



CCAS and SEAS Computing Facility



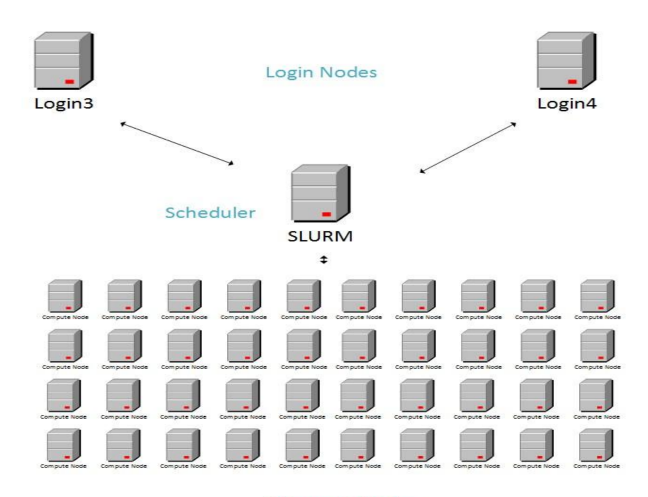
HPC Workshop 2

What we're covering:

- Review Cluster Architecture
- Working with SLURM
- Simple job submission script
- How to submit job script
- sinfo, salloc, squeue, scancel, sbatch, srun
- Tips for scripting submit files
- Reporting Job Errors



Cluster Architecture



Compute Nodes

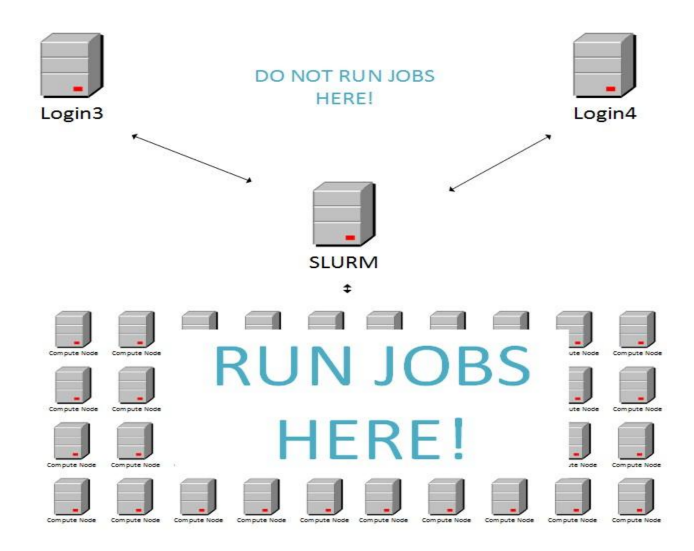


Cluster Architecture

- Login Node Server that acts as your interface to the cluster
- Scheduler Server that schedules jobs
- Compute Nodes Servers that run jobs

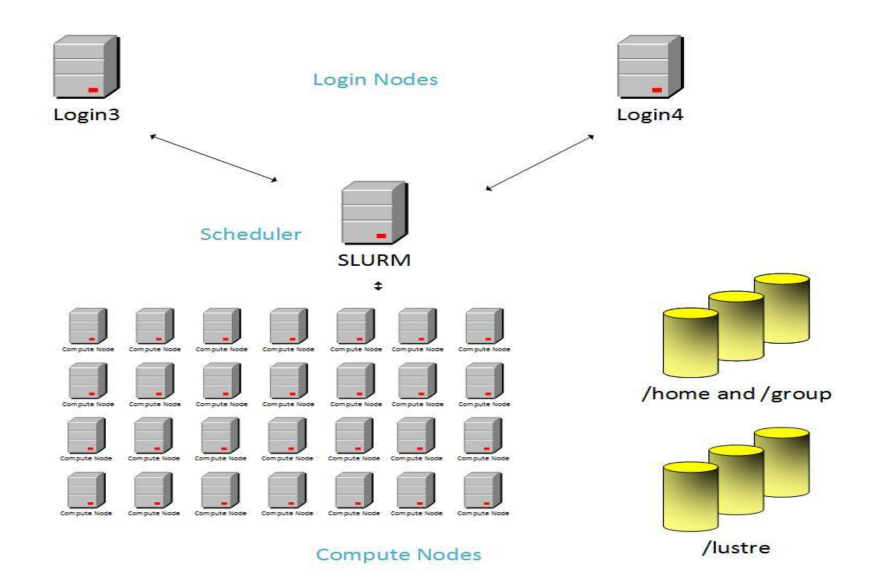


Cluster Architecture



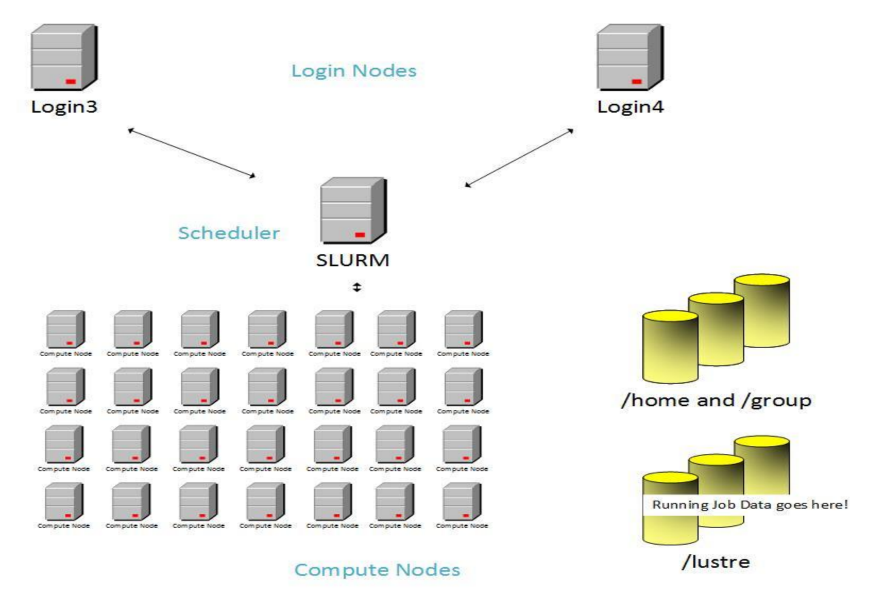


Cluster Architecture (Storage)





Cluster Architecture (Storage)





Working with SLURM

Overview: Slurm is an open source, fault-tolerant, and highly scalable cluster management and job scheduling system for large and small Linux clusters.

- Slurm requires no kernel modifications for its operation and is relatively self-contained. As a cluster workload manager, Slurm has three key functions:
 - First, it allocates exclusive access to resources (compute nodes) for users for some duration of time so they can perform work.
 - Second, it provides a framework for starting, executing, and monitoring work (normally a parallel job) on the set of allocated nodes.
 - Finally, it arbitrates contention for resources by managing a queue of pending work.



Working with SLURM

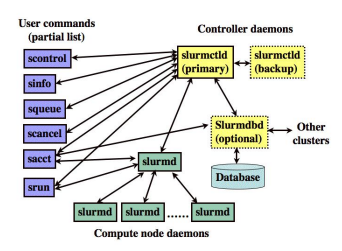
Why use SLURM?

- SLURM allows jobs to be scheduled so a user does not have to wait until a node is free to begin work. SLURM does the waiting for you!
- SLURM allows resources to be prioritized for groups that purchase shares on Colonial One.
- SLURM enables efficient use of the cluster since it constantly monitors resources in use and schedules jobs on unallocated resources as they free up
- SLURM runs a job based on a submit script.
- A submit script calls your job script to execute your calculations.



Architecture I

As depicted in Figure 1, Slurm consists of a **slurmd** daemon running on each compute node and a central **slurmctld** daemon running on a management node (with optional fail-over twin). The **slurmd** daemons provide fault-tolerant hierarchical communications. The user commands include: **sacct**, **salloc**, **sattach**, **sbatch**, **sbcast**, **scancel**, **scontrol**, **sinfo**, **smap**, **squeue**, and **srun**. All of the commands can run anywhere in the cluster.





Architecture II

The entities managed by these Slurm daemons, shown in Figure 2, include **nodes**, the compute resource in Slurm, **partitions**, which group nodes into logical (possibly overlapping) sets, **jobs**, or allocations of resources assigned to a user for a specified amount of time, and **job steps**, which are sets of (possibly parallel) tasks within a job. The partitions can be considered job queues, each of which has an assortment of constraints such as job size limit, job time limit, users permitted to use it, etc. Priority-ordered jobs are allocated nodes within a partition until the resources (nodes, processors, memory, etc.) within that partition are exhausted. Once a job is assigned a set of nodes, the user is able to initiate parallel work in the form of job steps in any configuration within the allocation. For instance, a single job step may be started that utilizes all nodes allocated to the job, or several job steps may independently use a portion of the allocation.

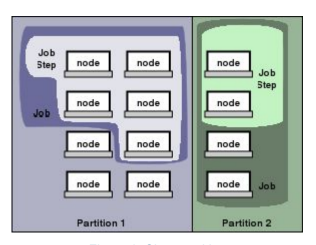


Figure 2. Slurm entities



Software, Modules & Login

- Common software is available to all users via the "module" system
- Can I install my own software?
 - Yes, provided it runs from your home/group directory
 - Users cannot use a package manager (yum) to install software
 - Colonial One users do not have root on Colonial One

Login to Colonial One using your GW NetID/email address & password

\$ ssh <netid>@login.colonialone.gwu.edu



sinfo

- lists node and partition information for the cluster
- useful for finding unallocated nodes

Example: type "sinfo" at the prompt

```
TIMELIMIT
PARTITION
             AVAIL
                               NODES
                                       STATE NODELIST
defq*
                up 14-00:00:0
                                 128
                                       alloc node[033-160]
short
                up 2-00:00:00
                                       alloc node[097-191]
                up 14-00:00:0
128qb
                                       alloc node[041-064]
256qb
                up 14-00:00:0
                                       alloc node[033-040]
                                       alloc node901
2tb
                up 14-00:00:0
                up 7-00:00:00
                                       alloc node[003-020,029-032]
gpu
qpu
                up 7-00:00:00
                                   10
                                        idle node[001-002,021-028] ← Free nodes!
                up 7-00:00:00
                                   22
                                       alloc node[003-020,029-032]
gpu-noecc
                up 7-00:00:00
                                        idle node[001-002,021-028] ← Free nodes!
gpu-noecc
                                   10
                up 7-00:00:00
                                   21
                                        idle node[333-353] ← Free nodes!
ivyqpu
                up 7-00:00:00
                                        idle node[333-353] ← Free nodes!
ivyqpu-noecc
                                   21
allqpu-noecc
                up 7-00:00:00
                                   22
                                       alloc node[003-020,029-032]
allqpu-noecc
                up 7-00:00:00
                                   31
                                        idle node[001-002,021-028,333-353] \leftarrow Free
nodes!
                                       alloc node[991-992]
debua
                      4:00:00
                up
debug-cpu
                    4:00:00
                                       alloc node992
                up
                     4:00:00
debug-gpu
                                       alloc node991
                up
```



salloc - Obtain a Slurm job allocation (a set of nodes), execute a command, and then release the allocation when the command is finished. You can use salloc to run interactive jobs:

```
salloc -N 1 -p ivygpu -t 5
```

srun - Run a parallel job on cluster managed by Slurm. Use srun to identify your allocated nodes after running salloc:

```
srun hostname #or
squeue -u <netid>
ssh nodename
```



squeue - view information about jobs located in the Slurm scheduling queue.

List jobs for your user account:

squeue -u <username>

Estimate when a job will start:

squeue -u username --start

List by job status:

squeue -u username -t RUNNING or PENDING



SLURM job status

Check Status of Job by User

```
[user@login4 ~]$ squeue -u <username>

JOBID PARTITION NAME USER ST TIME NODES NODELIST(REASON)

2325403 defq job_OpenMP.sh user PD 0:00 1 (Resources)
```

Jobs typically pass through several states in the course of their execution. The typical states are PENDING, RUNNING, SUSPENDED, COMPLETING, and COMPLETED. An explanation of each state follows.

```
CA = CANCELLED
```

CD = COMPLETED

CG = COMPLETING

 \mathbf{F} = FAILED

NF = NODE_FAIL

PD = PENDING

 \mathbf{R} = RUNNING

S = SUSPENDED

TO = TIMEOUT



sbatch - Submit a batch script to Slurm

scancel - used to signal or cancel jobs, job arrays or job steps.

sinfo - view information about Slurm nodes and partitions.



SLURM COMMANDS (cont.)

- salloc Obtain a Slurm job allocation (a set of nodes), execute a command,
 and then release the allocation when the command is finished.
- squeue view information about jobs located in the Slurm scheduling queue.
- scancel Used to signal jobs or job steps that are under the control of Slurm.
- sbatch Submit a batch script to Slurm.
- srun Run parallel jobs



hello-mpi.py

This small Python program will list an mpi process number and the node it is running on. Copy the text into a document called "hello-mpi.py" and save it.

```
#hello.py
from mpi4py import MPI
import socket

comm = MPI.COMM_WORLD
rank = comm.Get_rank()
size = MPI.COMM_WORLD.Get_size()
thishost = socket.gethostname()

print "hello world from process ", rank, " of ", size, "
on ", thishost < ----- THIS IS ALL ONE LINE</pre>
```



How to submit job script I

Load the current MPI module so we can run our program correctly:

```
module load openmpi/1.8/gcc/4.9.2/cpu module load python/2.7.6
```

Check your home directory to make sure your compiled program is there:

```
ls -la hello*
```



How to submit job script I

Create a SLURM script using an editor such as vi or emacs using steps 1 through 3. The script (or file) can be called anything you want but should end in .sh (i.e. submit.sh).

Step 1: Resource Specification

```
[user@login4 ~]$ nano submit.sh

#!/bin/sh
#SBATCH --time 5:00
#SBATCH -o testing%j.out
#SBATCH -e testing%j.err
#SBATCH -p defq -n 8
#SBATCH --mail-user=<username>@gwu.edu
#SBATCH --mail-type=ALL
module load openmpi/1.8/gcc/4.9.2/cpu
module load python/2.7.6
mpirun -n 8 python /home/<username>/hello.py

Step 2: Submit job
module load slurm
[user@login4 ~]$ sbatch submit.sh
Submitted batch job 2325403
```



How to submit job script I

SLURM will email you when you job has finished. Once the job is finished, SLURM will place a .out file and a .err file (if there are errors) in the directory with your submit script.

```
$> ls slurm*
slurm-2333627.out slurm-2333631.out slurm-2333633.out
slurm-2333629.out slurm-2333636.out
```

The .out file is the result of your job. Cat or less the file to show you the results.

```
$> cat slurm-2333636.out
Process 1 on node991.cm.cluster out of 8
Process 3 on node991.cm.cluster out of 8
Process 7 on node991.cm.cluster out of 8
Process 5 on node991.cm.cluster out of 8
Process 4 on node991.cm.cluster out of 8
Process 6 on node991.cm.cluster out of 8
Process 0 on node991.cm.cluster out of 8
Process 2 on node991.cm.cluster out of 8
```



Reporting Job Errors

- Email to hpchelp@gwu.edu
- Include .err file
- Include submit script
- Include what modules you loaded