PPCES 2024: MPI Lab

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Acknoledgements: Parts of these code example have been developed by: - Christian Iwainsky - Sandra Wienke - Hristo Iliev - Joachim Jenke

Synopsis

The purpose of this hands-on lab is to make you familiar with the basic concepts of MPI. Tasks 1–3 will introduce you to the principles of basic point-to-point communication. Task 4 will practice the concept and usage of collective communications and MPI in general.

Building and executing lab examples

Building

The lab examples are written in C and require a C99 compiler. To ensure the C99 support, the CFLAGS in common/make.def specify the flag -std=c99, which switches on the C99 language support for most compilers. You may need to adapt this to the compiler used on your system.

MPI uses compiler wrappers to take care of all MPI-related command-line arguments for the preprocessor, compiler, and linker. The names for these wrappers is not standardized, yet mpicc is a common name for the C compiler wrapper. Therefore, this is the default compiler for the lab examples.

On most systems, you should be able to build an example with just

% make

Yet, you can override the compiler wrapper used by specifying it directly with the make command. The following example chooses the Intel Compiler with the Intel MPI library.

% make MPICC=mpiicc

Executing

MPI does not specify a specific launch command to be used to launch an MPI application, however, many MPI libraries provide the command mpiexec. If you are on a local system (like your laptop), this command will likely allow you to start the MPI application. On HPC platforms however, execution of parallel application may be performed by a different launch command, and an existing mpiexec in the path may not work as desired. Always consult your local HPC platform documentation for the details on launching MPI applications.

The CLAIX system at RWTH Aachen University defines environment variables for the convenience of the user. For the sake of these lab exercises, the following command on the cluster frontends will launch your MPI application on a shared partition.

Note: The \ followed by a line break allows to spread a single command across multiple lines. It is used here for the sole purpose of clarity, and is not needed if all your command line arguments are on a single line.

All exercises contain a special make target that will launch you program with an appropriate number of MPI processes.

% make run

They also contain a special make target that will submit the program as a batch job.

% make batch

1. Hello, MPI!

Lecture Sessions: MPI Overview, MPI Concepts The purpose of this exercise is to get you familiar with the very basics of MPI programming. Start with the minimal program hello.(c|f90) in directory 1_helloMPI and insert the appropriate code at the TODO markers.

2. Ping Pong

Lecture Sessions: Blocking Point-to-Point Communication One basic MPI program using point-to-point communication is the "ping pong" between two MPI processes. A ping-pong program skeleton can be found in directory **2**_pingPong. Complete the source code parts marked with "TODO".

- a) Make the first process of the MPI program transmit its input to the second process. The second process should then print the received value and send it back with an opposite sign to the first process, which should again print the received value.
- b) Make each rank send an individually and randomly selected number of elements. Let the other process know in advance the size of the array by explicitly sending it as an additional message.
- c) What is the behaviour of the program for NPROCS=1 and NPROCS>2? Modify it to display an error message when started with too few processes and to execute properly with more than two processes.
- d) Implement part b) of the assignment without explicitly sending the number of elements.
- e) Bonus task: Implement a loop to send/receive messages with different sizes. How does the message size influence the time being spent in MPI functions? You may use MPI_Wtime() to measure wall-clock time and go with array size as high as 226 elements to make the impact of the data size clearly visible.

3. Send-Receive

Lecture Sessions: Blocking/Non-blocking Point-to-Point Communication When multiple MPI processes exchange messages concurrently using blocking communication, the application may run into a deadlock if the communication pattern is not properly implemented. Use the code example 3_sendReceive/send_receive.(c|f90) and identify why this application runs into a deadlock. Use different techniques to overcome this problem.

(Note: You can abort the program execution when running interactively using Ctrl-c.)

- a) Modify the original code to use MPI_Send() and MPI_Recv() such that it becomes a correct MPI program and completes execution.
- b) Modify the original code to use MPI_Sendrecv() or MPI_Sendrecv_replace to avoid the deadlock.
- c) Modify the original code to use non-blocking communication on one of the point-to-point communication calls to avoid the deadlock.
- d) Modify the original code to use non-blocking communication on both of the point-to-point communication calls to avoid the deadlock.
- **e)** Modify the code from **d)** to work with more than 2 MPI processes. In this case the messages should be exchanged between adjacent ranks. (Hint: Will a special treatment be needed for the last rank?)

4. Derived Datatypes

Messages can be exchanged with different datatype handles on sender and receiver side, if the respective type signature of the buffers at sender and receiver-side matches.

- a) Extend the skeleton file 6_datatypes/derived.c and create datatypes on the sender and receiver side to transpose a 10 x 10 matrix in flight, by reading the data column-wise (use MPI_Type_vector) on the sender side and receiving row-wise (use MPI_Type_contiguous) on the receiver side. (*Hint: It is easiest to send separate messages for each column of data)
- b) Extend your solution for (a) and create a full matrix type to be able to send the data with a single message. Use a MPI_Type_create_hindexed (which is similar to MPI_Type_indexed, but uses byte displacements) to combine several columns of data into a single message. Think about why you cannot

use MPI_Type_indexed for this easily. ## 5. Simple Collectives ##### Lecture Sessions: Blocking Collective Communication

MPI collective operations describe common communication patterns among multiple processes. Implement the collectives bcast_int, scatter_int, gather_int, alltoall_int, and reduce_sum_int present in the skeleton file 4_simpleCollectives/collectives.c. Note that these collectives are simplified to work on integer buffers.

The skeleton file already contains printf statements to help you verify the correctness of your implementation. You can provide the rank in MPI_COMM_WORLD of the process that should perform the print statements. To test your implementation execute the application with different arguments.

Here is an example for testing with four ranks:

```
% make run NPROCS=4  # rank 0 is default
% make run NPROCS=4 PROG_ARGS=1 # select rank 1 to output
% make run NPROCS=4 PROG_ARGS=2 # select rank 2 to output
% make run NPROCS=4 PROG_ARGS=3 # select rank 3 to output
```

6. Creating new communicators

Lecture Sessions: Derived Datatypes Create new communicators as described in the subtask

- a) Create a duplicate of MPI_COMM_WORLD using MPI_Comm_dup and query the processes' rank and size for this communicator in dupRank and dupSize, respectively.
- b) Split MPI_COMM_WORLD such that the resulting communicators hold the MPI processes with odd and even ranks in MPI_COMM_WORLD, respectively. Query the processes' rank and size for this communicator in oddevenRank and oddevenSize, respectively.
- c) Split MPI_COMM_WORLD such that the resulting communicators hold the MPI processes with the ranks in MPI_COMM_WORLD below have of the size of MPI_COMM_WORLD and equal and above, respectively. Reorder the ranks such that the lowest rank in MPI_COMM_WORLD has the highest rank in the resulting communicator. Query the processes' rank and size for this communicator in upperlowerRank and upperlowerSize, respectively.