

Introduction to HPC Using zcluster at GACRC

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Outline

- What is GACRC?
- What is HPC Concept?
- What is zcluster?
- How does zcluster operate?
- How to work with zcluster?



What is GACRC?

Who Are We?

- Georgia Advanced Computing Resource Center
- 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 highperformance computing (HPC) at UGA

Where Are We?

http://gacrc.uga.edu (Web)

http://wiki.gacrc.uga.edu (Wiki)

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



GACRC Users September 2015

Colleges & Schools	Depts	Pls	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	Pls	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



Concept of High Performance Computing (HPC)



- ✓ Serial problem can not be broken
- ✓ *Discrete* instructions executed *sequentially*
- Only 1 instruction executed at any moment on a single processor
- Problem broken into *parallel* parts can be solved *concurrently*
- ✓ Instructions executed *simultaneously* on *multiply* processors
- ✓ Synchronization/communication employed
- Shared-memory multithreaded job or MPI job (Message Passing Interface)



What is zcluster?

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





What is zcluster – General Information

GACRC zcluster is a Linux high performance computing (HPC) cluster:

- Operating System: 64-bit Red Hat Enterprise Linux 5 (RHEL 5)
- Login Node: zcluster.rcc.uga.edu
 Interactive Node: compute-14-7/9 Copy Node: copy.rcc.uga.edu
- Internodal Communication: 1Gbit network compute nodes \Leftrightarrow compute nodes compute nodes \Leftrightarrow storage systems

qlogin



What is zcluster – 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
- On zcluster: Sun Grid Engine (SGE)





What is zcluster – Computing Resources

	Queue Type	Queue Name	Nodes	Processor	Cores/Node	RAM(GB)/Node	Cores	NVIDIA GPU	
		45		12	48	540	NI / A		
	Regular	100-300	150	inter keon	8	16	1200	N/A	
		rec m128 20d	4		8	192	32		
	High Memory rcc-m128-30d rcc-m512-30d	10	Intel Xeon	12	256	120	N/A		
		rcc-m512-30d	2		32	512	64		
	Multi Core	rcc-mc-30d	6	AMD Opteron	32	64	192	N/A	
	Interactive	interq	2	AMD Opteron	48	132	96	N/A	
		rcc-sgpu-30d	2		8	48	16	4 Tesla S1070 cards	
	GPU	rcc-mgpu-30d	2	Intel Xeon	12	48	24	9 Tesla (Fermi) M2070 cards	
		rcc-kgpu-30d	4		12	96	24	32 Tesla (Kepler) K20Xm cards	

Total peak performance: 23 Tflops

ÎGACRC





- **Home directory** → */home/groupname/username*
 - Mounted and visible on all nodes, with a quota of ~100GB
 - Any directory on /home has snapshot backups
 - > Taken once a day, and maintained 4 daily ones and 1 weekly one
 - Name: .snapshot, e.g., /home/abclab/jsmith/.snapshot
 - Completely invisible, however, user can "cd" into it and then "ls":

zhuofei@zcluster:~\$ ls -a							
b	ash_profile .e	emacs.d	.fontconfig	.maple_history	MPIs	scripts	test.sh
b	ashrc .E	ENV_file	.gnuplot_history	.Mathematica	openMPs	serials	.viminfo
.bash_history do	wnloads ex	ke	.history	.mc	.profile	sht	.Xauthority
.bash_logout .e	emacs , f	flexlmrc	.lesshst	.mozilla	Pthreads	.ssh 🗲	.snapshot is NOT
zhuofei@zcluster:	~\$ cd .snapshot	🗩 🗲 can '	"cd" into .snapshot				shown here!
zhuofei@zcluster:~/.snapshot\$ ls 🗧 🗲 then "Is" to list its contents							
2015.06.21.00.00.	01.weekly 2015	5.06.27.01	.00.01.daily 20	15.06.28.01.00.0	1.daily 2	015.06.30.	01.00.01.daily
2015.06.26.01.00.	01.daily 2015	5.06.28.00	.00.01.weekly 20	15.06.29.01.00.0	1.daily		



- Local scratch → /lscratch/username
 - ➢ On local disk of each compute node → node-local storage
 - rcc-30d 8-core nodes: ~18GB, rcc-30d 12-core nodes: ~370GB
 - No snapshot backup
 - Usage Suggestion: If your job writes results to /lscratch, job submission script should move the data to your home or escratch before exit
- **Ephemeral Scratch** → /escratch4/zhuofei_Jun_22
 - > Create with make escratch command
 - Visible to all nodes with a quota of 4TB
 - No snapshot backup
 - > To be deleted after **37 days**



Filesystem	Role	Quota	Accessible from	Intended Use	Notes
/home/abclab/username	Home	100GB	zcluster.rcc.uga.edu (Login)	Highly static data being used frequently	Snapshots
/escratch4/username	Scratch	4TB	copy.rcc.uga.edu (Copy) Interactive nodes (Interactive) compute nodes (Compute)	Temporarily storing large data being used by jobs	Auto-deleted in 37 days
/lscratch/username	Local Scratch	18 ~ 370GB	Individual compute node	Jobs with heavy disk I/O	User to clean up
/project/abclab	Storage	Variable	copy.rcc.uga.edu (Copy)	Long-term data storage	Group sharing possible

- Note: 1. /usr/local : Software installation directory
 - /db : bioinformatics database installation directory
 - 2. To login to Interactive nodes, use qlogin from Login node



6 Main Function	On/From-Node	Related Filesystem
Login Landing	Login or Copy	/home/abclab/username (Home) (Always!)
Batch Job Submitting	Login or Interactive	/escratch4/username (Scratch) (<mark>Suggested!</mark>) /home/abclab/username (Home)
Interactive Job Running	Interactive	/escratch4/username (Scratch) /home/abclab/username (Home)
Data Archiving , Compressing and Transferring	Copy or Transfer	/escratch4/username (Scratch) /home/abclab/username (Home)
Job Data Temporarily Storing	Compute	/lscratch/username (Local Scratch) /escratch4/username (Scratch)
Long-term Data Storing	Copy or Transfer	/project/abclab



How does zcluster operate?

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How to work with zcluster?

Before we start:

- To get zcluster to be your best HPC buddy, go to GACRC Wiki (<u>http://wiki.gacrc.uga.edu</u>) GACRC Web (<u>http://gacrc.uga.edu</u>)
- To get the most effective and qualified support from us, go to GACRC Support (<u>https://wiki.gacrc.uga.edu/wiki/Getting_Help</u>)
- To work happily and productively, follow the cluster's Community Code of Conduct (CCOC)



How to work with it?

• Cluster's CCOC:

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 zcluster?

- Start with zcluster
- Connect & Login
- Transfer Files
- Softwares Installed
- Run Interactive Jobs
- Submit Batch Jobs
 - > How to submit *serial, threaded,* and *MPI* batch jobs
 - > How to check job status, cancel a job, etc.



How to work with zcluster – Start with zcluster

- > You need a User Account : username@zcluster.rcc.uga.edu
- Procedure: https://wiki.gacrc.uga.edu/wiki/User_Accounts
- A UGA faculty member (PI) may register a computing lab: http://help.gacrc.uga.edu/labAcct.php
- The PI of a computing lab may request user accounts for members of his/her computing lab: http://help.gacrc.uga.edu/userAcct.php
- > User receives an email notification once the account is ready
- User can use passwd command to change initial temporary password



How to work with zcluster – Connect & Login

• Open a connection: Open a terminal and ssh to your account

ssh zhuofei@zcluster.rcc.uga.edu

or

ssh -X zhuofei@zcluster.rcc.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 zcluster password

zhuofei@zcluster.rcc.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@zcluster:~\$ exit



How to work with zcluster – Transfer Files

User's local

Copy node (copy.rcc.uga.edu) of zcluster

On Linux, Mac or cygwin on Windows : scp [Source] [Target]
 E.g. 1: On local machine, do Local

 zcluster

scp file1 username@copy.rcc.uga.edu:~/subdir

scp *.dat username@copy.rcc.uga.edu:~/subdir

E.g. 2: On local machine, do zcluster \rightarrow Local

scp username@copy.rcc.uga.edu:~/subdir/file ./

scp username@copy.rcc.uga.edu:~/subdir/*.dat ./

• On Window: FileZilla, WinSCP, etc.



How to work with zcluster – Softwares Installed

- Perl, Python, Java, awk, sed, C/C++ and Fortran compilers
- Matlab, Maple, R
- Many Bioinformatics applications: NCBI Blast+, Velvet, Trinity, TopHat, MrBayes, SoapDeNovo, Samtools, RaxML, etc.
- RCCBatchBlast (RCCBatchBlastPlus) to distribute NCBI Blast (NCBI Blast+) searches to multiple nodes.
- > Many Bioinformatics Databases: NCBI Blast, Pfam, uniprot, etc.
- For a complete list of applications installed: https://wiki.gacrc.uga.edu/wiki/Software



How to work with zcluster – Run Interactive Jobs

• To run an interactive job, you need to open a session on an interactive node using qlogin command:

- Current maximum runtime is **12** hours
- When you are done, remember to exit the session!
- Detailed information, like interactive parallel job? Go to: https://wiki.gacrc.uga.edu/wiki/Running Jobs on zcluster



How to work with zcluster – Submit Batch Jobs

- Components you need to submit a batch job:
 - Software already installed on zcluster
 - Job submission script to run the software,
 - ✓ Specifying working directory
 - Exporting environment variables, e.g.,
 OMP_NUM_THREADS (OpenMP threads number)
 LD_LIBRARY_PATH (searching paths for shared libraries)
- Common commands you need:
 - qsub with specifying queue name, threads or MPI rank number
 - > qstat, qdel
 - qacct, qsj, etc.



How to work with zcluster – Batch Serial Job

- **Step 1**: Create a job submission script *sub.sh* running Samtools:
- #!/bin/bash → Linux shell (bash)
- cd \${HOME}/testdir \rightarrow Specify and enter (cd) the working directory ($\frac{1}{1000}$

time /usr/local/samtools/latest/samtools < command> [options] \rightarrow Run samtools with 'time' command to measure amount of

- time it takes to run the application
- Step 2: Submit it to the queue:





How to work with zcluster – Batch *Threaded* Job

• **Step 1**: Create a job submission script *sub.sh* running bowtie2:





How to work with zcluster – Batch MPI Job

- **Step 1**: Create a job submission script *sub.sh* running RAxML:
- #!/bin/bash
 cd \${HOME}/testdir
 export MPIRUN=/usr/local/mpich2/1.4.1p1/gcc 4.5.3/bin/mpirun
 \$ Define and export environment variable (MPIRUN)
 for convenient usage
 \$ MPIRUN_-np \$NSLOTS /usr/local/raxml/latest/raxmlHPC-MPI-SSE3 [options] → Run RAxML with 20 MPI processes (-np \$NSLOTS)
- **Step 2**: Submit it to the queue:





How to work with zcluster – Check and Cancel Jobs

• To check the status of all queued and running jobs: qstat

qstat	ightarrow shows your job in the pool
qstat –u "*"	ightarrow shows all the jobs in the pool
qstat –j 12345	→ shows detailed information, e.g., maxymem, about the job with JOBID 12345
qstat –g t	Iist all nodes used by your jobs

- To cancel a queued or running job: qdel
 - qdel −u zhuofei→ deleted all your jobsqdel 12345→ deletes your job with JOBID 12345
- To list detailed information about a job: qsj, qacct

qsj 12345→ shows information, e.g., maxymem, about the RUNNING job with JOBID 12345qacct -j 12345→ shows information, e.g., maxymem, about the ENDED job with JOBID 12345



Thank You!