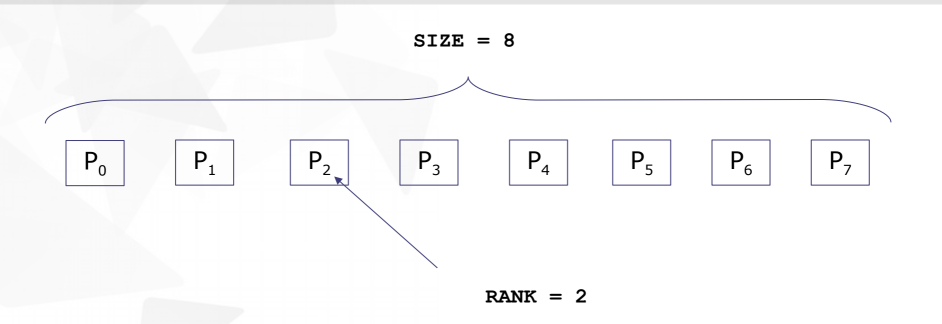
It identifies the group of all processes.

All MPI communication subroutines have a communicator argument. The Programmer could define many communicators at the same time

#### **Communicator Size and Process Rank**

```
How many processors are associated with a communicator?
C:
  MPI Comm size (MPI Comm comm, int *size)
Fortran:
  INTEGER COMM, SIZE, IERR
  CALL MPI COMM SIZE (COMM, SIZE, IERR)
OUTPUT: SIZE
What is the ID of a processor in a group?
C:
   MPI Comm rank(MPI Comm comm, int *rank)
Fortran:
   INTEGER COMM, RANK, IERR
   CALL MPI COMM RANK (COMM, RANK, IERR)
OUTPUT:
        RANK
```

#### **Communicator Size and Process Rank**



Size is the number of processors associated to the communicator

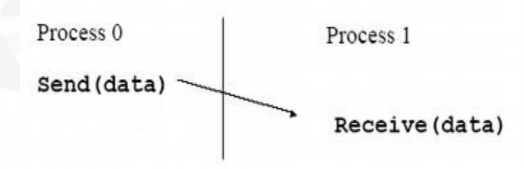
rank is the index of the process within a group associated to a communicator (rank = 0,1,...,N-1). The rank is used to identify the source and destination process in a communication

# **Communication Ingredients**

- To send a message via mail we typically have:
  - An envelope (with possibly some hints on the content itself...
     i.e., advertisement, bills, greetings....)
  - A message
  - A destination address
  - A sender address

For MPI it is exactly the same thing...

# MPI basic send/receive



#### questions:

How will "data" be described? datatypes
How will processes be identified? rank/comm
How will the receiver recognize messages? tag
What will it mean for these operations to complete?
blocking/non-blocking

# **Describing Data**

ones for subarrays

```
The data in a message to send or receive is described by a triple (address, count, datatype), where

An MPI datatype is recursively defined as:

predefined, corresponding to a data type from the language (e.g., MPI_INT, MPI_DOUBLE)

a contiguous array of MPI datatypes

a strided block of datatypes

an indexed array of blocks of datatypes

an arbitrary structure of datatypes

There are MPI functions to construct custom datatypes, in particular
```

# Fortran - MPI Basic Datatypes

MPI Data type	Fortran Data type
MPI_INTEGER	INTEGER
MPI_REAL	REAL
MPI_DOUBLE_PRECISION	DOUBLE PRECISION
MPI_COMPLEX	COMPLEX
MPI_DOUBLE_COMPLEX	DOUBLE COMPLEX
MPI_LOGICAL	LOGICAL
MPI_CHARACTER	CHARACTER (1)
MPI_PACKED	
MPI_BYTE	

# **C** - MPI Basic Datatypes

MPI Data type	C Data type	
MPI_CHAR	signed char	
MPI_SHORT	signed short int	
MPI_INT	signed int	
MPI_LONG	Signed log int	
MPI_UNSIGNED_CHAR	unsigned char	
MPI_UNSIGNED_SHORT	unsigned short int	
MPI_UNSIGNED	unsigned int	
MPI_UNSIGNED_LONG	unsigned long int	
MPI_FLOAT	float	
MPI_DOUBLE	double	
MPI_LONG_DOUBLE	long double	
MPI_BYTE		
MPI_PACKED		

### Data tag

- Messages are sent with an accompanying user-defined integer tag, to assist the receiving process in identifying the message
- Messages can be screened at the receiving end by specifying a specific tag, or not screened by specifying MPI\_ANY\_TAG as the tag in a receive

# Our First Sent Message....

#### The simplest call:

```
MPI_send( buffer, count, data_type, destination, tag, communicator)
```

#### where:

**BUFFER:** data to send

**COUNT:** number of elements in buffer.

DATA\_TYPE: which kind of data types in buffer?

**DESTINATION** the receiver

TAG: the label of the message

**COMMUNICATOR** set of processors involved

# ..and our First Received message.

#### The simplest call:

MPI\_recv( buffer, count, data\_type, source, tag, communicator, status)

Similar to send with the following differences:

- SOURCE is the sender; can be set as MPI\_any\_source (receive a message from any processor within the communicator)
- TAG the label of message: can be set as MPI\_any\_tag: receive any kind of message
- •STATUS integer array with information on message in case of error

### The status array

Status is a data structure allocated in the user's program.

```
In C:
   int recvd tag, recvd from, recvd count;
   MPI Status status;
   MPI Recv(..., MPI ANY SOURCE, MPI ANY TAG, ..., &status )
   recvd tag = status.MPI TAG;
   recvd from = status.MPI SOURCE;
   MPI Get count ( &status, datatype, &recvd count );
In Fortran:
  integer recvd tag, recvd from, recvd count
  integer status (MPI STATUS SIZE)
  call MPI RECV(..., MPI ANY SOURCE, MPI ANY TAG, .. status, ierr)
  tag recvd = status(MPI TAG)
  recvd from = status(MPI SOURCE)
  call MPI GET COUNT(status, datatype, recvd count, ierr)
```

### MPI: a FORTRAN example..

```
Program MPI
   Implicit None
   Include 'mpif.h'
   Integer
                                            :: rank
                                            :: buffer
   Integer
   Integer, Dimension( 1:MPI status size ) :: status
   Integer
                                            :: error
   Call MPI init( error )
   Call MPI comm rank ( MPI comm world, rank, error )
   If (rank == 0) Then
      buffer = 33
      Call MPI send( buffer, 1, MPI integer, 1, 10, &
                     MPI comm world, error )
   End If
   If (rank == 1) Then
      Call MPI recv( buffer, 1, MPI integer, 0, 10, &
                     MPI comm world, status, error )
      Print*, 'Rank ', rank, ' buffer=', buffer
      If( buffer /= 33 ) Print*, 'fail'
   End If
   Call MPI finalize( error )
End Program MPI
```

# **Blocking and Non-Blocking**

Q: When is a SEND instruction complete?

A: When it is safe to change the data that we sent.

Q: When is a RECEIVE instruction complete?

A: When it is safe to access the data we received.

With both communications (send and receive) we have two choices:

Start a communication and wait for it to complete:

**BLOCKING** approach

Start a communication and return control to the main program:

**NON-BLOCKING** approach

The Non-Blocking approach REQUIRES us to check for completion before we can modify/access the sent/received data!!!

# MPI\_send and MPI\_recv

MPI\_SEND() and MPI\_RECV() are blocking operations.

# Arere they really blocking?

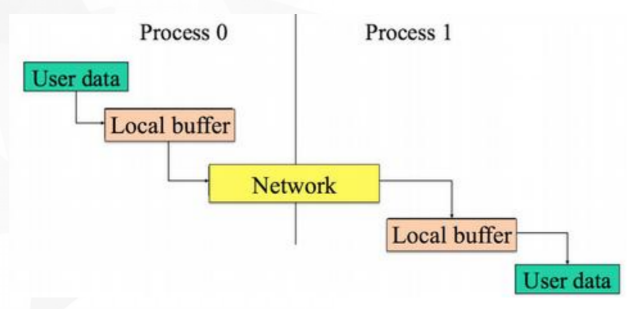
#### MPI\_SEND() and MPI\_RECV() are blocking operations.

However: often a system buffer is used that allows small messages to be non-blocking send-recv handshakes, but large messages will be blocking.

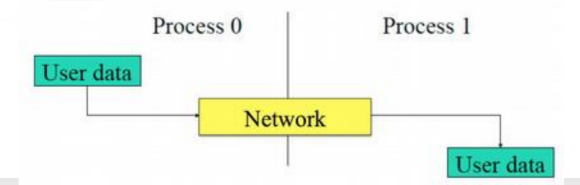
MPI implementation (not the MPI standard) decides this Blocking communication can be unsafe and may lead to deadlocks.

# Again on this

Small messages make use of system-supplied buffer



Large message are really blocking:



# Pros and Cons of Non-Blocking Send/ Receive

- Non-Blocking communications allows the separation between the initiation of the communication and the completion.
- Advantages:
  - between the initiation and completion the program could do other useful computation (latency hiding).
- Disadvantages:
  - the programmer has to insert code to check for completion.

#### **Communication mode**

- 4 different send types:
  - Standard: let MPI decide the best strategy...
  - Synchronous: it is complete when the receiver acknowledged the reception of the message
  - Buffered: it is complete when the data has been copied to a local buffer
  - Ready: requires a receiver to be already waiting for the message
- 1 single receive type

#### **Communication Modes and MPI Subroutines**

Mode	Completion Condition	Blocking subroutine	Non-blocking subroutine
Standard send	Message sent (receive state unknown)	MPI_SEND	MPI_ISEND
receive	Completes when a message has arrived	MPI_RECV	MPI_IRECV
Synchronous send	Only completes when the receive has completed	MPI_SSEND	MPI_ISSEND
Buffered send	Always completes, irrespective of receiver	MPI_BSEND	MPI_IBSEND
Ready send	Always completes, irrespective of whether the receive has completed	MPI_RSEND	MPI_IRSEND

# **Non-Blocking Send and Receive**

```
Fortran:
MPI ISEND (buf, count, type, dest, tag, comm, req, ierr)
MPI IRECV(buf, count, type, dest, tag, comm, req, ierr)
buf
        array of type type see table.
        (INTEGER) number of element of buf to be sent
count
type (INTEGER) MPI type of buf
dest (INTEGER) rank of the destination process
        (INTEGER) number identifying the message
tag
        (INTEGER) communicator of the sender and receiver
comm
        (INTEGER) output, identifier of the communications handle
req
        (INTEGER) output, error code (if ierr=0 no error occurs)
ierr
```

### Non-Blocking Send and Receive

C: int MPI Isend(void \*buf, int count, MPI Datatype type, int dest, int tag, MPI Comm comm, MPI Request \*req); int MPI Irecv (void \*buf, int count, MPI Datatype type, int dest, int tag, MPI Comm comm, MPI Request \*req);

# **Waiting and Testing for Completion**

```
Fortran:
```

```
MPI_WAIT(req, status, ierr)
```

A call to this subroutine cause the code to wait until the communication pointed by req is complete.

**Req** INTEGER) input/output, identifier associated to a communications event (initiated by MPI\_ISEND or MPI\_IRECV).

**Status** (INTEGER) array of size

MPI\_STATUS\_SIZE, if req was associated to a call to MPI\_IRECV, status contains informations on the received message, otherwise status could contain an error code.

**ierr** (INTEGER) output, error code (if **ierr=o** no error occours).

C:

int MPI\_Wait(MPI\_Request \*req, MPI\_Status \*status);

# **Waiting and Testing for Completion**

```
Fortran:
MPI TEST(req, flag, status, ierr)
A call to this subroutine sets flag to .true. if the
communication pointed by req is complete, sets
flag to .false. otherwise.
      (INTEGER) input/output, identifier associated to a communications event
(initiated by MPI ISEND or MPI IRECV).
Flag (LOGICAL) output, .true. if communication req
has completed .false. otherwise
Status (INTEGER) array of size MPI STATUS SIZE, if
req was associated to a call to MPI IRECV, status contains informations
 on the received message, otherwise status could contain an error code.
ierr
      (INTEGER) output, error code (if ierr=0 no error occours).
C:
int MPI Wait(MPI Request *req, int *flag, MPI Status *status);
```

#### MPI: a case study

#### Problem: exchanging data between two processes



#### **DEADLOCK**

#### Solution A

# USE BUFFERED SEND: bsend send and go back so the deadlock is avoided

#### **NOTES:**

1. Requires a copy therefore is not efficient for large data set memory problems

#### **Solution B**

# Use non blocking SEND : isend send go back but now is not safe to change the buffer

#### **NOTES:**

- 1 A handle is introduced to test the status of message.
- 2. More efficient of the previous solutions

#### Solution C

#### Exchange send/recv order on one processor

#### NOTES:

efficient and suggested!

#### **Exercises and tutorials**

- Compile/Run and understand usage of MPI programs
  - mpi pi.c
  - hello\_world.c/f90
  - mpi\_env\_call.c
  - send\_message.f90
- First MPI exercise: fix deadlock problem
- Second MPI exercise: play with MPI\_send call on mpi\_pi.c
- Third MPI exercise: implement the sum of N number using MPI paradigm

# **END OF FIRST PART**