# Lecture 29 — Clusters & Cloud Computing

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ECE 459 Winter 2019 1/29

# Part I

# **MPI & Clusters**

ECE 459 Winter 2019 2/29

#### **Continuing to Switch Gears**

So far, we've seen how to make things fast on one computer:

- threads:
- compiler optimizations;
- GPUs.

To get a lot of bandwidth, though, you need lots of computers, (if you're lucky and the problem allows!)

Today: programming for performance with multiple computers via MPI.

ECE 459 Winter 2019 3 / 29

# **Key Idea: Explicit Communication**

Mostly we've seen shared-memory systems; complication: must manage contention.

Recently, GPU programming: explicitly copy data.

Message-passing: yet another paradigm.

ECE 459 Winter 2019 4/29

#### **Message Passing Interface:**

A language-independent communation protocol for parallel computers.

- Use it to run the same code on a number of nodes (different hardware threads; or servers in a cluster).
- Provides explicit message passing between nodes.
- Is the dominant model for high performance computing (de-facto standard).

ECE 459 Winter 2019 5 / 29

# High Level View of MPI

MPI is a type of SPMD (single process, multiple data).

Idea: have multiple instances of the same program, all working on different data.

The program could be running on the same machine, or a cluster of machines.

MPI facilitates communication of data between processes.

ECE 459 Winter 2019 6 / 29

```
// Initialize MPI
int MPI Init(int *argc. char **argv)
// Determine number of processes within a communicator
int MPI Comm size (MPI Comm comm. int *size)
// Determine processor rank within a communicator
int MPI Comm rank (MPI Comm comm. int *rank)
// Exit MPI (must be called last by all processors)
int MPI Finalize()
// Send a message
int MPI Send (void *buf, int count, MPI Datatype datatype,
              int dest, int tag, MPI Comm comm)
// Receive a message
int MPI Recy (void *buf.int count. MPI Datatype datatype.
              int source, int tag, MPI_Comm comm,
              MPI Status *status)
```

ECE 459 Winter 2019 7/29

#### **MPI Function Notes**

- MPI\_Comm: a communicator, often MPI\_COMM\_WORLD for global channel.
- MPI\_Datatype: just an enum, e.g. MPI\_FLOAT\_INT, etc.
- dest/source: "rank" of the process (in a communicator) to send a message to/receive a message from; you may use MPI\_ANY\_SOURCE in MPI\_Recv.
- Both MPI\_Send and MPI\_Recv are blocking calls—
   see man MPI\_Send or man MPI\_Recv for more details.
- The tag allows you to organize your messages, so you can filter all but a specific tag.

ECE 459 Winter 2019 8 / 29

```
As with OpenCL kernels:
  first, figure out what "current" process is supposed to compute.
// http://www.dartmouth.edu/~rc/classes/intro_mpi/
#include <stdio.h>
#include <mpi.h>
int main (int argc, char * argv[])
  int rank. size:
  /* start MPT */
  MPI_Init (&argc, &argv);
  /* get current process id */
  MPI_Comm_rank (MPI_COMM_WORLD, &rank);
  /* get number of processes */
  MPI_Comm_size (MPI_COMM_WORLD, &size);
  printf("Hello world from process %d of %d\n", rank, size);
  MPI_Finalize():
  return 0;
```

# Longer MPI Example (from Wikipedia)

#### Here's a common example:

- The "master" (rank 0) process creates some strings and sends them to the worker processes.
- The worker processes modify their string and send it back to the master.

#### Source:

http://en.wikipedia.org/wiki/Message\_Passing\_Interface.

ECE 459 Winter 2019 10 / 29

```
"Hello World" MPI Test Program
#include <mpi.h>
#include <stdio.h>
#include <string.h>
#define BUFSIZE 128
#define TAG 0
int main(int argc, char *argv[])
  char idstr[32];
  char buff[BUFSIZE];
  int numprocs;
  int myid;
  int i;
  MPI_Status stat;
```

ECE 459 Winter 2019 11/29

#### Example Code (2)

```
/* all MPI programs start with MPI_Init; all 'N'
 * processes exist thereafter
 */
MPI_Init(&argc,&argv);

/* find out how big the SPMD world is */
MPI_Comm_size(MPI_COMM_WORLD, &numprocs);

/* and this processes' rank is what? */
MPI_Comm_rank(MPI_COMM_WORLD, &myid);

/* At this point, all programs are running equivalently;
 * the rank distinguishes the roles of the programs in
 * the SPMD model, with rank 0 often used specially...
 */
```

ECE 459 Winter 2019 12 / 29

ECE 459 Winter 2019 13 / 29

```
else
  /* receive from rank 0: */
  MPI_Recv(buff, BUFSIZE, MPI_CHAR, 0, TAG,
           MPI_COMM_WORLD, &stat);
  sprintf(idstr, "Processor %d", myid);
  strncat(buff, idstr, BUFSIZE-1);
  strncat(buff, "reporting for duty", BUFSIZE-1):
  /* send to rank 0: */
  MPI Send (buff, BUFSIZE, MPI CHAR, 0, TAG,
           MPI COMM WORLD):
/* MPI Programs end with MPI Finalize; this is a weak
 * synchronization point.
 */
MPI Finalize():
return 0;
```

ECE 459 Winter 2019 14/29

#### Compiling with OpenMPI

```
// Wrappers for gcc (C/C++)
mpicc
mpicxx

// Compiler Flags
OMPI_MPICC_CFLAGS
OMPI_MPICXX_CXXFLAGS

// Linker Flags
OMPI_MPICC_LDFLAGS
```

OMPI MPICXX LDFLAGS

OpenMPI does not recommend that you set the flags yourself. To see them, try:

```
# Show the flags necessary to compile MPI C applications shell$ mpicc —showme:compile
```

# Show the flags necessary to link MPI C applications shells mpicc —showme: link

ECE 459 Winter 2019 15 / 29

# **Compiling and Running**

Starts num\_processors instances of the program using MPI.

```
jon@riker examples master % mpicc hello_mpi.c
jon@riker examples master % mpirun —np 8 a.out
0: We have 8 processors
0: Hello 1! Processor 1 reporting for duty
0: Hello 2! Processor 2 reporting for duty
0: Hello 3! Processor 3 reporting for duty
0: Hello 4! Processor 4 reporting for duty
0: Hello 5! Processor 5 reporting for duty
0: Hello 6! Processor 6 reporting for duty
0: Hello 6! Processor 7 reporting for duty
```

■ By default, MPI uses the lowest-latency communication resource available; shared memory, in this case.

#### MPI Matrix Multiplication Example

Highlights of: http://www.nccs.gov/wp-content/training/mpi-examples/C/matmul.c. To compute the matrix product AB:

- Initialize MPI.
- 2 If the current process is the master task (task id 0):
  - 1 Initialize matrices.
  - 2 Send work to each worker task: row number (offset); number of rows; row contents from A; complete contents of matrix B.

- Wait for results from all worker tasks (MPI\_Recv).
- 4 Print results.
- For all other tasks:
  - Receive offset, number of rows, partial matrix A, and complete matrix B, using MPT Recy:

- 2 Do the computation.
- 3 Send the results back to the sender.

ECE 459 Winter 2019 17 / 29

# Other Things MPI Can Do

We can use nodes on a network (by using a hostfile).

We can even use MPMD:

■ multiple processes, multiple data

```
% mpirun -np 2 a.out : -np 2 b.out
```

This launches a single parallel application.

- All in the same MPI\_COMM\_WORLD; but
- Ranks 0 and 1 are instances of a.out, and
- Ranks 2 and 3 are instances of b.out.

You could also use the -app flag with an appfile instead of typing out everything.

#### **Performance Considerations**

Your bottleneck for performance here is message-passing.

Keep the communication to a minimum!

In general, the more machines/farther apart they are, the slower the communication.

Each step from multicore machines to GPU programming to MPI triggers an order-of-magnitude decrease in communication bandwidth and similar increase in latency.

ECE 459 Winter 2019 19 / 29

#### Part II

# **Cloud Computing**

ECE 459 Winter 2019 20 / 29

# Using a Cluster

#### Historically:

- find \$\$\$;
- buy and maintain pile of expensive machines.

Not anymore!

We'll talk about Amazon's Elastic Compute Cloud (EC2) and principles behind it.

ECE 459 Winter 2019 21/29

#### **Evolution of servers**

You want a server on the Internet.

- Once upon a time: had to get a physical machine hosted (e.g. in a rack).
   Or, live with inferior shared hosting.
- Virtualization: pay for part of a machine on that rack. A win: you're usually not maxing out a computer, and you'd be perfectly happy to share it with others, as long as there are good security guarantees. All users can get root access.
- Clouds enable you to add more machines on-demand. Instead of having just one virtual server, spin up dozens (or thousands) of server images when you need more compute capacity. Servers typically share persistent storage, also in the cloud.

ECE 459 Winter 2019 22 / 29

#### **Paying for Computes**

#### Cloud computing:

pay according to the number of machines, or instances, that you've started up.

Providers offer different instance sizes; sizes vary according to the number of cores, local storage, and memory.

Some instances even have GPUs!

ECE 459 Winter 2019 23 / 29

#### Launching Instances

Need more computes? Launch an instance!

Input: Virtual Machine image.

Mechanics: use a command-line or web-based tool.

New instance gets an IP address and is network-accessible.

You have full root access to that instance.

ECE 459 Winter 2019 24/29

#### What to Launch?

#### Amazon provides public images:

- different Linux distributions;
- Windows Server; and
- OpenSolaris (maybe not anymore?).

You can build an image which contains software you want, including Hadoop and OpenMPI.

ECE 459 Winter 2019 25 / 29

#### Cleanup

Presumably you don't want to pay forever for your instances.

When you're done with an instance:

■ shut it down, stop paying for it.

All data on instance goes away.

ECE 459 Winter 2019 26 / 29

# **Data Storage**

To keep persistent results:

- mount a storage device, also on the cloud (e.g. Amazon Elastic Block Storage); or,
- connect to a database on a persistent server (e.g. Amazon SimpleDB or Relational Database Service); or,
- you can store files on the Web (e.g. Amazon S3).

ECE 459 Winter 2019 27 / 29

#### Summary

- MPI is a powerful tool for highly parallel computing across multiple machines.
- MPI Programming is similar to a more powerful version of fork/join; but you have to manage communication more explicitly.

■ Saw cloud computing basics.

ECE 459 Winter 2019 28 / 29

# Movie Hour, featuring NoSQL Bane

Let's take a humorous look at cloud computing: James Mickens' session from Monitorama PDX 2014.

https://vimeo.com/95066828

ECE 459 Winter 2019 29 / 29