# DISTRIBUTED AND CLOUD COMPUTING

LAB 1: MPI INTRODUCTION AND SETUP

(Module: Message Passing)

### **Overview**

- During the lab:
  - Introduction to technical concepts in Distributed and Cloud Computing
  - Application of concepts and Q&A
- Programming language: C, Java, Python, Go
- Blackboard site:
  - Distributed and Cloud Computing Fall 2025: bb.sustech.edu.cn

### **Distributed Computing Intuition**

Doing a task alone v.s. doing a task in a group?

Can you think of one good thing and one bad thing for each?

# **Distributed Computing Intuition**

#### Alone:

- + Management is simple
- You have to do everything

### Group:

- + Work is split amongst many people (ideally!)
- You have to communicate and assign tasks

Same things apply to computers!

# **MPI – Message Passing Interface**



- Core requirements for all parallel systems: Communication
  - Shared Memory, or
  - Message Passing
- MPI: a message passing standard to allow for distributed/parallel computation
  - o Many implementations [open-mpi, mpich, ...] with various languages [C, C++, FORTRAN, ...]
- This course: open-MPI with C
- MPI is extremely common for High Performance Computing (HPC) applications
  - o If sth runs on a supercomputer (and does not use GPUs) there's a good chance it uses MPI

# MPI – Message Passing Interface



- MPI implements an interface for parallel process communication
  - Abstracts the low-level details of processes communication
  - Allows the programmer to focus on the problem at hand (the parallel application)!
- You can think of MPI as a postal service and the processes as people sending letters
  - When you want to send a letter, you just drop it in the mailbox!





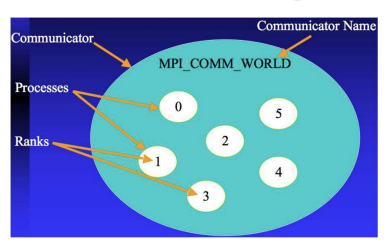
- MPI processes are managed by MPI and run concurrently (at the same time)
  - On the CPU cores of a computer or server
  - On the cores of many CPUs each in a cluster of network computers

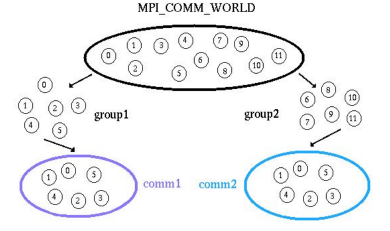
#### • Each process:

- Runs an instance of the parallel application (e.g. large weather model)
- Has its own local memory (does not share memory with any other process)
- Can use the MPI interface to send messages to other processes



- MPI communicators (neighbourhoods)
  - Group processes
  - Assign them unique IDs called ranks (house addresses)
- "MPI\_COMM\_WORLD" is the default communicator
  - Contains all MPI processes generated for the application
- Customized communicators can be defined by the user
  - Can contain just a subset of the processes
  - Useful for managing more complex communication patterns





# Setting Up the Environment for Open MPI

### **WINDOWS**

#### Opt. 1: WSL2 (RECOMMENDED)

- Open the windows store and search 'Ubuntu' and install
- Open ubuntu app and set up username and password when prompted

#### Opt. 2: Ubuntu Virtual Machine

- Download a VM hypervisor
  - VirtualBox is free and recommended
- Download an ISO of the latest ubuntu LTS
- o There are plenty of guides online to help you

### **MAC OS**

#### • Opt. 1: Install locally (RECOMMENDED)

Through homebrew or building from source

#### Opt. 2: Ubuntu Virtual Machine

- not recommended: macs natively support openMPI
- Similar to Windows
- o Intel mac: VirtualBox
- Apple silicone: VMware fusion

# Setting Up the Environment for Open MPI

### **Ubuntu (Debian)**

• Run the following commands in the terminal:

```
sudo apt-get update
sudo apt install gcc (c compiler if not already installed)
sudo apt-get install openmpi-bin openmpi-doc libopenmpi-dev
```

#### **MAC OS**

Get xcode-command line tools: xcode-select --install

#### Opt. 1: Homebrew (RECOMMENDED)

- Install homebrew packet manager: <a href="https://brew.sh/">https://brew.sh/</a>
- Run: brew install open-mpi

If download is **slow** or blocked set up the brew sustech mirror: <a href="https://mirrors.sustech.edu.cn/help/homebrew.html#introduction">https://mirrors.sustech.edu.cn/help/homebrew.html#introduction</a>

#### Opt. 2: Build openMPI (NOT RECOMMENDED)

- First answer at:
   <a href="https://www.open-mpi.org/fag/?category=building">https://www.open-mpi.org/fag/?category=building</a>
- Download the zipped source code and unzip
- Go the unzipped directory and run the configuration
- Make all install

```
#include <mpi.h>
#include <stdio.h>
int main(int argc, char* argv[]) {
   // Initialization
   MPI Init (NULL, NULL);
    // Get the number of processes
    int world size;
   MPI Comm size (MPI COMM WORLD, &world size);
    // Get the rank of the process
    int world rank;
   MPI Comm rank (MPI COMM WORLD, &world rank);
    // Get the name of the processor
    char processor name[MPI MAX PROCESSOR NAME];
    int name len;
   MPI Get processor name (processor name, &name len);
    // Print off a hello world message
    printf("Hello world from processor %s, rank %d out of %d processor n",
            processor name, world rank, world size);
    // Finalize the MPI environment.
   MPI Finalize();
```

### **Open MPI - Boiler Plate Code**

THIS CODE NEVER CHANGES AND IS ALWAYS USED WHEN WORKING WITH MPI

# **Open MPI - Useful Functions**

```
// Get the number of processes
int world_size;
MPI_Comm_size(MPI_COMM_WORLD, &world_size);

// Get the rank of the process
int world_rank;
MPI_Comm_rank(MPI_COMM_WORLD, &world_rank);

Sets world_size equal equal to the total number of MPI processes

Sets world_rank equal equal to the rank of the current process
```

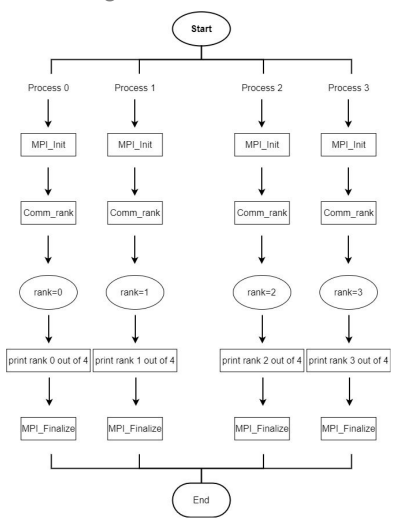
```
// Get the name of the processor
char processor_name[MPI_MAX_PROCESSOR_NAME];
int name_len;
MPI_Get_processor_name(processor_name,
&name_len);
```

Retrieves the name of the processor

(usually, the name you have given to the PC like GeorgesLaptop)

```
#include <mpi.h>
#include <stdio.h>
int main(int argc, char* argv[]) {
    // Initialization
   MPI Init(NULL, NULL);
    // Get the number of processes
    int world size;
    MPI Comm size (MPI COMM WORLD, &world size);
    // Get the rank of the process
    int world rank;
    MPI Comm rank (MPI COMM WORLD, &world rank);
    // Get the name of the processor
    char processor name[MPI MAX PROCESSOR NAME];
    int name len;
    MPI Get processor name (processor name, &name len);
    // Print off a hello world message
    printf("Hello world from processor %s, rank %d out of %d
         processors\n", processor name, world rank, world size);
    // Finalize the MPI environment.
    MPI Finalize();
```

#### **Diagram of MPI execution**



https://github.com/wesleykendall/mpitutoria l/blob/gh-pages/tutorials/mpi-hello-world/ code/mpi\_hello\_world.c

### **Compilation and Execution**

To compile, use mpicc, which is just a wrapper around gcc: mpicc source.c -o executable\_name

Then run the program use mpirun:

mpirun -np X ./executable\_name "-np X" : tells MPI how many processes to create

#### The result:

```
george@DESKTOP-E24BUDU:~/DistSys24/MPI/w1$ mpicc mpi_intro.c -o intro
george@DESKTOP-E24BUDU:~/DistSys24/MPI/w1$ mpirun -np 8 ./intro
Hello world from processor DESKTOP-E24BUDU, rank 7 out of 8 processors
Hello world from processor DESKTOP-E24BUDU, rank 0 out of 8 processors
Hello world from processor DESKTOP-E24BUDU, rank 4 out of 8 processors
Hello world from processor DESKTOP-E24BUDU, rank 5 out of 8 processors
Hello world from processor DESKTOP-E24BUDU, rank 6 out of 8 processors
Hello world from processor DESKTOP-E24BUDU, rank 1 out of 8 processors
Hello world from processor DESKTOP-E24BUDU, rank 2 out of 8 processors
Hello world from processor DESKTOP-E24BUDU, rank 3 out of 8 processors
george@DESKTOP-E24BUDU:~/DistSys24/MPI/w1$
```

### **Compilation and Execution**

- Every process runs the same program! What's the point?
  - Ranks!

Every process has a unique rank which we can use to assign it different tasks!

```
#include <mpi.h>
#include <stdio.h>
int main(int argc, char* argv[]) {
    // Initialization
    MPI Init(NULL, NULL);
    // Get the number of processes
    int world size;
    MPI_Comm_size(MPI_COMM_WORLD, &world_size);
    // Get the rank of the process
    int world rank;
    MPI Comm rank (MPI COMM WORLD, &world rank);
    // Get the name of the processor
    char processor name[MPI MAX PROCESSOR NAME];
    int name len;
    MPI Get processor name (processor name, &name len);
    if(world rank % 2 == 0) {
     // Print a hello world message
     printf("Hello world from processor %s, rank %d out of %d
         processors\n", processor name, world rank, world size);
    // Finalize the MPI environment.
    MPI Finalize();
```

```
george@DESKTOP-E24BUDU:~/DistSys24/MPI/w1$ mpicc mpi_intro_plus.c -o plus
george@DESKTOP-E24BUDU:~/DistSys24/MPI/w1$ mpirun -np 8 ./plus
Hello world from processor DESKTOP-E24BUDU, rank 2 out of 8 processors
Hello world from processor DESKTOP-E24BUDU, rank 4 out of 8 processors
Hello world from processor DESKTOP-E24BUDU, rank 6 out of 8 processors
Hello world from processor DESKTOP-E24BUDU, rank 0 out of 8 processors
george@DESKTOP-E24BUDU:~/DistSys24/MPI/w1$
```

### **Compilation and Execution - Extra**

#### Process to CPU core mapping

- MPI runs new processes in what we call slots
- By default, MPI identifies the number of CPU cores and creates an equal number of slots
- In this case, due to multithreading, there are 16 threads available for processing
- Going over that number is possible, but processes are not fully parallel (CPU task scheduling)

```
george@DESKTOP-E24BUDU:~/DistSys24/MPI/w1$ mpirun -np 16 ./intro
```

There are not enough slots available in the system to satisfy the 16 slots that were requested by the application:

```
./intro
```

Either request fewer slots for your application, or make more slots available for use.

A "slot" is the Open MPI term for an allocatable unit where we can launch a process. The number of slots available are defined by the environment in which Open MPI processes are run:

```
george@DESKTOP-E24BUDU:~/DistSys24/MPI/w1$ mpirun -np 16 --oversubscribe ./intro
Hello world from processor DESKTOP-E24BUDU, rank 5 out of 16 processors
Hello world from processor DESKTOP-E24BUDU, rank 8 out of 16 processors
Hello world from processor DESKTOP-E24BUDU, rank 9 out of 16 processors
Hello world from processor DESKTOP-E24BUDU, rank 10 out of 16 processors
Hello world from processor DESKTOP-E24BUDU, rank 3 out of 16 processors
Hello world from processor DESKTOP-E24BUDU, rank 13 out of 16 processors
Hello world from processor DESKTOP-E24BUDU, rank 14 out of 16 processors
Hello world from processor DESKTOP-E24BUDU, rank 15 out of 16 processors
Hello world from processor DESKTOP-E24BUDU, rank 0 out of 16 processors
Hello world from processor DESKTOP-E24BUDU, rank 11 out of 16 processors
<u>Hello world from processor DESKTOP-E24BUDU, rank 12 out of 16 processors</u>
Hello world from processor DESKTOP-E24BUDU, rank 4 out of 16 processors
Hello world from processor DESKTOP-E24BUDU, rank 7 out of 16 processors
Hello world from processor DESKTOP-E24BUDU, rank 1 out of 16 processors
Hello world from processor DESKTOP-E24BUDU, rank 6 out of 16 processors
Hello world from processor DESKTOP-E24BUDU, rank 2 out of 16 processors
```

# **Configuration of MPI Run**

Official documentation available at:

https://www.open-mpi.org/doc/v4.0/man1/mpirun.1.php

Structure of mpirun: mpirun [MPI options] cprogram> [program arguments]

**IMPORTANT:** Be careful to add **MPI options BEFORE** the program! Otherwise, they will be treated as program arguments and will not take effect!