Lab Guide 7

Introduction to MPI

Objectives:

- basic message passing concepts
- concept of a single program on multiple data (SPMD)
- pipeline parallelism pattern

Introduction

This lab session aims to introduce the basic concepts of MPI programming, starting from a basic program with two processes, where one process sends a message to another process. The basic program will be extended to support any number of processes (a pipeline of N processes) and any number of messages among processes.

The program should be compiled in the cluster frontend using the mpicc -O2 prog.c command. To run the program, use the sbatch mpi.sh. The mpi.sh file should specify the required resources and should run the MPI program. The following example requests two PUs during 1 second and spawns two MPI processes:

```
[search7edu]$ cat mpi.sh
#!/bin/bash
#SBATCH --time=1:00
#SBATCH --ntasks=2
#SBATCH --partition=cpar
mpirun -np 2 ./a.out
```

Note that the number of requested resources (--ntasks) must be the same as the number of resources used in the mpirun command.

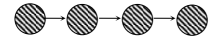
Exercise 1 - Pipeline of processes

Compile and run the following MPI program, where the process with rank 0 sends a message (integer value 123456) to the process with rank 1:

```
#include <mpi.h>
#include <stdio.h>
int main( int argc, char *argv[]) {
  int rank, msq;
  MPI Status status;
  MPI Init(&argc, &argv);
  MPI Comm rank ( MPI COMM WORLD, &rank ); // gets this process rank
  /* Process 0 sends and Process 1 receives */
  if (rank == 0) {
        msg = 123456;
        MPI_Send( &msg, 1, MPI INT, 1, 0, MPI COMM WORLD);
  }
  else if (rank == 1) {
        MPI Recv ( &msg, 1, MPI INT, 0, 0, MPI COMM WORLD, &status );
        printf( "Received %d\n", msg);
  MPI Finalize();
  return 0;
}
```

Modify the program to implement a pipeline of processes (using the SPMD model):

a) Start by modifying the program to support a pipeline with four processes: process with rank 0 sends the message that is successively processed (e.g., printed) by each process in the pipeline.



- b) Modify the program developed in a) to implement a pipeline with an arbitrary number of processes **specified as a parameter in the command** mpirun -np xx. Note that MPI is based on the SPMD style of parallel programming: the same process will be spawned xx times. The number of processes spawned by the mpirun command can be retrieved with the MPI Comm size call.
- c) Modify the program developed in b) to process 10 messages: the process with rank 0 should send 10 messages to the next in the pipeline; each other process should receive a message, process it (e.g., print) and send it to the next one in the pipeline.

Exercise 2 (optional) - Farm of processes and collective operations

Modify the original program to implement a directive-like behaving as "work sharing". A master process has a set of tasks to process, where each task will perform a given operation and produce as result an integer. Each worker receives the required data to process its task - which is a message with the argument (an integer) — and returns the processed task to the master.

Implement the following variations:

- a) Static scheduling: set the number of tasks to process equal to the number of MPI worker processes (one task per worker).
- b) Dynamic scheduling: set the number of tasks as 10x the number of MPI worker processes; faster processes should get more tasks.
- c) <u>Collective operations</u>: a message is broadcasted to all workers and then a reduce with the sum operation joins the results from all workers.