

Introduction to HPC Using the New Cluster (Sapelo) at GACRC

Georgia Advanced Computing Resource Center

University of Georgia

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Outline

- What is GACRC?
- What is the new cluster (Sapelo) at GACRC?
- How does Sapelo operate?
- How to work with Sapelo?

What is GACRC?

Who Are We?

- Georgia **A**dvanced **C**omputing **R**esource **C**enter
- Collaboration between the Office of Vice President for Research (**OVPR**) and the Office of the Vice President for Information Technology (**OVPIT**)
- Guided by a faculty advisory committee (GACRC-AC)

Why Are We Here?

- To provide computing hardware and network infrastructure in support of high-performance computing (**HPC**) at UGA

Where Are We?

- <http://gacrc.uga.edu> (Web) <http://wiki.gacrc.uga.edu> (Wiki)
- <http://gacrc.uga.edu/help/> (Web Help)
- https://wiki.gacrc.uga.edu/wiki/Getting_Help (Wiki Help)

GACRC Users September 2015

Colleges & Schools	Depts	PIs	Users
Franklin College of Arts and Sciences	14	117	661
College of Agricultural & Environmental Sciences	9	29	128
College of Engineering	1	12	33
School of Forestry & Natural Resources	1	12	31
College of Veterinary Medicine	4	12	29
College of Public Health	2	8	28
College of Education	2	5	20
Terry College of Business	3	5	10
School of Ecology	1	8	22
School of Public and International Affairs	1	3	3
College of Pharmacy	2	3	5
	40	214	970
Centers & Institutes	9	19	59
TOTALS:	49	233	1029

GACRC Users September 2015

Centers & Institutes	PIs	Users
Center for Applied Isotope Study	1	1
Center for Computational Quantum Chemistry	3	10
Complex Carbohydrate Research Center	6	28
Georgia Genomics Facility	1	5
Institute of Bioinformatics	1	1
Savannah River Ecology Laboratory	3	9
Skidaway Institute of Oceanography	2	2
Center for Family Research	1	1
Carl Vinson Institute of Government	1	2
	19	59

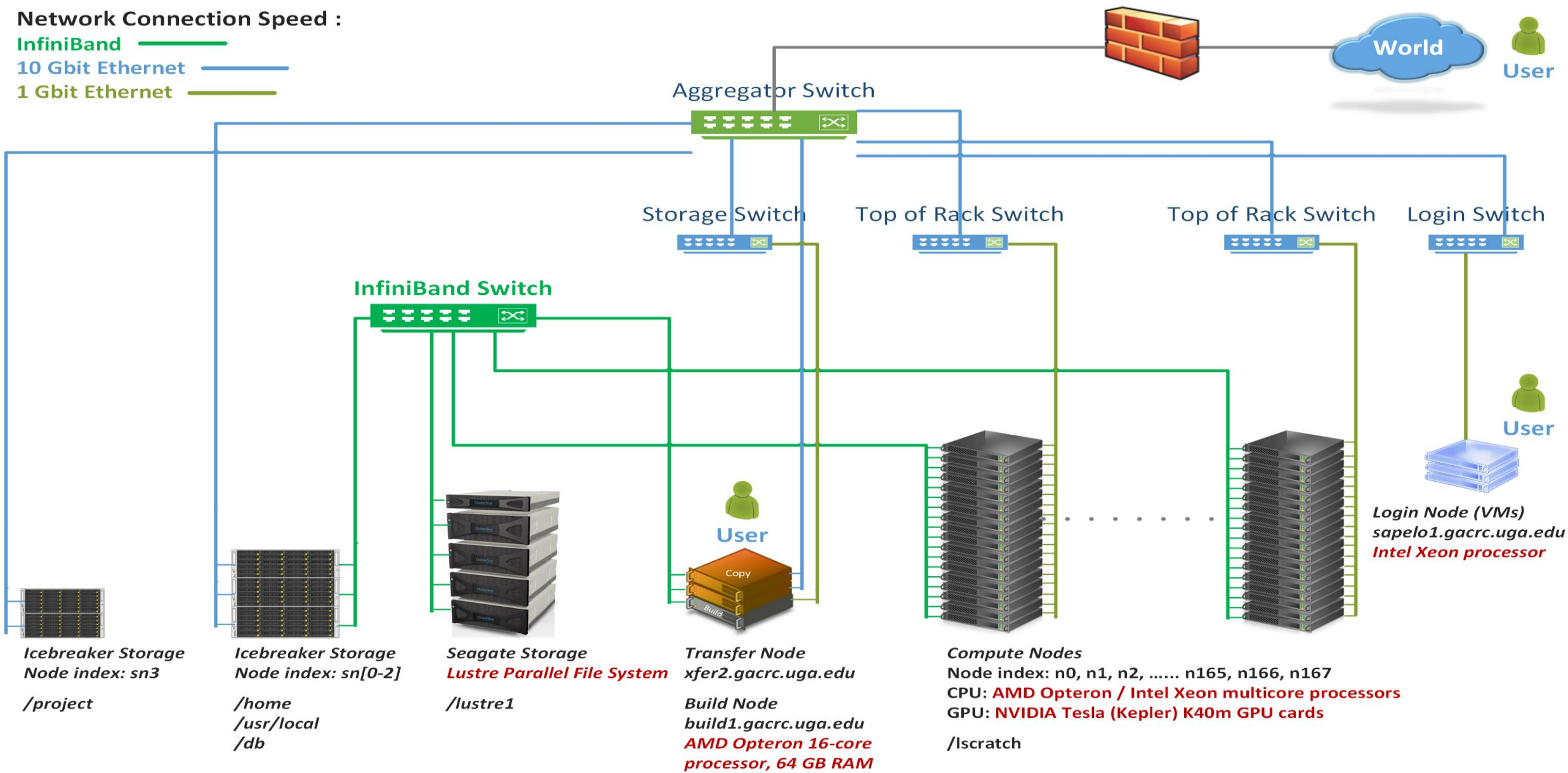
What is the new cluster (Sapelo) at GACRC?

- Cluster Structural Diagram
- General Information
- Computing Resources
- Storage Environment

The New GACRC Linux HPC Cluster Structural Diagram

Network Connection Speed :

- InfiniBand** —
- 10 Gbit Ethernet** —
- 1 Gbit Ethernet** —



Icebreaker Storage
Node index: sn3
/project

Icebreaker Storage
Node index: sn[0-2]
/home
/usr/local
/db

Seagate Storage
Lustre Parallel File System
/lustre1


Transfer Node
xfer2.gacrc.uga.edu
Build Node
build1.gacrc.uga.edu
AMD Opteron 16-core processor, 64 GB RAM

Compute Nodes
Node index: n0, n1, n2, n165, n166, n167
CPU: AMD Opteron / Intel Xeon multicore processors
GPU: NVIDIA Tesla (Kepler) K40m GPU cards
/lscratch

Login Node (VMs)
sapelo1.gacrc.uga.edu
Intel Xeon processor

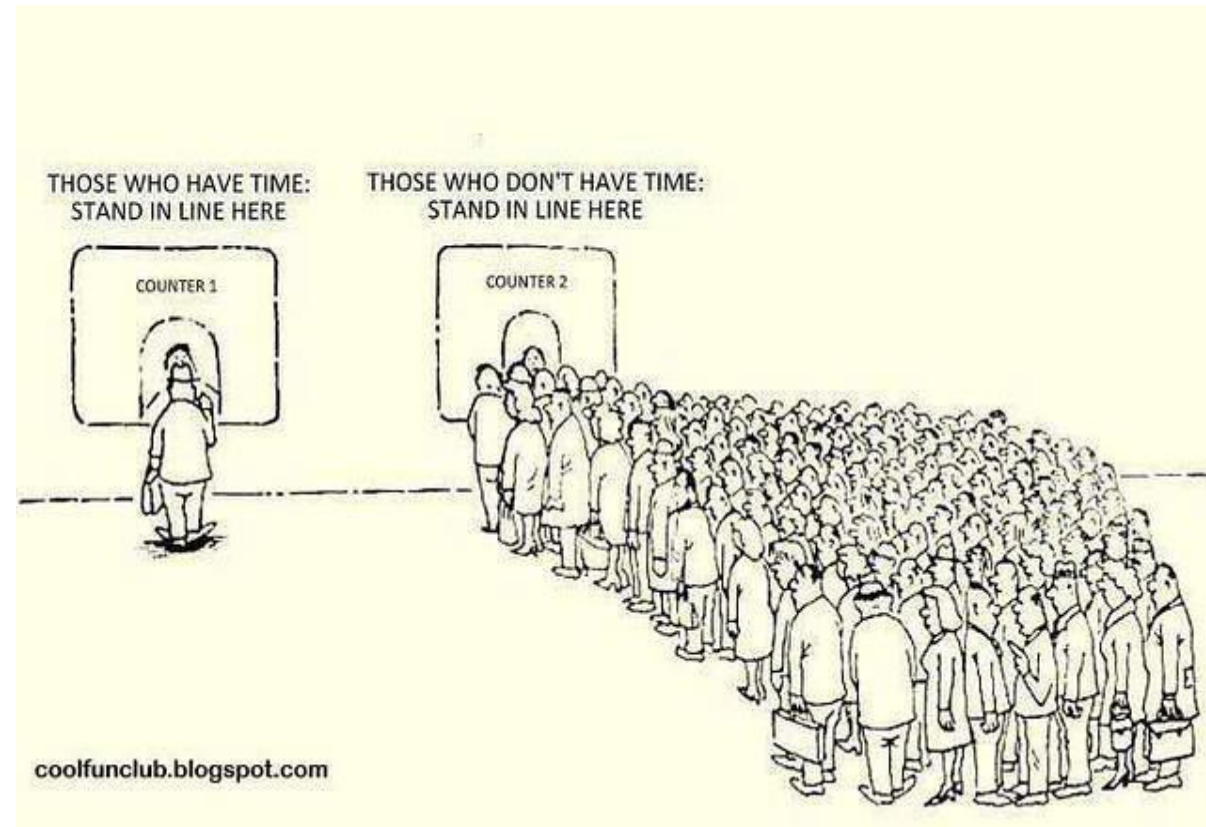
What is the new cluster – General Information

Sapelo is a Linux high performance computing (HPC) cluster:

- OS: 64-bit CentOS 6.5
- You can login to:
 - Login (username@sapelo1.gacrc.uga.edu)** : login, edit script, submit jobs
 - Transfer (username@xfer2.gacrc.uga.edu)** : transfer, compress data
 - Build (username@build1.gacrc.uga.edu)** : compile, test
- **Login**  **Interactive Node** : run interactively, edit script, submit jobs
- Communication: **InfiniBand network**
 compute nodes ↔ compute nodes ↔ storage systems, e.g., /home and /lustre1

What is the new cluster – General Information

- Batch-queueing System:
 - Jobs can be started (submitted), monitored, and controlled
 - Determine which compute node is the best place to run a job
 - Determine appropriate execution priority for a job to run
- Sapelo:
 - Torque Resource Manager
 - Moab Workload Manager



What is the new cluster – Computing Resources

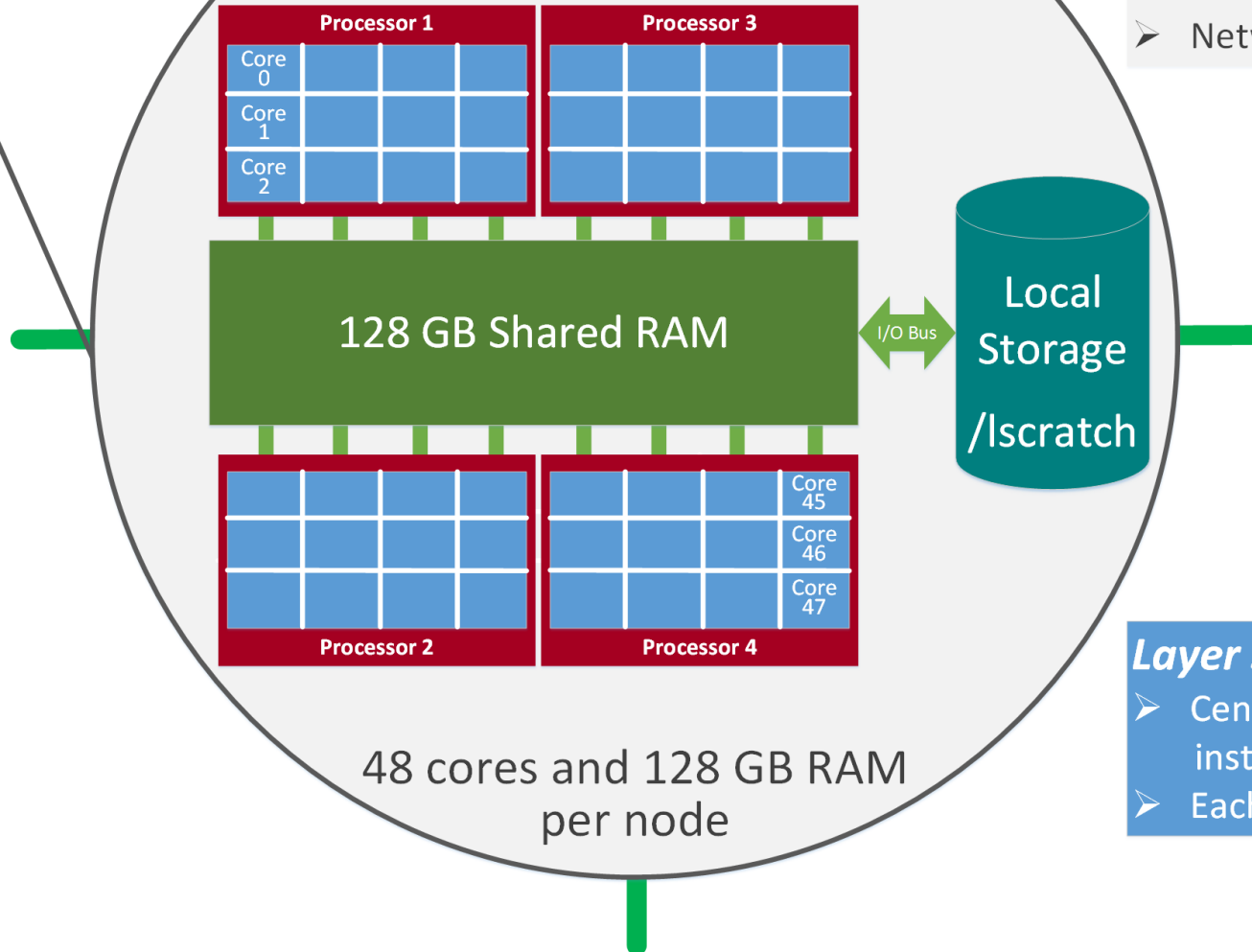
Queue	Node	Total	Processor	CPUs /Node	RAM (GB) /Node	Max RAM (GB) /Single-node Job	GPU	GPU Cards /Node	InfiniBand
batch	AMD	120	AMD Opteron	48	128	126	N/A	N/A	Yes
	HIGHMEM	3	AMD Opteron	48	512 (2)	504	N/A	N/A	Yes
					1024 (1)	997			
	GPU	2	Intel Xeon	16	128	126	NVIDIA K40m	8	Yes
abcnode (buy-in)	2	AMD Opteron	48	256	252	N/A	N/A	Yes	

Peak Performance per Node: 500Gflops/Node

Home: /home/username: **100GB**

Global scratch: /lustre1/username: **NO quota limit, auto-moved to /project if no modification in 30 days!**

Node 167



48 cores and 128 GB RAM
per node

Layer 1: Node

- A standalone “computer in a box”
- Multiple processors, e.g. 4, sharing memory
- Local disk storage, network interface, etc.
- Networked into a cluster

Layer 2: Processor

- A single computing component
- Multicore processor, e.g. 12 cores

Layer 3: Core

- Central processing unit (CPU) reading and executing instructions independently
- Each core is assigned to a software thread

What is the new cluster – Storage Environment

4 Filesystems	Role	Quota	Accessible from	Intended Use	Notes
/home/username	Home	100GB	sapelo1.gacrc.uga.edu (Login) Interactive nodes (Interactive) xfer2.gacrc.uga.edu (Transfer) build1.gacrc.uga.edu (Build) compute nodes (Compute)	Highly static data being used frequently	Snapshots
/lustre1/username	Global Scratch	No Limit	Interactive nodes (Interactive) xfer2.gacrc.uga.edu (Transfer) compute nodes (Compute)	Temporarily storing large data being used by jobs	Auto-moved to /project if 30 days no modification
/lscratch/username	Local Scratch	250GB	Individual compute node	Jobs with heavy disk I/O	User to clean up
/project/abclab	Storage	Variable	xfer2.gacrc.uga.edu (Transfer)	Long-term data storage	Group sharing possible

Note: /usr/local/apps : Software installation directory
 /db : Bioinformatics database installation directory

What is the new cluster – Storage Environment

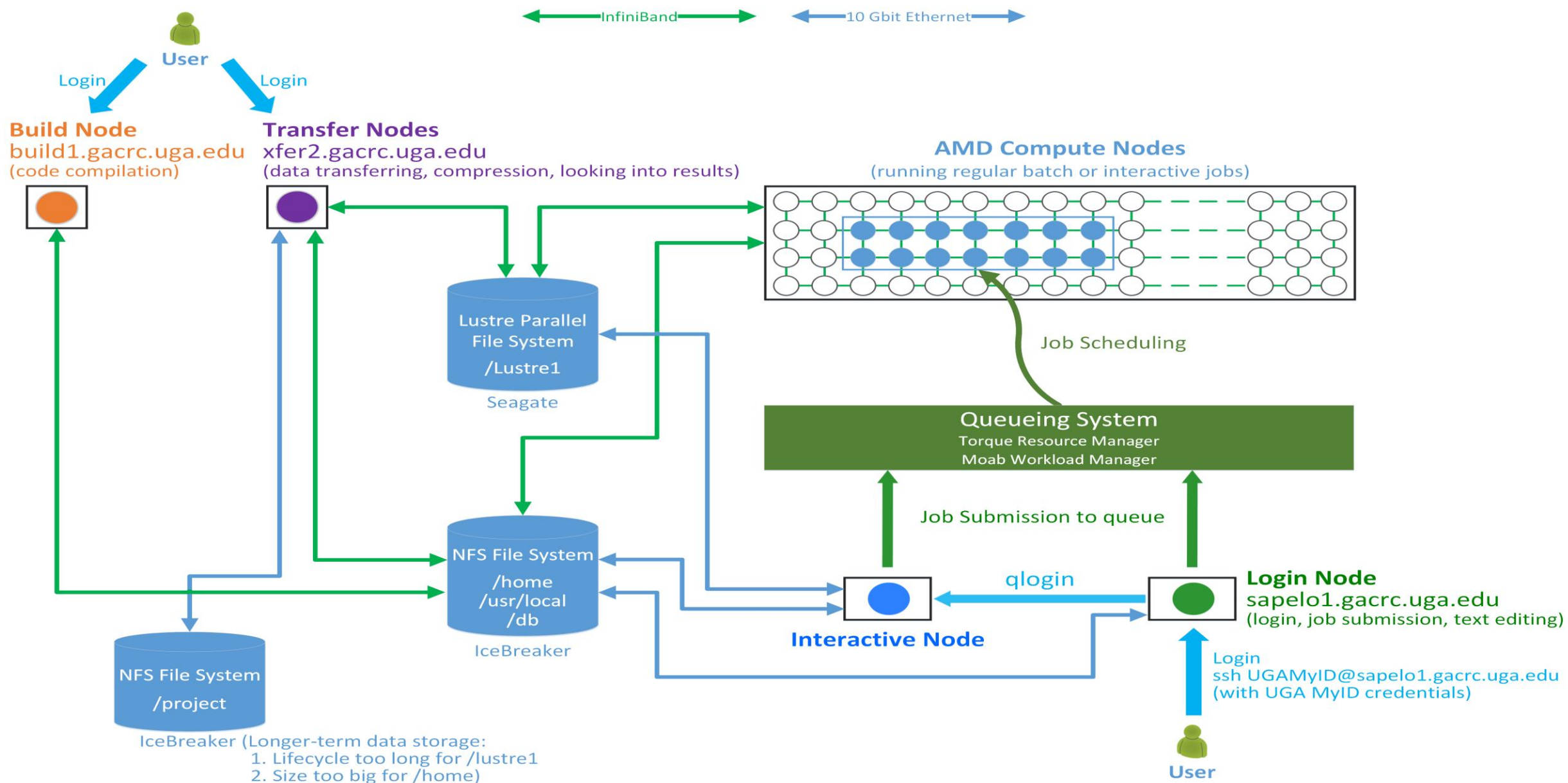
7 Main Functions	On/From-Node	Related Filesystem
Login Landing	Login or Transfer or Build	/home/username (Home) (Always!)
Batch Job Submitting	Login	/home/username (Home)
	Interactive	/lustre1/username (Scratch) (Suggested!) /home/username (Home)
Interactive Job Running	Interactive	/lustre1/username (Scratch) /home/username (Home)
Data Transferring, Archiving , and Compressing	Transfer	/lustre1/username (Scratch) /home/username (Home)
Job Data Temporarily Storing	Compute	/lscratch/username (Local Scratch) /lustre1/username (Scratch)
Long-term Data Storing	Transfer or Copy	/project/abclab
Code Compilation	Build	/home/username (Home)

How does Sapelo operate?

Next Page



The New GACRC Linux HPC Cluster Operational Diagram



IceBreaker (Longer-term data storage:
 1. Lifecycle too long for /lustre1
 2. Size too big for /home)

How to work with Sapelo?

Before we start:

- To get zcluster to be your best HPC buddy

GACRC Wiki: <http://wiki.gacrc.uga.edu>

GACRC Support: https://wiki.gacrc.uga.edu/wiki/Getting_Help

How to work with Sapelo?

To submit your ticket to GACRC?

➤ **Job Troubleshooting:**

Please tell us details of your question or problem, including but not limited to:

- ✓ Your user name
- ✓ Your job ID
- ✓ Your working directory
- ✓ The queue name and command you used to submit the job

➤ **Software Installation:**

- ✓ Specific name and version of the software
- ✓ Download website
- ✓ Supporting package information if have

Note:

It's **USER's** responsibility to make sure the **correctness of datasets** being used by jobs!



How to work with Sapelo?

- On cluster, you are not alone..... Each user is sharing finite resources, e.g., CPU cycles, RAM, disk storage, network bandwidth, with other researchers. *What you do may affect other researchers on the cluster.*

6 rules of thumb to remember:


- **NO jobs running on login node**
- NO multi-threaded job running with only 1 core requested
- NO large memory job running on regular nodes
- NO long job running on interactive node
- NO small memory job running on large memory nodes
- Use the copy node for file transfer and compression



How to work with Sapelo?

- Start with the Cluster
- Connect & Login
- Software Packages
- Run Jobs
 - How to submit a job
 - How to make a job submission script
 - How to check job status, cancel a job, etc.

How to work with Sapelo – Start with the Cluster

- You need a **User Account**: **UGAMyID@sapelo1.gacrc.uga.edu**
To create your account correctly, you must provide us with your **official MyID**,  not a UGA MyID alias!
- To get a user account, follow **4** steps:

1. New user training (<http://gacrc.uga.edu/help/training/>)
2. Tell us your **Name**, **UGA MyID**, **Lab name** and **PI's name**, via GACRC Support (https://wiki.gacrc.uga.edu/wiki/Getting_Help)
3. **Invitation letter** with instructions to start account initialization
4. **Welcome letter** with whole package of information about your account created successfully

How to work with Sapelo – Connect & Login

- Open a connection: Open a terminal and `ssh` to your account

```
ssh zhuofei@sapelo1.gacrc.uga.edu
```

or

```
ssh -X zhuofei@sapelo1.gacrc.uga.edu
```

⁽¹⁾ `-X` is for X windows application running on the cluster to be forwarded to your local machine

⁽²⁾ If using Windows, use **SSH client** to open connection, get from UGA download software page)

- Logging in: You will be prompted for your **UGA MyID password**

```
zhuofei@sapelo1.gacrc.uga.edu's password: █
```

⁽³⁾ On Linux/Mac, when you type in the password, the prompt blinks and does not move)

- Logging out: `exit` to leave the system

```
[zhuofei@75-104 ~]$ exit
```

How to work with Sapelo – Software Packages

- The cluster uses **environment modules** to define the various paths for software packages
- Current number of modules installed is ~200 and expanding daily!
- **module avail** to list all modules available on the cluster:

```
[zhuofei@75-104 ~]$ module avail
```

----- /usr/local/modulefiles -----				
Core/StdEnv	exabayes/1.4.1	java/jdk1.8.0_20	openmpi/1.6.5/gcc/4.4.7	rsem/latest
Data/cache/moduleT.new	examl/3.0.11	java/latest (D)	openmpi/1.6.5/pgi/14.9	rsem/1.2.20 (D)
Data/cache/moduleT (D)	expat/latest	lammps/5Sep14	openmpi/1.8.3/gcc/4.4.7	samtools/latest
Data/system.txt	expat/2.0.1 (D)	lammps/16Aug13 (D)	openmpi/1.8.3/gcc/4.7.4	samtools/0.1.19
R/3.1.2	fastqc/latest	moab/7.2.10	openmpi/1.8.3/gcc/4.8.0 (D)	samtools/1.1
bedops/latest	fastqc/0.11.3 (D)	moab/8.1.1 (D)	openmpi/1.8.3/intel/14.0	samtools/1.2 (D)
bedops/2.4.14 (D)	gcc/4.7.4	moabs/1.3.2	openmpi/1.8.3/intel/15.0.2 (D)	scripture/latest
boost/1.47.0/gcc447	gcc/4.8.0 (D)	mvapich2/2.0.0/gcc/4.4.7	openmpi/1.8.3/pgi/14.9	scripture/03202015 (D)
boost/1.57.0/gcc447	gmap-gsnap/latest	mvapich2/2.0.0/pgi/14.9	orca/3.0.3	sparsehash/latest
boost/1.57.0_thread/gcc447	gmap-gsnap/2014-12-24 (D)	ncbiblast+/2.2.29	perl/latest	sparsehash/2.0.2 (D)
bowtie/latest	gnuplot/5.0.0	netcdf/3.6.3/gcc/4.4.7	perl/5.20.1	tophat/latest
bowtie/1.1.1 (D)	gsl/1.16/gcc/4.4.7	netcdf/3.6.3/intel/14.0	perl/5.20.2 (D)	tophat/2.0.13 (D)
bowtie2/latest	hdf5/1.8.14/gcc/4.4.7	netcdf/3.6.3/intel/15.0.2 (D)	pgi/14.9	trinity/latest
bowtie2/2.2.4 (D)	hdf5/1.8.14/intel/15.0.2	netcdf/4.1.3/gcc/4.4.7	pgi/14.10 (D)	trinity/r20140717
cuda/5.0.35/gcc/4.4.7	hdf5/1.8.14/pgi/14.9	netcdf/4.1.3/intel/15.0.2	python/2.7.8-ucs4	trinity/2.0.6 (D)
cuda/6.5.14/gcc/4.4.7	imb/3.2	netcdf/4.1.3/pgi/14.10	python/2.7.8	zlib/gcc447/1.2.8
cufflinks/latest	intel/14.0	netcdf/4.3.2/gcc/4.4.7	python/3.4.3 (D)	
cufflinks/2.2.1 (D)	intel/15.0.2 (D)	netcdf/4.3.2/pgi/14.9	raxml/8.1.20	

How to work with Sapelo – Software Packages

- `module list` to list which modules currently loaded:

```
[zhuofei@75-104 ~]$ module list
Currently Loaded Modules:
  1) StdEnv   2) moab/7.2.10
```

- `module load` to load the needed modules:

```
[zhuofei@75-104 ~]$ module load ncbiblast+/2.2.29
[zhuofei@75-104 ~]$ module load python/2.7.8
[zhuofei@75-104 ~]$ module load R/3.1.2
[zhuofei@75-104 ~]$ module list
Currently Loaded Modules:
  1) StdEnv   2) moab/7.2.10   3) ncbiblast+/2.2.29   4) python/2.7.8   5) R/3.1.2
```

- `module unload` to remove the specific module:

```
[zhuofei@75-104 ~]$ module unload R/3.1.2
[zhuofei@75-104 ~]$ module list
Currently Loaded Modules:
  1) StdEnv   2) moab/7.2.10   3) ncbiblast+/2.2.29   4) python/2.7.8
```

How to work with Sapelo – Run Jobs

- Components you need to run a job:
 - **Software** already loaded. If not, used `module load`
 - **Job submission script** to run the software, and specify computing resources:
 - ✓ Number of nodes and cores
 - ✓ Amount of memory
 - ✓ Type of nodes
 - ✓ Maximum wallclock time, etc.
- Common commands you need:
 - `qsub, qdel`
 - `qstat -f, showjobs, showq` etc.

How to work with Sapelo – Run Jobs

- How to submit a job? **Easy!**

```
[zhuofei@75-104 MPIs]$ qsub sub.sh
```

qsub is to
submit a job

sub.sh is your **job submission script**
specifying:

- ✓ Number of nodes and cores
- ✓ Amount of memory
- ✓ Type of nodes
- ✓ Maximum wallclock time, etc.

- How to make a job submission script? **Next Page!**

How to work with Sapelo – Run Jobs

- Example 1: **Serial job script** *sub.sh* running NCBI Blast +

<code>#PBS -S /bin/bash</code>	→ Linux shell (bash)
<code>#PBS -q batch</code>	→ Queue name (batch)
<code>#PBS -N testBlast</code>	→ Name of the job (testBlast)
<code>#PBS -l nodes=1:ppn=1:AMD</code>	→ Number of nodes (1), number of cores/node (1), node type (AMD)
<code>#PBS -l mem=20gb</code>	→ Maximum amount of physical memory (20 GB) used by the job
<code>#PBS -l walltime=48:00:00</code>	→ Maximum wall clock time (48 hours) for the job, default 6 minutes
 <code>cd \$PBS_O_WORKDIR</code>	 → Use the directory from which the job is submitted as the working directory
 <code>module load ncbiblast+/2.2.29</code>	 → Load the module of ncbiblast+, version 2.2.29
 <code>time blastn [options] > outputfile</code>	 → Run blastn with ‘time’ command to measure the amount of time it takes to run the application

How to work with Sapelo – Run Jobs

- Example 2: **Threaded job script** *sub.sh* running NCBI Blast + with **4** threads

```

#PBS -S /bin/bash
#PBS -q batch
#PBS -N testBlast
#PBS -l nodes=1:ppn=4:AMD
#PBS -l walltime=480:00:00
#PBS -l mem=20gb

#PBS -M jSmith@uga.edu
#PBS -m ae
#PBS -j oe

cd $PBS_O_WORKDIR

module load ncbiblast+/2.2.29

time blastn -num_threads 4 [options] > outputfile

```

→ Number of nodes (**1**), number of cores/node (**4**), node type (**AMD**)
Number of threads (4) = Number of cores requested (4)

- Email address to receive a notification for computing resources
- Send email notification when job aborts (**a**) or terminates (**e**)
- Standard error file (**testBlast.e1234**) will be merged into standard out file (**testBlast.o1234**)

→ Run blastn with 4 threads (**-num_threads 4**)

How to work with Sapelo – Run Jobs

- Example 3: **MPI job script** *sub.sh* running RAxML with **50** MPI processes

```
#PBS -S /bin/bash
```

```
#PBS -q batch
```

```
#PBS -N testRAxML
```

```
#PBS -l nodes=2:ppn=48:AMD
```

→ Number of nodes (2), number of cores/node (48), node type (AMD)

```
#PBS -l walltime=480:00:00
```

Total cores requested = $2 \times 48 = 96$

```
#PBS -l mem=20gb
```

We suggest, Number of MPI Processes (50) ≤ Number of cores requested (96)

```
#PBS -j oe
```

```
cd $PBS_O_WORKDIR
```

```
module load raxml/8.1.20
```

→ To run raxmlHPC-MPI-AVX, MPI version using OpenMPI 1.8.3/Intel 15.0.2

```
module load intel/15.0.2
```

```
module load openmpi/1.8.3/intel/15.0.2
```



```
mpirun -np 50 raxmlHPC-MPI-AVX [options] > outputfile
```

→ Run raxmlHPC-MPI-AVX with 50 MPI processes (**-np 50**)

```
#PBS -S /bin/bash
#PBS -q batch
#PBS -N testRAxML
#PBS -l nodes=2:ppn=27:AMD
#PBS -l walltime=480:00:00
#PBS -l mem=20gb
#PBS -j oe
```

→ ppn number (27) fewer than 48 MUST be a multiplier of 3!

```
cd $PBS_O_WORKDIR
```

```
# Context Sharing
CONTEXTS=$(/usr/local/bin/set_contexts.sh $PBS_NUM_PPN)
if [[ "$?" -eq "0" ]]; then
    export PSM_SHAREDCONTEXTS_MAX=$CONTEXTS
fi
```

} New lines copied from GACRC Wiki

```
module load raxml/8.1.20
module load intel/15.0.2
module load openmpi/1.8.3/intel/15.0.2
```

```
mpirun -np 50 raxmlHPC-MPI-AVX [options] > outputfile → Run raxmlHPC-MPI-AVX with 50 MPI processes (-np 50)
```

How to work with Sapelo – Run Jobs

- How to check job status? **qstat**

```
[jSmith@75-104 MPIs]$ qstat
Job ID          Name           User           Time Use  S Queue
-----
481929.pbs      testJob1       jSmith         900:58:0  C batch
481931.pbs      testJob2       jSmith         04:00:03  R batch
481934.pbs      testJob3       jSmith         0         Q batch

Job status:
R : job is running
C : job completed (or crashed) and is not longer running. Jobs stay in this state for 1h
Q : job is pending, waiting for resources to become available
```

- How to cancel *testJob3* with jobID 481934? **qdel**

```
[zhuofei@75-104 MPIs]$ qdel 481934
[jSmith@75-104 MPIs]$ qstat
Job ID          Name           User           Time Use  S Queue
-----
481929.pbs      testJob1       jSmith         900:58:0  C batch
481931.pbs      testJob2       jSmith         04:00:03  R batch
481934.pbs      testJob3       jSmith         0         C batch
```

← Stay on list 1 hr

How to work with Sapelo – Run Jobs

- How to check computing resources?

Option 1: `qstat -f JobID` for *running jobs* or *finished jobs in 1 hour*

Option 2: `showjobs JobID` for *finished jobs over 1 hour, but ≤ 7 days*

Option 3: Email notification from *finished jobs (completed, canceled, or crashed)*,
if using:

```
#PBS -M jSmith@uga.edu
```

```
#PBS -m ae
```

How to work with it – Run Jobs

- **qstat -f JobID** for *running jobs* or *finished jobs in 1 hour*

```
[zhuofei@75-104 MPIs]$ qstat -f 699847
Job Id: 699847.pbs.scn
  Job_Name = testJob
  Job_Owner = zhuofei@uga-2f0f976.scn
resources_used.cput = 00:11:55
resources_used.energy_used = 0
resources_used.mem = 411572kb
resources_used.vmem = 6548528kb
resources_used.walltime = 00:01:36
job_state = C
queue = batch
.
Error_Path = uga-2f0f976.scn:/home/zhuofei/MPIs/testJob.e699847
exec_host = n165/0-23
Output_Path = uga-2f0f976.scn:/home/zhuofei/MPIs/testJob.o699847
.
Resource_List.mem = 5gb
Resource_List.nodect = 1
Resource_List.nodes = 1:ppn=24:AMD
Resource_List.walltime = 10:00:00
.
Variable_List = PBS_O_QUEUE=batch,PBS_O_HOME=/home/zhuofei, ..... ,
                PBS_O_WORKDIR=/home/zhuofei/MPIs,
```


How to work with it – Run Jobs

- **showjobs JobID** for *finished jobs over 1 hour, but \leq 7 days*

```
[zhuofei@75-104 MPIs]$ showjobs 699847
Job Id       : 699847.pbs.scn
Job Name     : testJob
Output File  : uga-2f0f976.scn:/home/zhuofei/MPIs/testJob.o699847
Error File   : uga-2f0f976.scn:/home/zhuofei/MPIs/testJob.e699847
Working Directory : /home/zhuofei/MPIs
Home Directory : /home/zhuofei
Submit Arguments : sub.sh
User Name    : zhuofei
Group Name   : rccstaff
Queue Name   : batch
Wallclock Limit : 10:00:00
Walloclock Duration: 00:01:36
CPUTime      : 00:11:55
Memory Used   : 401.9Mb
Memory Limit  : 5gb
vmem Used    : 6.2Gb
Submit Time   : Wed Nov  4 12:02:22 2015
Start Time    : Wed Nov  4 12:03:41 2015
End Time      : Wed Nov  4 12:04:45 2015
Exit Code     : 0
Master Host   : n165
```

How to work with it – Run Jobs

- Email notification from *finished jobs* (**completed**, **canceled**, or **crashed**)

```

PBS Job Id: 700009.pbs.scm
Job Name:   testJob
Exec host:  n1/4-27
Execution terminated
Exit_status=0
resources_used.cput=00:05:12
resources_used.energy_used=0
resources_used.mem=410984kb
resources_used.vmem=6548516kb
resources_used.walltime=00:00:59
Error_Path: uga-
2f0f976.scm:/home/zhuofei/MPIs/testJob.o700009
Output_Path: uga-
2f0f976.scm:/home/zhuofei/MPIs/testJob.o700009
    
```

```

PBS Job Id: 700097.pbs.scm
Job Name:   testJob
Exec host:  n1/4-27
Execution terminated
Exit_status=271
resources_used.cput=00:11:22
resources_used.energy_used=0
resources_used.mem=412304kb
resources_used.vmem=6548524kb
resources_used.walltime=00:00:41
Error_Path: uga-
2f0f976.scm:/home/zhuofei/MPIs/testJob.o700097
Output_Path: uga-
2f0f976.scm:/home/zhuofei/MPIs/testJob.o700097
    
```

How to work with it – Run Jobs

- How to check queue status?
showq

```
[zhuofei@75-104 MPIs]$ showq
active jobs-----
JOBID                USERNAME           STATE  PROCS   REMAINING          STARTTIME
481914                brant              Running  1      20:46:21   Fri Jun 12 11:32:23
481915                brant              Running  1      20:48:56   Fri Jun 12 11:34:58
481567                becton             Running 288    2:04:15:48 Wed Jun 10 15:01:50
481857                kkim               Running  48     9:18:21:41 Fri Jun 12 09:07:43
481859                kkim               Running  48     9:18:42:21 Fri Jun 12 09:28:23
.
108 active jobs      5141 of 5740 processors in use by local jobs (89.56%)
                    121 of 122 nodes active          (99.18%)
eligible jobs-----
JOBID                USERNAME           STATE  PROCS   WCLIMIT          QUEUETIME
481821                joykai             Idle   48     50:00:00:00     Thu Jun 11 13:41:20
481813                joykai             Idle   48     50:00:00:00     Thu Jun 11 13:41:19
481811                joykai             Idle   48     50:00:00:00     Thu Jun 11 13:41:19
481825                joykai             Idle   48     50:00:00:00     Thu Jun 11 13:41:20
.
50 eligible jobs
blocked jobs-----
JOBID                USERNAME           STATE  PROCS   WCLIMIT          QUEUETIME
0 blocked jobs
Total jobs: 158
```

Thank You!