# Iowa High Performance Computing Summer School 2013 Getting Online

Welcome to the fifth annual Iowa High Performance Computing Summer School, May 20–22, 2013. I have compiled these notes as a reference to help you get online on Helium, the 3508-core shared cluster here at the University of Iowa, and on Moffett, the 4536-core SiCortex 5832 at the Rosen Center for Advanced Computing at Purdue University.

#### 1 ITS Computer Training Room Workstations

For this course, we will be using the Macs in the ITS Computer Training Room, 2523 UCC. We use the Macs, rather than Windows machines, because the Mac OS X operating system is based on Unix, and therefore is a good platform for running Xwindows to connect remotely to the computer cluster. Here I outline how to get set up on the local workstation for easy use during this course.

- 1. If the computer at your workstation displays a Windows username and password input, tap the "Scroll Lock" key twice rapidly to toggle to the Mac.
- 2. Log onto the workstation using your HawkID and password.
- 3. Open a Finder window by clicking on the Finder (face icon) at the left hand side of the Dock at the bottom of the screen.
- 4. Navigate to the Applications/Utilities/ directory and double click on X11 to start Xwindows. This will bring up an xterm terminal window which we will use to connect to the remote machines.
- 5. Useful software on the Mac workstations:
  - (a) Emacs: This is a very useful editor for writing code, among other things. It can be found in the Applications directory.

*NOTE*: We have had problems with this application hanging up. If, after starting **Emacs** from the Applications directory, a window does not appear, follow these steps: (1) In the **xterm** window type **ps aux** | grep Emacs

The second number after your username is the process ID, pid. (2) Issue the following command to kill Emacs: kill pid

where pid is the integer number from the previous command. (3) Then launch Emacs again from the Applications directory, and it should work.

- (b) gnuplot: Handy plotting program. Can be launched from the xterm window by typing gnuplot.
- (c) Mission Control: Virtual desktop for the Mac operating system. Can be found in the dock at the bottom of the screen in the position second from the left side. If you press and hold the option key, a tab appears at the right that allows you to add virtual desktops. When not in the Mission Control view, you can navigate left or right from one virtual desktop to another using the key combinations control-→ and control-→.

### 2 Logging onto and Running Parallel Programs on Helium

The directions in this section will get you logged onto and running on Helium.

- 1. Helium is a 3508-core shared cluster computer on the University of Iowa campus. The 359 compute nodes consist of the following configuration:
  - (A) 200 compute nodes of: Dual Quad Core Intel(R) Xeon(R) CPU X5550 @ 2.67GHz Processors 24 Gb DDR3 1333MHz Memory 1Tb Storage Most of the interconnect is Infiniband, although some of the nodes are connected only by Gigabit Ethernet.
    (B) 159 compute nodes of:

Dual Hex-Core Intel(R) Xeon(R) Processors 24 Gb DDR3 Memory 1Tb Storage

- 2. Online documentation for Helium can be found at https://www.icts.uiowa.edu/confluence/display/ICTSit/User+Documentation
- 3. Access to Helium uses Active Directory and requires your current HawkID and password. Accounts on Helium have been set up for you for use during this course.
- 4. To get connected to Helium, be sure you have Xwindows running on your machine and pull up an xterm window. Connect to helium using ssh,

ssh -X username@helium.hpc.uiowa.edu

where username is your HawkID, and enter your *password* at the prompt. This will put you into your home directory on Helium. Note that the -X option enables Xforwarding so that you can pull up an Xwindow of an application running on the remote machine (Helium) on the monitor of your local machine.

Note that if you are connecting to Helium from off-campus (not within the uiowa.edu domain), you must specify that you want to connect on port 40 instead of the default port 22, using ssh -p 40 -X username@helium.hpc.uiowa.edu

- 5. Module Set Up: The first time, and only the first time, that we log onto Helium, we need to set up our .bashrc file so that the software modules that we need are loaded automatically.
  - (a) To view the modules that are already loaded, use the command module list, and to see available modules, use module avail.
  - (b) To add the necessary lines to the .bashrc file, in your home directory, open the .bashrc file using emacs in the background using emacs .bashrc &
    A window will open showing the contents of your .bashrc file. At the end of the file, after then line # User specific aliases and functions add the following two lines module load intel\_11.1.072 module load openmpi\_intel\_1.4.3 Next hit CTRL-x CTRL-s to save the changes, and CTRL-x CTRL-c to quit emacs.
  - (c) Now you may exit your login shell and follow the previous instruction to log in via **ssh** again. Once you have logged on again, you may use **module list** to verify that the necessary modules are loaded.
- 6. We are going to copy an example parallel code, HYDRO, to Helium using scp. Follow the instructions below:

- (a) Create a directory named hydro in your home directory on Helium mkdir hydro
- (b) Go to the IHPC 2013 website at http://www.physics.uiowa.edu/~ghowes/teach/ihpc13/index.html and follow the Examples link. Download the tar file of HYDRO, hyd120605.tar, to a directory on your local machine.
- (c) Open a new xterm window on your machine and navigate to the directory in which you just put hyd120605.tar.
- (d) Now, we will copy this tar file over to your home directory on Helium using scp scp hyd120605.tar username@helium.hpc.uiowa.edu:~/hydro/ and enter your password at the prompt.
  Note that if you are off-campus, specification of port 40 uses -P 40, so use scp -P 40 hyd120605.tar username@helium.hpc.uiowa.edu:~/hydro/
- (e) In the window on Helium, go into the hydro directory and unpack the tar file tar -xvf hyd120605.tar
- 7. Compiling the parallel code HYDRO:
  - (a) The tar file for HYDRO contains a Makefile that allows for easy compilation of the code on different platforms (different computers). HYDRO is written in Fortran90 using MPI for parallelization.
  - (b) Compile the code by typing make This will produce an executable hydro.e
- 8. Running parallel programs on Helium
  - (a) Helium uses the Sun Grid Engine (SGE) as a scheduler for parallel jobs. More information can be found online at https://www.icts.uiowa.edu/confluence/display/ICTSit/Basic+Job+Submission
  - (b) The examples below will use HYDRO with the sample input file sample1.in. The code requires the first argument after the executable to be the input file, thus the command will be hydro.e sample1.in
  - (c) We can choose to run either interactively or in batch mode. Interactive runs generally run immediately (if resources are available), whereas running in batch mode puts the job into a queue to be run when resources become available. As we write our first parallel codes today, we will generally run in interactive mode, since we want the results right away (and since we have resources reserved for this course). Typically, when running your codes at a national supercomputing center, you will almost exclusively run in batch mode.
  - (d) BATCH MODE: The usual method for running on a shared cluster or at a national supercomputing center is to run in batch mode, submitting your jobs using a script.
    - i. An example shell script for running on Helium is included in the tar archive, sample1\_helium.sh, for submitting a batch job to SGE to run HYDRO (for the input file sample1.in) follows: #!/bin/sh

```
#$ -q IHPC
#$ -pe 12cpn 24
#$ -l h_rt=00:10:00
#$ -N sample1
#$ -o sample1.log
#$ -j y
#$ -V
#$ -V
#$ -cwd
```

# Job Submission script

```
#$ -1 ib=1
```

```
echo "Job begin:"'date'
echo "Run sample1 on Helium, 24 proc"
mpirun -n 24 hydro.e sample1.in
echo "Job end:"'date'
```

ii. Each of the options on the lines above specifies a different aspect of the run:
-q IHPC specifies the IHPC training queue for the job (another option is all.q)
-1 h\_rt=00:10:00 specifies the time limit in HH:MM:SS format

-pe 12cpn 24 requests 24 cores using 12 cores per node

- -N sample1 specifies the name name of the job
- -o sample1.log specifies the name of the file to send the standard output
- -j y merges the stdout and stderr output,
- -V imports environmental variables to the parallel SGE environment
- -cwd places the output files in the current working directory.
- -l ib=1 requests Infiniband nodes only

The echo lines above simply write to the log file a few useful comments, but are not necessary.

- iii. To submit the job, use
- qsub sample1\_helium.sh
- iv. You may then check to see that the job is in the queue or running using qstat -u username. This will produce output that looks like

job-ID prior name user state submit/start at queue slots ja-task-ID

736050 0.50944 sample1 ghowes r 05/31/2012 21:59:58 IHPC@compute-6-177.local 16 The job-ID in this example is 736050.

v. If you are having to wait a long time for your job to start, you can change the processor request line in the script above to

-pe orte 24 (requests any 24 cores using OpenMPI)

This command will take any 24 available cores, but since those cores may be on many different nodes, performance will likely suffer.

- vi. To delete a job, submit a qdel command followed by the job-ID qdel 736050
- (e) INTERACTIVE MODE: The qlogin command is used to create an interactive session via SGE. Unlike many schedulers, using qlogin with SGE is unusually complicated, requiring the user to perform a number of steps that are normally handled by the scheduler for batch jobs. More details about this can be found in the online documentation at

https://www.icts.uiowa.edu/confluence/display/ICTSit/Qlogin+for+interactive+sessions The most important issue here is that it is up to the user to ensure that any MPI job started using qlogin employs *only* the hosts allocated by SGE. Therefore, it is very important to follow the directions below carefully to set up an interactive session using qlogin:

i. Begin an interactive session using 16 cores using the command qlogin -l ib=1 -pe orte 16 -q IHPC -l h\_rt=02:00:00 The options for this commmand specify the following: -l ib=1 requests Infiniband nodes -pe orte 16 requests 16 cores using OpenMPI -q IHPC employs nodes in the IHPC training queue (all.q is another option) -l h\_rt=01:00:00 specifies an interactive session time in hh:mm:ss

- ii. The qlogin command will produce a result something like Your job 734954 ("QLOGIN") has been submitted waiting for interactive job to be scheduled ... Your interactive job 734954 has been successfully scheduled. Establishing builtin session to host compute-3-63.local ... As long as cores are available, you should not have to wait very long. The two important pieces of information here are the job number (734954) on line 2 and the masterq host (compute-3-63.local) on the last line.
- iii. You may query the schedule to see only your own running jobs using the qstat command, qstat which will produce output something like the following, qstat job-ID prior name user state submit/start at queue slots ja-task-ID

734954 0.50948 QLOGIN ghowes r 05/31/2012 14:19:15 all.q@compute-3-63.local 16 Here you can see that your interactive session is running on the queue IHPC using compute-3-63.local as the masterq host.

- iv. To see the hosts that have been assigned to your interactive job using SGE, you can use the following command, cat /opt/gridengine/default/spool/compute-3-63/active\_jobs/734954.1/pe\_hostfile where you replace 734954 with your job number from line 2 and compute-3-63 with your masterq host. A sample of the output follows: compute-3-63.local 8 all.q@compute-3-63.local UNDEFINED compute-3-64.local 8 all.q@compute-3-64.local UNDEFINED
- v. The tricky part about using qlogin for an interactive session is that you need to specify the hosts in a way that OpenMPI will understand. To do so, you can use the command cat /opt/gridengine/default/spool/compute-3-63/active\_jobs/734954.1/pe\_hostfile | awk '{print \$1,"slots="\$2}' > hostfile to create the file hostfile which consists of the following lines: compute-3-63.local slots=8 compute-3-64.local slots=8
- vi. Now that you have everything set up, you are ready to perform and interactive run using the mpirun command. First, however, you will need to navigate to the directory that contains the parallel code that you want to run (note that launching qlogin will have landed you back in your home directory, which is probably not the directory in which you want to work). Thus, change directory to /hydro cd hydro
- vii. Run the test case sample1.in using the following command: mpirun -n 16 hydro.e sample1.in In this case, because you are running interactively, the output of HYDRO for this run will be sent to standard output (the screen).
- viii. IMPORTANT: When you are finished running your interactive session, you should exit so that you free up the reserved cores for other jobs exit

# 3 Logging onto and Running Parallel Programs on Moffett

The directions in this section will get you logged onto and running on Moffett.

1. Moffett is a 4536-core SiCortex 5832 located at the Rosen Center for Advanced Computing at Purdue University. You will be set up with a guest account on this machine during your participation in this course. Your guest account *username* and *password* will be given to you on paper. Detailed instructions on the use of Moffett can be found online at

http://www.rcac.purdue.edu/userinfo/resources/moffett/newuser.cfm

2. To get connected to Moffett, be sure you have Xwindows running on your machine and pull up an xterm window. Connect to moffett using ssh,

ssh -X username@moffett.rcac.purdue.edu

and enter your *password* at the prompt. This will put you into your home directory on Moffett. Note that the  $-\mathbf{X}$  enables X forwarding so that you can pull up an Xwindow of an application running on the remote machine (Moffett) on the monitor of your local machine.

- 3. We are going to copy and example parallel code, HYDRO, to Moffett using scp. Follow the instructions below:
  - (a) Create a directory named hydro in your home directory on Moffett mkdir hydro
  - (b) Go to the IHPC 2013 website at http://www.physics.uiowa.edu/~ghowes/teach/ihpc13/index.html and follow the Examples link. Download the tar file of HYDRO, hyd120605.tar, to a directory on your local machine.
  - (c) Open a new xterm window on your machine and navigate to the directory in which you just put hyd120605.tar.
  - (d) Now, we will copy this tar file over to Moffett using scp scp hyd120605.tar username@moffett.rcac.purdue.edu:~/hydro/ and enter your password at the prompt.
  - (e) In the window on Moffett, go into the hydro directory and unpack the tar file tar -xvf hyd120605.tar
- 4. Compiling the parallel code HYDRO:
  - (a) The tar file for HYDRO contains a Makefile that will allows for easy compilation of the code on different platforms (different computers). This code is written in Fortran90 using MPI for parallelization.
  - (b) First, open up the Makefile using emacs running in the background emacs Makefile &
    Edit the Makefile by changing the SYSTEM options variable SYSTEM=SiCortex and save the changes (using CTRL-x CTRL-s).
  - (c) Compile the code by typing make This will produce an executable hydro.e
- 5. Running parallel programs on Moffett
  - (a) Unlike many parallel computers which use Portable Batch System, or PBS, for job scheduling, Moffett uses the Simple Linux Utility for Resource Management, or SLURM. Although the syntax of the commands differs between these two systems, in general the operation is similar. Here we will give you a few of the basics on running parallel codes using SLURM on Moffett. Much more detail can be found online at http://www.rcac.purdue.edu/userinfo/resources/moffett/newuser.cfm

- (b) The examples below will use HYDRO with the sample input file sample1.in. The code requires the first argument after the executable to be the input file, thus the command will be hydro.e sample1.in
- (c) We can choose to run either interactively or in batch mode. Interactive runs generally run immediately (if resources are available), whereas running in batch mode puts the job into a queue to be run when resources become available. As we write our first parallel codes today, we will generally run in interactive mode, since we want the results right away (and since we have resources reserved for this course). Typically, when running your codes at a national supercomputing center, you will almost exclusively run in batch mode.
- (d) INTERACTIVE MODE: To run interactively, the following syntax applies srun -p <partition> -n <tasks> <executable> [args]
  Here, we select the default partition on Moffett, scx-comp. The number of tasks is the number of MPI processes, or equivalently the number of cores you choose to use. In the case of running HYDRO on 16 cores, we would use srun -p scx-comp -n 16 ./hydro.e sample1.in
  In this case, because you are running interactively, the output of HYDRO for this run will be sent to standard output (the screen).
- (e) To check the queue, use the command squeue
- (f) To cancel a job that has been submitted or kill a job that is running, use scancel <jobid>
- (g) BATCH MODE: The usual method for running on a shared cluster or at a national supercomputing center is to run in batch mode, submitting your jobs using a script. The scripts for SLURM are different from those using PBS, so here we will provide an example for using SLURM on Moffett. Here is a shell script named sample1\_moffett.sh for running the sample1.in run with HYDRO using 16 cores

```
#!/bin/sh
#SBATCH -n 16
#SBATCH -t 00:10:00
#SBATCH -o sample1.log
#SBATCH -e sample1.err
#SBATCH -J sample1
#SBATCH -p scx-comp
echo Running hydro on 16 processors
srun ./hydro.e sample1.in
echo hydro run complete
```

Each of the options on the lines above specifies a different aspect of the run: -n specifies the number of cores,

-t specifies the time limit in HH:MM:SS format,

-o specifies the name of the file to send the standard output,

-e specifies the name of the file to send the error output,

-J specifies the name of the submitted job,

-p specifies the partition on which to submit the job.

The echo lines above simply write to the log file a few useful comments, but are not necessary.

To submit the job, use

sbatch sample1\_moffett.sh

You may then check to see that the job is in the queue or running using squeue.

## 4 Compiling Your Own Parallel Code

To compile an MPI code on either Helium or Moffett, you use mpif90 for Fortran 90 or mpicc for C. Examples are below.

- FORTRAN 90: If your code is name program.f90, you can compile using mpif90 -o program.e program.f90 where the option -o program.e names the resulting executable program.e (the default behavior is to name the resulting executable a.out).
- 2. C: If your code is name program.c, you can compile using mpicc -o program.e program.c where the option -o program.e names the resulting executable program.e (the default behavior is to name the resulting executable a.out).
- 3. Note that on Helium, you must be sure that the following modules are loaded intel\_11.1.072 openmpi\_intel\_1.4.3 You should have this set up in your .bashrc file, but if not you can load the modules yourself by issuing the module load commands yourself, for example, module load intel\_11.1.072.