#### **RIT Computer Engineering Cluster**

The RIT Computer Engineering cluster contains 12 computers for parallel programming using MPI. One computer, *cluster-head.ce.rit.edu*, serves as the master controller or head node for the cluster and is accessible from the Internet. There is also a backup headnode, cluster-secondary, which also goes by the alias of *cluster.ce.rit.edu*. The other 12 machines, named cluster-node-01, through cluster-node-12, are attached to a private LAN segment and are visible only to each other and the cluster head node.

The hardware for each cluster node consists of the following:

- Supermicro AS-1012G Server Chassis
  - o Supermicro H8SGL-F
- Singe Socket 12 Core AMD Opteron 2.4GHz
- 3x 8GB PC3-8500
- 60GB Crucial SSD

To connect to the cluster, simply use a SSH or SFTP client to connect to:

```
<username>@cluster.ce.rit.edu
```

using your DCE login information (username and password).

The head node only supports secure connections using SSH and SFTP; normal Telnet and FTP protocols simply won't work.

#### **SSH Clients**

Putty, a very small and extremely powerful SSH client, is available from:

```
http://www.chiark.greenend.org.uk/~sgtatham/putty/
```

or from the mirror sight:

```
http://www.putty.nl/
```

This SSH client supports X11 forwarding, so if you use an XWindow emulator such as Exceed, ReflectionX, or Xming, you may open graphical applications remotely over the SSH connection. The website also includes a command line secure FTP client.

WinSCP is an excellent graphical FTP/SFTP/SCP client for Windows. It is available from:

```
http://winscp.net/eng/index.php
```

Xming X Server is a free X Window Server for Windows. It is available from:

```
http://www.straightrunning.com/XmingNotes/
```

# Using Message Passing Interface on the RIT Computer Engineering Cluster

MPI is designed to run Single Program Multiple Data (SPMD) parallel programs on homogeneous cluster or supercomputer systems. MPI uses shell scripts and the remote shell to start, stop, and run parallel programs remotely. Thus, MPI programs terminate cleanly, and require no additional housekeeping or special process management.

# **Summary**

Before using these commands, you will need to load the MPI module for use. Run the following command:

```
module load openmpi-x86 64
```

Compile using: mpicc [linking flags]
Run programs with: mpirun -np # executable

# **Specifying Machines**

The cluster is currently configured to execute jobs that are sent to the scheduler. You have no access to the compute nodes to run your jobs. Therefore, there is no way for you to specify which machines you want the processes to run on. In summary, the scheduler will handle this for you.

It is also prohibited to use mpirun on the head node. Any students that does this will have their processes immediately killed without warning.

### Compiling

To compile a MPI program, use the mpicc script. This script is a preprocessor for the compiler, which adds the appropriate libraries as appropriate. As it is merely an interface to the compiler, you may need to add the appropriate -1 library commands, such as -1m for the math functions. In addition, you may use -c and -o to produce object files or rename the output.

For example, to compile the test program:

```
[abc1234@phoenix mpi]$ mpicc greetings.c -o greetings
```

### **Running MPI Programs**

Use the mpirun program to execute parallel programs. The most useful argument to mpirun is -np, followed by the number of machines required for execution and the program name. The following is the output of the command to run the program. Your results will vary.

```
[jml1554@cluster-node-01 MultipleProcessorSystems]$ mpirun -np 3 greetings Process 2 of 3 on cluster-node-01 done Process 1 of 3 on cluster-node-01 done Greetings from process 1! Greetings from process 2! Process 0 of 3 on cluster-node-01 done
```

#### General syntax for mpirun is

```
mpirun -np <np> program
```

While this will work for the general case, it will not work for you since you don't have access to the compute nodes. This command will need to be placed in a script that is passed to the scheduler. More information about this process can be found on the course website and in the Job Submission document.

# **Programming Notes**

- All MPI programs require MPI Init and MPI Finalize.
- All MPI programs generally use MPI Comm rank and MPI Comm size.
- Printing debug output prefixed with the process's rank is extremely helpful.
- Printing a program initialization or termination line with the machine's name (using MPI Get processor name) is also suggested.
- If you're using C++, or C with C++ features (such as declarations other than at the start of the declaration) try using mpicc instead of mpicc.

## **CE Cluster Scheduling**

As mentioned above, the CE cluster uses a scheduler called SLURM. The purpose of SLURM is to adequately maintain the resources that are provided by the compute nodes. As you are developing your applications, you will need to be familiar with some of the basic SLURM commands that are outlined below.

*sinfo* – used to display the current state of the cluster

#### Example:

```
[jml1554@cluster-secondary ~]$ sinfo
PARTITION AVAIL TIMELIMIT NODES STATE NODELIST
class* up 4:00:00 10 idle cluster-node-[01-10]
```

*squeue* – used to display the current job queue; with the –u option, you can provide a username to view the jobs

#### Example:

sbatch – used to submit a job to the queue; a number of options can be used in two forms: one the command line, or in the script. In either case you need to use a script to submit your work. The easier of the two ways is to have the options embedded in the script as shown below. Make sure that you give your script execute permissions: chmod +x test.sh

# Script Example: test.sh

```
#!/bin/bash
# When the #SBATCH appears at the start of a line, it will
# be interpreted by the scheduler as a command for it
# Tell the scheduler that we want to use 13 cores for our job
#SBATCH -n 13
# Give the location of the stdout and stderr to be directed to
#SBATCH -o test.out
#SBATCH -e test.err
# Give the job a name
# You should give a unique name to each job to make it easily identifiable
#SBATCH -J Test
# Other options can be provided. Refer to the SLURM documentation for more parameters.
# SLURM: https://computing.llnl.gov/linux/slurm/
# You may also refer to http://mps.ce.rit.edu for more information
# Your commands go below this line
# This command MUST be in your script, otherwise the job will not run properly.
module load openmpi-x86 64
# This is where you need to provide the mpirun command
# $SLURM_NPROCS is set by SLURM when it handles the job. This value will be equal
# to the number given to -n from above. In this case, it will be 13.
# This should NOT be changed to a number; it will ensure that you are using only
# what you neeed.
mpirun -np $SLURM NPROCS greetings
```

# To submit the job to SLURM, use the following command:

sbatch test.sh

# You will see the following output if your job is submitted successfully:

Submitted batch job 5749

After your job completes, you can view the output from the text files test.out and/or test.err using any text editor. A fragment of the output file is provided below.

```
[jml1554@cluster-secondary MultipleProcessorSystems]$ more test.out
Process 2 of 10 on cluster-node-01 done
Process 8 of 10 on cluster-node-01 done
Process 4 of 10 on cluster-node-01 done
Process 5 of 10 on cluster-node-01 done
Process 10 of 10 on cluster-node-01 done
Greetings from process 1!
Greetings from process 2!
Process 12 of 13 on cluster-node-02 done
```

### Below is a sample program which you can compile and run:

```
// greetings.c
//
#include <stdio.h>
#include <string.h>
#include "mpi.h"
main( int argc, char *argv[] )
       // General identity information
       int my_rank;  // Rank of process
       int p;
                               // Number of processes
       // Size of local processor name
       int my_name_len;
       // Message packaging
       int source;
       int dest;
       int tag=0;
       char message[100];
       MPI Status status;
       //
       // Start MPI
       MPI_Init( &argc, &argv );
       // Get rank and size
       MPI_Comm_rank( MPI_COMM_WORLD, &my_rank );
       MPI Comm_size( MPI_COMM_WORLD, &p );
       MPI_Get_processor_name( my_name, &my_name_len );
       if ( my rank != 0 )
               // Create the message
               sprintf( message, "Greetings from process %d!", my rank );
               // Send the message
               dest = 0;
               MPI Send( message, strlen(message)+1, MPI CHAR,
                       dest, tag, MPI COMM WORLD );
       else
       {
               for( source = 1; source < p; source++ )</pre>
                       MPI Recv( message, 100, MPI CHAR, source,
                               tag, MPI COMM WORLD, &status );
                       printf( "%s\n", message );
               }
       }
       // Print the closing message
       printf( "Process %d of %d on %s done\n", my_rank, p, my_name );
       MPI Finalize();
}
```