# Lecture 29 — Clusters & Cloud Computing

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# Part I

# **MPI & Clusters**

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### More, More, More

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!)

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Today: programming for performance with multiple computers via MPI.

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# 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.

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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).

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### 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.



MPI facilitates communication of data between processes.

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```
// 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 Recv (void *buf, int count, MPI Datatype datatype,
              int source, int tag, MPI_Comm comm,
              MPI Status *status)
```

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### 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.

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```
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 MPI */
  MPI_Init (&argc, &argv);^^I
  /* get current process id */
  MPI_Comm_rank (MPI_COMM_WORLD, &rank);^^I
  /* get number of processes */
  MPI_Comm_size (MPI_COMM_WORLD, &size);^^I
  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.

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# Example Code (1)

```
/*
 "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;
```

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# 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...
 */
```

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# Example Code (3)

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# Example Code (4)

```
else
  /* receive from rank 0: */
  MPI_Recv(buff, BUFSIZE, MPI_CHAR, 0, TAG,
           MPI COMM WORLD, &stat);
  sprintf(idstr. "Processor %d". mvid):
  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;
```

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### 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
shell$ mpicc --showme:link
```

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### Compiling and Running

```
mpirun -np <num_processors> <program>
mpiexec -np <num_processors> <program> # a synonym
```

#### 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 7! Processor 7 reporting for duty
```

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

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# Matrix Multiplication



Matrix Multiplication

**Neural Network** 

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### MPI Matrix Multiplication Example

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

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

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

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

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### 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.

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### 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.

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### Part II

# **Cloud Computing**



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### 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.

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### Evolution of servers

You want a server on the Internet.

- Once upon a time: physical machine; shared hosting.
- Virtualization:
- Clouds

Servers typically share persistent storage, also in the cloud.

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# Paying for Computes

Cloud computing: pay by the number of instances that you've started up.

Providers offer different instance sizes: vary in cores, memory, GPU...

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### 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.

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### 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.

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

When you're done with an instance:



All data on instance goes away.

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#### 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).

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### Clusters vs. Laptops

Key idea: scaling to big data systems introduces substantial overhead.

Up next: Laptop vs. 128-core big data systems.

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### Don't Guess, Measure

Are big data systems obviously good? Have we measured (the right thing)?

The important metric is not just scalability; absolute performance matters a lot.

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Don't want: scaling up to  ${\bf n}$  systems to deal with complexity of scaling up to  ${\bf n}$ .



Or, as Oscar Wilde put it: "The bureaucracy is expanding to meet the needs of the expanding bureaucracy."

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### Methodology

Compare: competent single-threaded implementation vs. top big data systems.

Domain: graph processing algorithms— PageRank and graph connectivity (bottleneck is label propagation).

Subjects: graphs with billions of edges (a few GB of data.)

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System	cores	twitter_rv	uk_2007_05			
Spark	128	857s	1759s			
Giraph	128	596s	1235s			
GraphLab	128	249s	833s			
GraphX	128	419s	462s			
Single thread	1	300s	651s			

#### Label propagation to fixed-point (graph connectivity)

System	cores	twitter_rv	uk_2007_05
Spark	128	1784s	8000s+
Giraph	128	200s	8000s+
GraphLab	128	242s	714s
GraphX	128	251s	800s
Single thread	1	153s	417s

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### Takeaways

- "If you are going to use a big data system for yourself, see if it is faster than your laptop."
- "If you are going to build a big data system for others, see that it is faster than my laptop."

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### 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

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