

# GACRC Sapelo2 Cluster New User Training

Georgia Advanced Computing Resource Center (GACRC)

Enterprise Information Technology Services(EITS)

The University of Georgia



#### Outline

- GACRC
- Sapelo2 Cluster
  - Cluster Diagram and Overview
  - Five Directories
  - Five Computational Partitions
  - Software Environment
- Batch Job Submission Workflow
- Useful Commands: squeue --me, sacct, sacct-gacrc, qlogin
- GACRC Wiki and User Support
- Appendices



#### **GACRC**

- A high-performance-computing (HPC) center at the UGA
- Provide to the UGA research and education community an advanced computing environment:
  - HPC computing and networking infrastructure located at the Boyd Data Center
  - Comprehensive collection of scientific, engineering and business applications
  - Consulting and training services

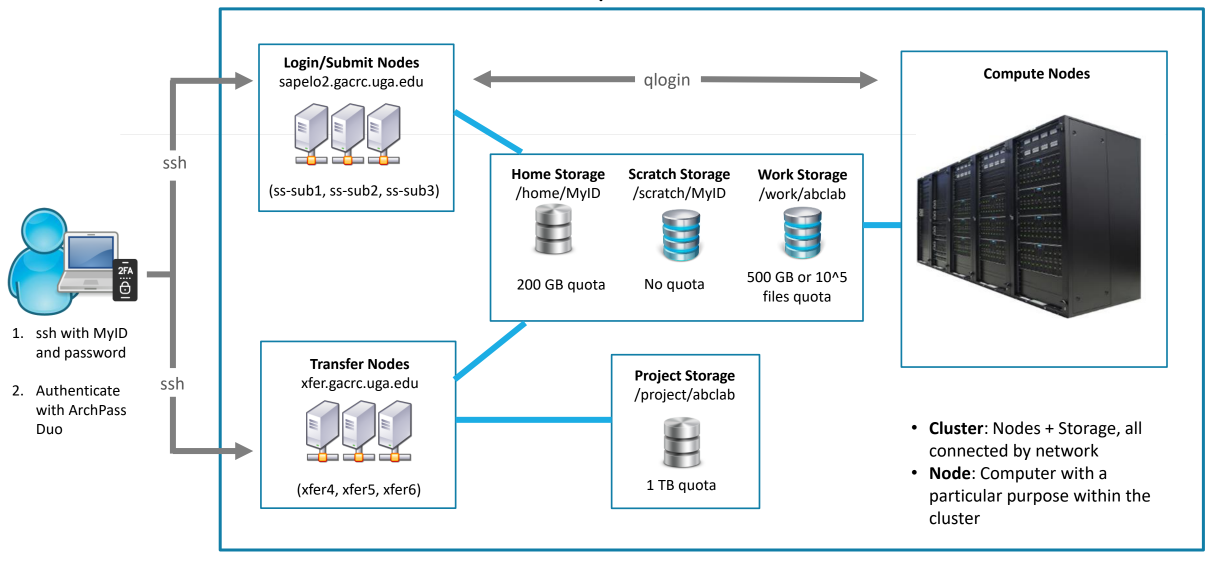
Wiki: <a href="http://wiki.gacrc.uga.edu">http://wiki.gacrc.uga.edu</a>

Help and Support: <a href="http://help.gacrc.uga.edu">http://help.gacrc.uga.edu</a>

Web Site: <a href="http://gacrc.uga.edu">http://gacrc.uga.edu</a>

Kaltura Channel: <a href="https://kaltura.uga.edu/channel/GACRC/176125031">https://kaltura.uga.edu/channel/GACRC/176125031</a>

#### Sapelo2 Cluster



Note: You need to connect to the UGA network using VPN when accessing from outside of the UGA main campus.

UGA VPN: <a href="https://eits.uga.edu/access\_and\_security/infosec/tools/vpn/">https://eits.uga.edu/access\_and\_security/infosec/tools/vpn/</a>



## Five Directories <a href="https://wiki.gacrc.uga.edu/wiki/Disk Storage">https://wiki.gacrc.uga.edu/wiki/Disk Storage</a>

Directory	Name	Quota	Accessible from	Intended Use	Backed- up	Important Notes	
/home/MyID	Home	200GB	Login Transfer Compute	Static data, e.g.  1. Scripts, source codes  2. Local software	Yes	Not for storing data of your jobs!	
/scratch/MyID	Scratch	No Limit	Login Transfer Compute	Temporary files needed for currently running jobs	No	Clean up when your job finishes! Subject to "30-day purge" policy	
/work/abclab	Work	500GB 10 <sup>5</sup> files	Login Transfer Compute	Input files needed for repeated jobs	No	Clean up when your job finishes! Group sharing is possible	
/project/abclab	Project	1TB (initial)	Transfer	Temporary data parking	Yes	Group sharing is possible	
/Iscratch	Local Scratch	200GB - 800GB	Compute	Jobs with heavy disk I/O operations	No	Clean up when job exits from node!	

### Scratch File System 30-Day Purge Policy

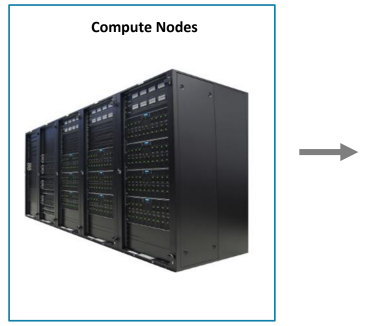
https://wiki.gacrc.uga.edu/wiki/Disk Storage#Scratch file system

Any file that is not accessed or modified by a compute job in a time period no longer than 30 days will be automatically deleted off the /scratch file system.

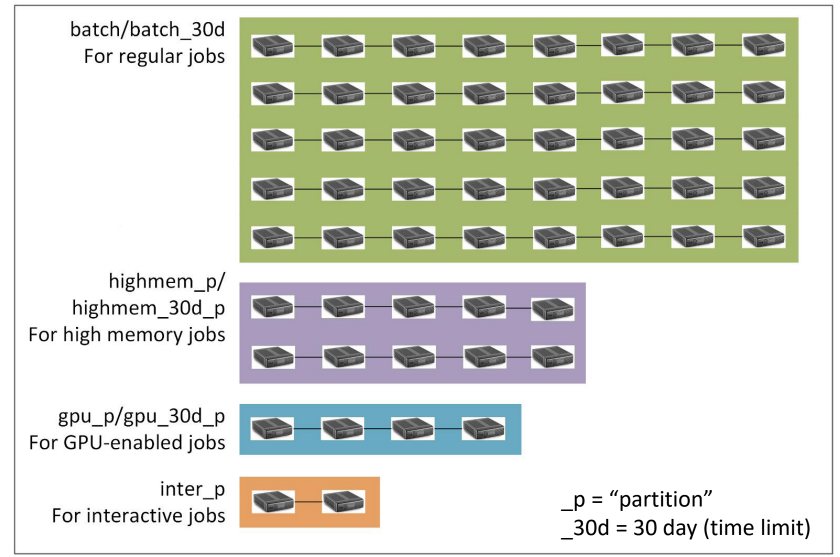
Measures circumventing this policy will be monitored and actively discouraged.

- ➤ You are suggested to copy files from /scratch to /project or outside of GACRC
- You should first move all unnecessary files and folders to /scratch/trash/\$USER
- > The fastest way to save old files is to copy them to /project area, using the fpsync utility on xfer.gacrc.uga.edu
- ➤ When you archive data using tar on /scratch, please do not use z option (compression option). After you archive data with tar, you can use gzip to compress it.

### Computational Partitions



Compute nodes are divided into groups called **partitions**. A **partition** is a collection of compute nodes for a particular computing need.



Computational Partitions <a href="https://wiki.gacrc.uga.edu/wiki/Job\_Submission">https://wiki.gacrc.uga.edu/wiki/Job\_Submission</a> partitions on Sapelo2 https://wiki.gacrc.uga.edu/wiki/Systems#Sapelo2

Туре	Partition	Time limit	Max jobs Running	Max jobs Submit	Notes		
	batch		250	10,000	Regular nodes		
Regular	highmem_p	7 days	days 15 100 For running high memory jobs		For running high memory jobs		
	gpu_p		18	20	For running GPU-enabled jobs		
	batch_30d						
Long-term	highmem_30d_p	30 days	1	2	30-day partition counterparts		
	gpu_30d_p		2				
Interactive	inter_p	2 days	3	20	Regular nodes, for interactive jobs.		
Buy-in	name_p	variable			Partitions that target different groups' buy-in nodes. The name string is specific to each group.		

Partition	Total Nodes	Max Mem(GB) /Single-node job	Cores /Node	Processor Type	GPU Cards /Node					
			64	AMD FDVC						
		120	32	AIVID LI TC						
batch batch_30d	~275		48	AMD Opteron	N/A					
		180	32 Intel Xeon Skylake							
		58	28	Intel Xeon Broadwell						
		500	32	AMD EPYC						
		300	48	AMD Opteron						
highmem_p highmem_30d_p	~30		64	AMD EPYC	N/A					
G		990	28	Intel Xeon Broadwell						
			48	AMD EPYC  AMD Opteron  Intel Xeon Skylake  Intel Xeon Broadwell  AMD EPYC  AMD Opteron  AMD EPYC  AMD Opteron  AMD EPYC  Intel Xeon Broadwell  AMD Opteron  AMD EPYC  AMD Opteron						
		180	<b>32</b>	Intel Xeon Skylake	1 NVDIA P100					
gpu_p gpu_30d_p	~10	120	16	Intol Voor						
8ba_30a_b		90	Intel Xeon 12							
name_p	variable									

#### Software Environment



#### https://wiki.gacrc.uga.edu/wiki/Software

- Approximately 900 software modules are installed (as of March 2021)
- Most modules are compiled EasyBuild (EB) toolchains GCC-8.3.0 or foss-2019b.
- Name format: Name/Version-Toolchain, e.g., Python/3.8.2-GCCcore-8.3.0 (case-sensitive)
- Module commands:
  - > ml spider pattern: Search module names matching a pattern
  - > ml moduleName: Load a module into your working environment
    - DO NOT LOAD/USE MODULES ON THE LOGIN/SUBMIT NODES! (ss-sub1, ss-sub2, ss-sub3, etc...)
  - > ml av : List all available software modules installed on cluster
  - ml : List modules currently loaded
  - > ml -moduleName: Remove a module from working environment
  - ml purge: Remove all modules from working environment



### Important Tip using Software

https://wiki.gacrc.uga.edu/wiki/Available Toolchains and Toolchain Compatibility

- When you load more than one software modules, toolchain compatibility is the most important thing
  you need to pay attention to
- If you load more than one module and some toolchains are incompatible, your job will end up with failing dependencies or Lmod errors, such as:

Lmod has detected the following error:

These module(s) exist but cannot be loaded as requested



ml Python/3.7.4-GCCcore-8.3.0 ml Perl/5.28.0-GCCcore-7.3.0



ml Python/3.7.4-GCCcore-8.3.0 ml Perl/5.30.0-GCCcore-8.3.0



ml Beast/2.6.3-foss-2019b ml Perl/5.28.0-GCCcore-7.3.0



ml Beast/2.6.3-foss-2019b ml Perl/5.30.0-GCCcore-8.3.0



#### Job Submission Workflow

https://wiki.gacrc.uga.edu/wiki/Running Jobs on Sapelo2

- 1. Log on to Login node using MyID and password, and two-factor authentication with Archpass Duo: ssh MyID@sapelo2.gacrc.uga.edu
- 2. On Login node, change directory to your scratch space: cd /scratch/MyID
- 3. Create a working subdirectory for a job: mkdir ./workDir
- 4. Change directory to workDir: cd ./workDir
- 5. Transfer data from local computer to <u>workDir</u>: use <u>scp</u> or <u>WinSCP</u> to connect Transfer node

  Transfer data on cluster to <u>workDir</u>: log on to Transfer node and then use <u>cp</u> or <u>mv</u>
- 6. Make a job submission script in workDir: nano ./sub.sh
- 7. Submit a job from workDir: sbatch ./sub.sh
- 8. Check job status: squeue --me or Cancel a job: scancel jobID



#### Step1: Log on to Login node - Mac/Linux using ssh

https://wiki.gacrc.uga.edu/wiki/Connecting

- 1. Open Terminal utility
- 2. Type command line: ssh MyID@sapelo2.gacrc.uga.edu
- 3. You will be prompted for your MyID password
- 4. Sapelo2 access requires ID verification using two-factor authentication with Archpass
  - Duo. If you are not enrolled in Archpass Duo, please refer to

https://eits.uga.edu/access and security/infosec/tools/archpass duo/ on how to enroll



### Step1 (Cont.) - Mac/Linux using ssh

Use Terminal utility on Mac or Linux!



[zhuofei@localhost ~]\$ [zhuofei@localhost ~]\$ ssh zhuofei@sapelo2.gacrc.uga.edu ← Log on Password: Input MyID password! Enter a passcode or select one of the following options: 1. Duo Push to XXX-XXX-5758 2. Phone call to XXX-XXX-5758 3. Phone call to XXX-XXX-1925 4. SMS passcodes to XXX-XXX-5758 Passcode or option (1-5): 1 Select Duo authentication option! Success. Logging you in... Last login: Tue Sep 15 11:22:42 2020 from 128.192.75.65 zhuofei@ss-sub1 ~\$ ← I am on login node ss-sub1!



### Step1 (Cont.) - Windows using PuTTY

- Download and install PuTTY: <a href="https://www.putty.org/">https://www.putty.org/</a>
- 2. Detailed downloading and installation instructions:

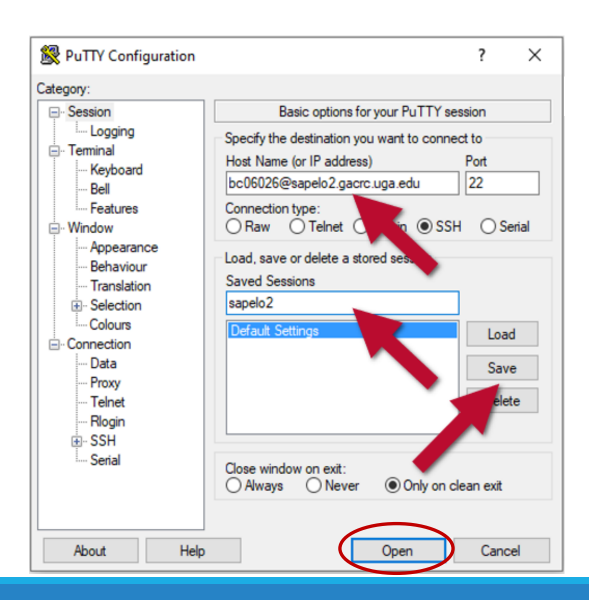
https://wiki.gacrc.uga.edu/wiki/How to Install and Configure PuTTY

3. Detailed configuring and usage instructions:

https://wiki.gacrc.uga.edu/wiki/How to Install and Configure PuTTY#Configuring PuTTY

### Step1 (Cont.) - Windows using PuTTY





The first time you connect to a login node, PuTTY will give you this security alert window. Please click "Yes" or "Accept"

Sapelo2.gacrc.uga.edu - PuTTY

PuTTY Security Alert

The server's host key is not cached in the registry. You

think it is.

connection.

Yes

have no guarantee that the server is the computer you

If you want to carry on connecting just once, without

If you do not trust this host, hit Cancel to abandon the

No

ssh-ed25519 255 e5:4b:18:93:5c:03:66:ff:39:45:cf:fb:90:95:97:db

Cancel

The server's ssh-ed25519 key fingerprint is:

PuTTY's cache and carry on connecting.

adding the key to the cache, hit No.

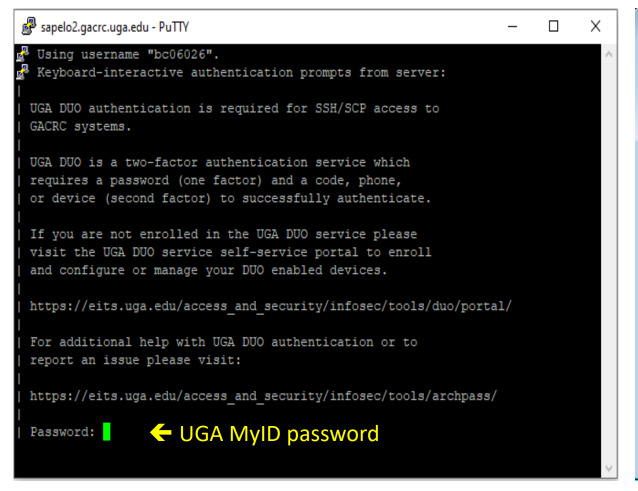
If you trust this host, hit Yes to add the key to

Help

### Step1 (Cont.) - Windows using PuTTY



Next you will enter your UGA MyID password and initiate DUO authentication procedure:



```
sapelo2.gacrc.uga.edu - PuTTY
                                                                      - - X
 requires a password (one factor) and a code, phone,
 or device (second factor) to successfully authenticate.
 If you are not enrolled in the UGA DUO service please
 visit the UGA DUO service self-service portal to enroll
  and configure or manage your DUO enabled devices.
 https://eits.uga.edu/access and security/infosec/tools/duo/portal/
 For additional help with UGA DUO authentication or to
 report an issue please visit:
 https://eits.uga.edu/access and security/infosec/tools/archpass/
 Duo two-factor login for zhuofei
 Enter a passcode or select one of the following options:
   1. Duo Push to XXX-XXX-5758
   2. Phone call to XXX-XXX-5758
   Phone call to XXX-XXX-1925
   4. Phone call to XXX-XXX-3535
   5. SMS passcodes to XXX-XXX-5758
                                      ← DUO option
 Passcode or option (1-5): 1
```



### Step2: On Login node change directory to global scratch

Once you logged on, your current directory will be your <u>home directory</u>

Use cd command to change your current directory to /scratch/MyID

Use ls command to take a look in /scratch/MyID

```
zhuofei@ss-sub1 zhuofei$ ls
user_test
```



#### Step3 - 4: Create and cd to a working subdirectory

• Use mkdir command to make a subdirectory in /scratch/MyID

```
zhuofei@ss-sub1 zhuofei$ mkdir workDir
zhuofei@ss-sub1 zhuofei$ ls
user_test workDir
```

Use cd command to change your current directory to /scratch/MyID/workDir



### Step5: Transfer data from local computer to workDir - Mac/Linux

https://wiki.gacrc.uga.edu/wiki/Transferring Files#Using scp

- 1. You need to connect to the cluster's <u>Transfer node</u> (xfer.gacrc.uga.edu)
- 2. Open Terminal utility on <u>local computer</u> to use scp (-r) [Source] [Target]

#### E.g. 1: working on local computer, from Local → workDir on cluster

```
scp ./file zhuofei@xfer.gacrc.uga.edu:/scratch/zhuofei/workDir/
scp -r ./folder/ zhuofei@xfer.gacrc.uga.edu:/scratch/zhuofei/workDir/
```

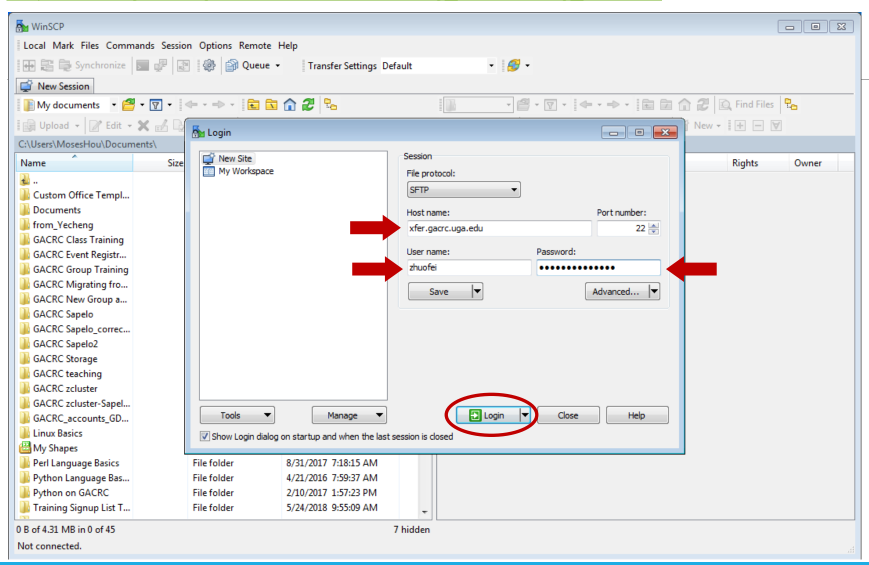
#### *E.g. 2*: working on local computer, from workDir on cluster → Local

```
scp zhuofei@xfer.gacrc.uga.edu:/scratch/zhuofei/workDir/file .
scp -r zhuofei@xfer.gacrc.uga.edu:/scratch/zhuofei/workDir/folder/ .
```

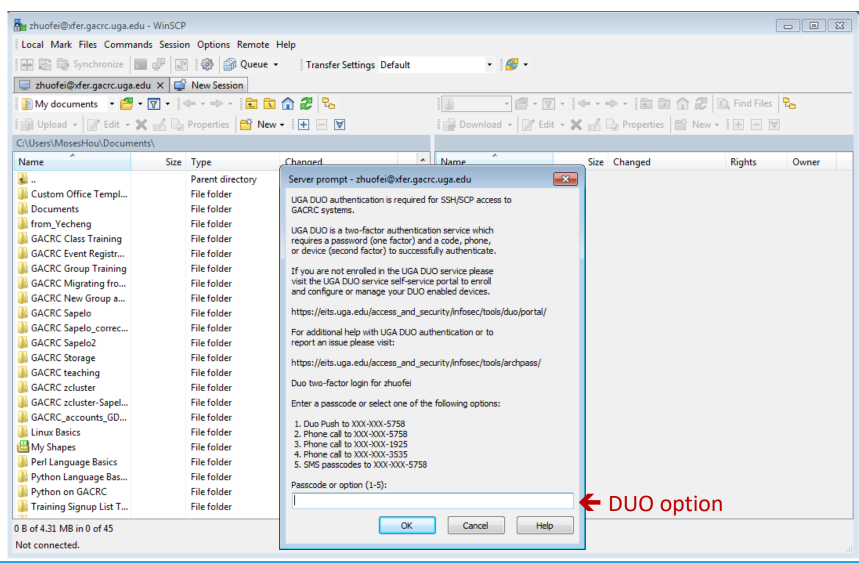


- 1. You need to connect to the cluster's <u>Transfer node</u> (xfer.gacrc.uga.edu)
- 2. Use WinSCP on <u>local computer</u>
  - WinSCP can be downloaded from <a href="https://winscp.net/eng/index.php">https://winscp.net/eng/index.php</a>
  - Default installation procedure is simple
- 3. Alternative FileZilla <a href="https://wiki.gacrc.uga.edu/wiki/Transferring-Files#Using-FileZilla">https://wiki.gacrc.uga.edu/wiki/Transferring-Files#Using-FileZilla</a>

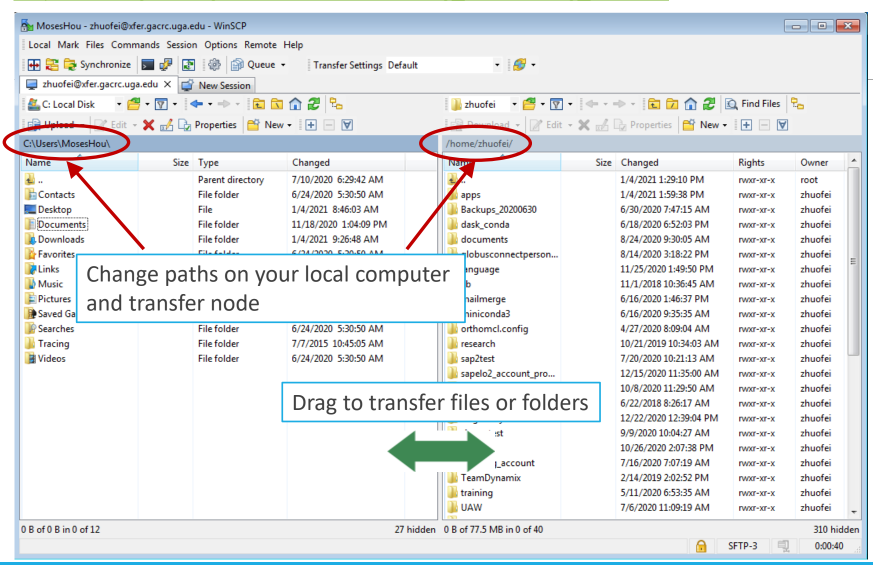














#### Step5 (Cont.): Transfer data on cluster to workDir

- Log on to Transfer node (xfer.gacrc.uga.edu)
  - Mac/Linux: ssh MyID@xfer.gacrc.uga.edu (page 15-16)
  - ✓ Windows: use PuTTY to log in MyID@xfer.gacrc.uga.edu (page 17-19)
- Landing folder: /home/MyID (Home)
- You can transfer data between following directories on cluster using cp or mv:
  - 1. /home/MyID (Home)
  - /scratch/MyID (Scratch)
  - /work/abclab (Work)
  - 4. /project/abclab (Project)
- Most file systems on Transfer are auto-mounted upon the first time full-path access, e.g.,
   cd /project/abclab/



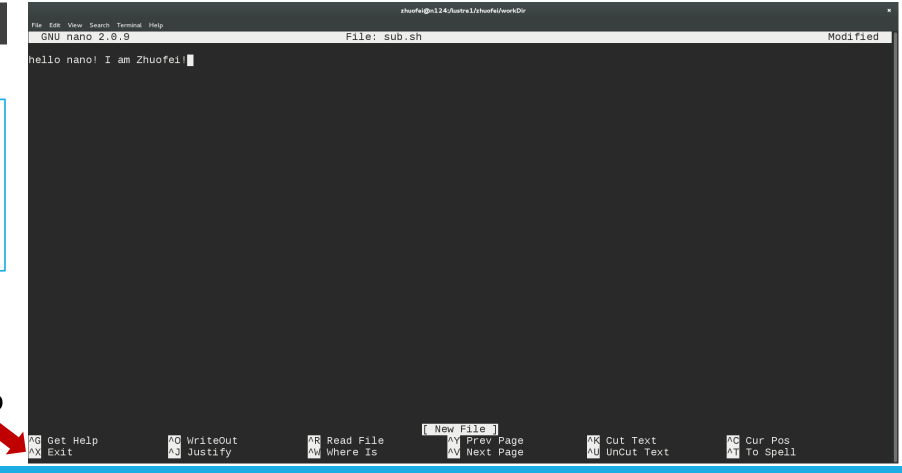
#### Step6: Make a job submission script in workDir

https://wiki.gacrc.uga.edu/wiki/Sample batch job submission scripts on Sapelo2

\$ nano sub.sh

nano is a simple text editor on Linux. You are welcome to use other editors like vim or emacs.

Ctrl-x to save file and quit from nano





```
#!/bin/bash
#SBATCH --job-name=testBowtie2
                                       # Job name (testBowtie2)
#SBATCH --partition=batch
                                       # Queue name (batch)
#SBATCH --ntasks=1
                                        # Run single task using one CPU core on a single node
                                       # Job memory limit (4 GB)
#SBATCH --mem=4G
                                       # Time limit hrs:min:sec or days-hours:minutes:seconds
#SBATCH --time=1:00:00
                                       # Do not load any users' explicit environment variables
#SBATCH --export=NONE
#SBATCH --output=%x %j.out
                                       # Standard output, testBowtie2 1234.out
                                       # Standard error log, testBowtie2 1234.err
#SBATCH --error=%x %j.err
#SBATCH --mail-type=END,FAIL
                                       # Mail events (BEGIN, END, FAIL, ALL)
#SBATCH --mail-user=username@uga.edu
                                        # Where to send mail
cd $SLURM SUBMIT DIR
                                       # Change directory to job submission directory
                                        # Load software module and run bowtie2 below
ml Bowtie2/2.4.1-GCC-8.3.0
bowtie2 -x ./index/lambda virus -U ./myreads.fq -S output.sam
```

^R Read File

^W Where Is

To run the demo example, please copy ^U UnCut Tex these files into your working dir:

cp -r /usr/local/training/Sapelo2/\* .

^0 WriteOut

^J Justify

^G Get Help

Exit

^K Cut Text

^Y Prev Page

^V Next Page



#### Step7: Submit a job from workDir using sbatch

https://wiki.gacrc.uga.edu/wiki/Running Jobs on Sapelo2#How to submit a job to the batch partition

```
zhuofei@ss-sub1 workDir$ pwd
/scratch/zhuofei/workDir
zhuofei@ss-sub1 workDir$ ls
index myreads.fq sub.sh
zhuofei@ss-sub1 workDir$ sbatch sub.sh
Submitted batch job 32860
```

#### sub.sh is a job submission script to

- 1. specify computing resources:
- 2. load software using ml *moduleName*
- 3. run any Linux commands you want to run
- 4. run the software



#### Step8: Check job status using squeue --me

https://wiki.gacrc.uga.edu/wiki/Monitoring Jobs on Sapelo2

```
zhuofei@ss-sub1 workDir$ squeue --me
JOBID PARTITION
                   NAME
                            USER
                                                        NODELIST (REASON)
                                  ST
                                         TIME
                                               NODES
32862
                         zhuofei
                                         0:00
         batch testBowt
                                  PD
                                                        (None)
32861
                         zhuofei R
                                         0:05
                                                       c5 - 19
         batch testBowt
32860
         batch testBowt
                         zhuofei R
                                         4:37
                                                       c5-19
32859
                                                       b1-2
         batch testBowt
                         zhuofei
                                         9:29
                                  CD
```

R: job is Running

PD: job is PenDing, waiting for resources to become available

**CD**: job is CompleteD and is not longer running

CA: job is CAnceled and is not longer running

F: job is Failed (crashed) on the node and is not longer running

Note: "TIME" is the elapsed wall-clock time of your job running on cluster, instead of the CPU time.



### Step8: Check job status using squeue --me -l

https://wiki.gacrc.uga.edu/wiki/Monitoring Jobs on Sapelo2

zhuofei@ss-sub1 workDir\$ squeueme -l										
Tue Sep 15 15:00:51 2020										
JOBID	PARTITION	NAME	USER	STATE	TIME	TIME_LIMI	NODES	NODELIST (REASON)		
32866	batch	testBowt	zhuofei	RUNNING	0:14	1:00:00	1	rc6-10		
32865	batch	testBowt	zhuofei	RUNNING	0:30	1:00:00	1	rc6-10		
32864	batch	testBowt	zhuofei	RUNNING	0:33	1:00:00	1	rc6-10		

### Step8 (Cont.): Cancel job using scancel

https://wiki.gacrc.uga.edu/wiki/Running Jobs on Sapelo2#How to cancel .28delete.29 a running or pending job

zhuofei@ss-sub1 workDir\$ squeueme -l Tue Sep 15 15:08:27 2020									
JOBID PARTITION NAME	USER STATE	TIME TIME LIMI	NODES NODELIST (REASON)						
32869 batch testBowt	zhuofei RUNNING	$0:01  1:\overline{0}0:00$	1 c5-19						
32868 batch testBowt	zhuofei RUNNING	0:05 1:00:00	1 c5-19						
32867 batch testBowt	zhuofei RUNNING		1 c5-19						
zhuofei@ss-sub1 workDir\$									
zhuofei@ss-sub1 workDir\$	scancel 32867								
zhuofei@ss-sub1 workDir\$	squeueme -l								
Tue Sep 15 15:08:45 2020									
JOBID PARTITION NAME	USER STATE	TIME TIME_LIMI	NODES NODELIST (REASON)						
32867 batch testBowt	zhuofei COMPLETI								
32869 batch testBowt	zhuofei RUNNING	0:19 1:00:00	1 c5-19						
32868 batch testBowt	zhuofei RUNNING	0:23 1:00:00	1 c5-19						
zhuofei@ss-sub1 workDir\$	squeueme -l								
Tue Sep 15 15:08:50 2020									
JOBID PARTITION NAME	USER STATE	TIME TIME_LIMI	NODES NODELIST (REASON)						
32869 batch testBowt	zhuofei RUNNING	0:19 1:00:00	1 c5-19						
32868 batch testBowt	zhuofei RUNNING	0:23 1:00:00	1 c5-19						

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```
bc06026@b1-24 workdir$ sacct
   JobID JobName Partition Account AllocCPUS State ExitCode
          testBowti+
                     batch gacrc-ins+
                                       1 RUNNING
                                                     0:0
3326893
3326893.bat+ batch
                        gacrc-ins+ 1 RUNNING
                                                   0:0
3326893.ext+
                    gacrc-ins+ 1 RUNNING
                                                   0:0
             extern
bc06026@b1-24 workdir$ sacct -X
   JobID JobName Partition Account AllocCPUS State ExitCode
                     batch gacrc-ins+ 1 RUNNING
3326893
          testBowti+
                                                     0:0
bc06026@b1-24 workdir$ sacct -X --format jobid, state
   JobID
          State
3326893
           RUNNING
bc06026@b1-24 workdir$
```

#### zhuofei@ss-sub1 workDir\$ sacct-gacrc

JobID JobName User Partition NodeList AllocNodes NTasks NCPUS ReqMem MaxVMSize State CPUTime Elapsed Timelimit ExitCode WorkDir

275	bowti+ z	huofei	batch	rc6-10	1		1	4Gn		COMPLETED	00:09:13	00:09:13	01:00:00	0:0 /scratch/zhu
275.batch	batch			rc6-10	1	1	1	4Gn	396868K	COMPLETED	00:09:13	00:09:13		0:0
275.extern	extern			rc6-10	1	1	1	4Gn	142616K	COMPLETED	00:09:13	00:09:13		0:0
276	amberjob	shtsai	gpu_p	c4-23	1		1	10Gn		CANCELLED+	00:03:19	00:03:19	02:00:00	0:0 /scratch/sht
276.batch	batch			c4-23	1	1	1	10Gn	221140K	CANCELLED	00:03:20	00:03:20		0:15
276.extern	extern			c4-23	1	1	1	10Gn	169800K	COMPLETED	00:03:19	00:03:19		0:0
277	mpitest	shtsai	batch	c2-[11-12]	2		24	600Mc		COMPLETED	04:01:12	00:10:03	02:00:00	0:0 /scratch/sht
277.batch	batch			c2-11	1	1	12	600Mc	221268K	COMPLETED	02:00:36	00:10:03		0:0
277.extern	extern			c2-[11-12]	2	2	24	600Mc	169800K	COMPLETED	04:01:12	00:10:03		0:0
277.0	orted			c2-12	1	1	1	600Mc	265640K	COMPLETED	00:00:01	00:00:01		0:0
278	bash	shtsai	inter_p	c2-4	1		1	2Gn		RUNNING	00:13:37	00:13:37	12:00:00	0:0 /scratch/sht
278.extern	extern			c2-4	1	1	1	2Gn		RUNNING	00:13:37	00:13:37		0:0
278.0	bash			c2-4	1	1	1	2Gn		RUNNING	00:13:37	00:13:37		0:0

#### zhuofei@ss-sub1 workDir\$ sacct-gacrc-v 47939

JobID 47939

JobName testBowti+

User zhuofei

Partition batch

NodeList c1-3

AllocNodes :

State RUNNING

CPUTime 00:00:28

Elapsed 00:00:28

Timelimit 01:00:00

ExitCode 0:0

WorkDir /scratch/zhuofei/workDir\_sapelo2

NTasks

NCPUS 1

ReqMem 4Gn

zhuofei@ss-sub1 workDir\$ seff 37259

Job ID: 37259

Cluster: tc2

User/Group: zhuofei/gacrc-instruction

State: COMPLETED (exit code 0)

Cores: 1

CPU Utilized: 00:09:45

CPU Efficiency: 99.66% of 00:09:47 core-walltime

Job Wall-clock time: 00:09:47

Memory Utilized: 197.34 MB

Memory Efficiency: 4.82% of 4.00 GB



#### Obtain Job Details

https://wiki.gacrc.uga.edu/wiki/Running Jobs on Sapelo2#How to check resource utilization of a running or finished job

Option 1: squeue --me for details of a pending or running jobs

Option 2: sacct or sacct-gacrc (-v) for details of computing resource usage of

a <u>running or finished</u> job

Option 3: seff for details of computing resource usage of a finished job

Option 4: Email notification from finished jobs (completed, canceled, or crashed), if using:

```
#SBATCH --mail-user=username@uga.edu
```

#SBATCH --mail-type=ALL



# Interactive jobs

https://wiki.gacrc.uga.edu/wiki/Running Jobs on Sapelo2#How to open an interactive session

https://wiki.gacrc.uga.edu/wiki/Running Jobs on Sapelo2#How to run an interactive job with Graphical User Interface capabilities

Description	Command
Start an interactive session	qlogin
Start an interactive session with X forwarding	xqlogin

qlogin	srunpty -p inter_pmem=2Gnodes=1ntasks-per-node=1time=12:00:00job-name=qlogin bash -l
xqlogin	srunptyx11 -p inter_pmem=2Gnodes=1ntasks-per-node=1time=12:00:00job-name=xqlogin bash -l



# GACRC Wiki <a href="http://wiki.gacrc.uga.edu">http://wiki.gacrc.uga.edu</a> Kaltura channel <a href="https://kaltura.uga.edu/channel/GACRC/176125031">https://kaltura.uga.edu/channel/GACRC/176125031</a>

System: <a href="https://wiki.gacrc.uga.edu/wiki/Systems#Sapelo2">https://wiki.gacrc.uga.edu/wiki/Systems#Sapelo2</a>

Connection: <a href="https://wiki.gacrc.uga.edu/wiki/Connecting#Connecting">https://wiki.gacrc.uga.edu/wiki/Connecting#Connecting to Sapelo2</a>

Software: <a href="https://wiki.gacrc.uga.edu/wiki/Software">https://wiki.gacrc.uga.edu/wiki/Software</a> on Sapelo2

Running Jobs: https://wiki.gacrc.uga.edu/wiki/Running Jobs on Sapelo2

Monitoring Jobs: <a href="https://wiki.gacrc.uga.edu/wiki/Monitoring Jobs on Sapelo2">https://wiki.gacrc.uga.edu/wiki/Monitoring Jobs on Sapelo2</a>

Sample scripts: <a href="https://wiki.gacrc.uga.edu/wiki/Sample-batch-job-submission-scripts-on-Sapelo2">https://wiki.gacrc.uga.edu/wiki/Sample-batch-job-submission-scripts-on-Sapelo2</a>

Transferring Files: <a href="https://wiki.gacrc.uga.edu/wiki/Transferring-Files">https://wiki.gacrc.uga.edu/wiki/Transferring-Files</a>

Linux Commands: <a href="https://wiki.gacrc.uga.edu/wiki/Command-List">https://wiki.gacrc.uga.edu/wiki/Command-List</a>

Training: <a href="https://wiki.gacrc.uga.edu/wiki/Training">https://wiki.gacrc.uga.edu/wiki/Training</a>



## GACRC Help and Support

https://wiki.gacrc.uga.edu/wiki/Getting Help

## 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

When you ask GACRC to test or troubleshoot your jobs, Please make sure of the correctness of your datasets being used!

## **GACRC Service Catalog**

Georgia Advanced Computing Resource Center (GACRC) service catalog.

If you would like to reach out to GACRC and do not have a UGA MyID, please send an email to gacrc-help@uga.edu, and we will respond promptly.

### Categories (3)

#### **Services For Users**

General user support, request software installation or update, request training.

#### Services for PIs

For PIs only: Lab registration, user account creation/modification, class account requests, storage quota modifications.

#### For GACRC Staff

For GACRC's internal use only.

## **Services For Users**

General user support, request software installation or update, request training.

## Services (3)

### **General Support**

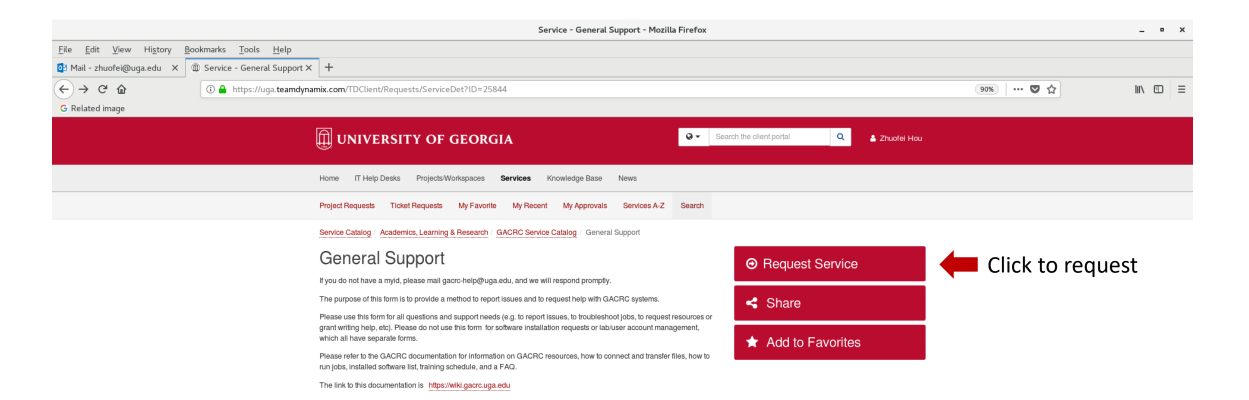
Report issues and request help with GACRC systems, except for software installation requests and account/lab creation requests.

### Software Installation/Update

Request software and common application database (e.g. NCBI blast databases) installation and upgrade.

### **Training**

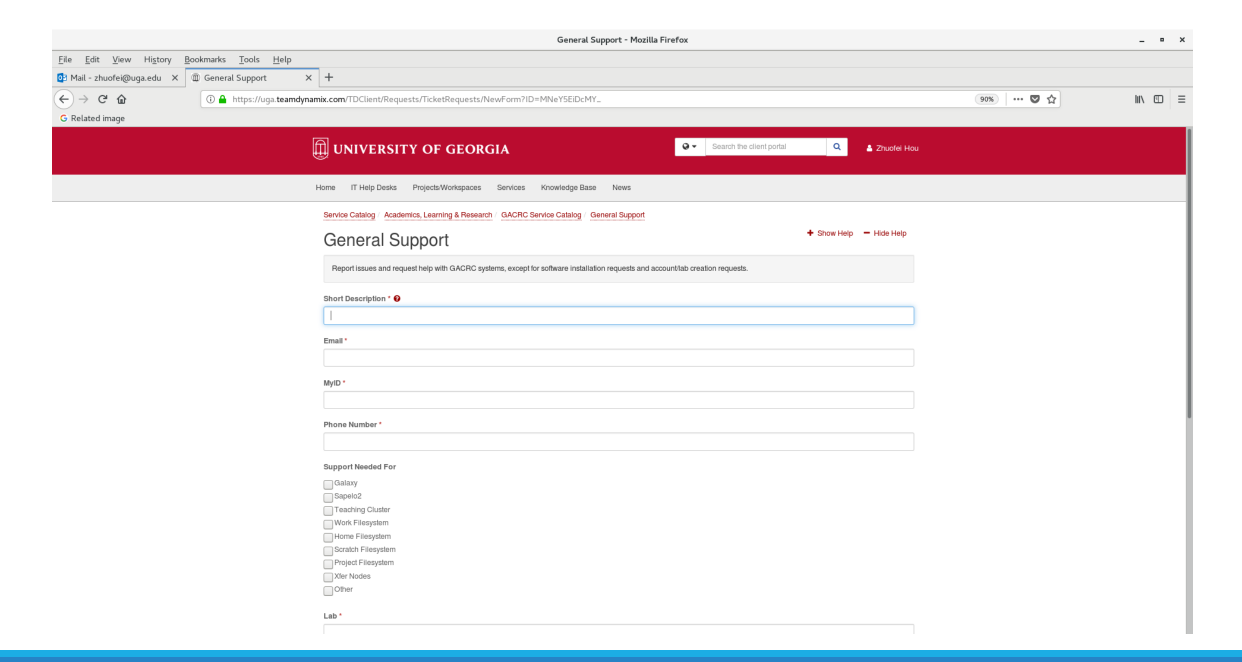
Request support related to training provided by the GACRC.



This site is operated by Enterprise Information Technology Services (EITS) at the University of Georgia.

Privacy | Accessibility | Website Feedback

https://uga.teamdynamix.com/TDClient/Requests/ServiceCatalogSearch





# Slurm job states

Code	State	Meaning
R	Running	Job is running on compute node(s)
PD	Pending	Job is waiting for compute node(s)
CD	Completed	Job completed
CG	Completing	Job is completing
CA	Canceled	Job was canceled
F	Failed	Job terminated with non-zero exit code
NF	Node Fail	Job terminated due to failure of node(s)



# Commands for submitting and canceling jobs

Description	Slurm Command
Submit a batch job to queue	sbatch sub.sh
Delete a job from queue	scancel <jobid></jobid>
Cancel all your job(s)	scancel -u <username></username>
Cancel all your pending job(s)	scancel -t PENDING -u <username></username>
Cancel your job(s) by job name	scancelname <jobname></jobname>
Cancel an element (index) of an array job (jobID)	scancel <jobid>_<index></index></jobid>



# Commands for monitoring jobs

Description	Slurm Command
Command	squeue or squeue -l
Job status of all your jobs	squeueme or squeueme -l
Job status of a job	squeue -j <jobid></jobid>
Job status of job(s) from a user	squeue -u <username></username>
Job status with details	scontrol show job <job id=""> scontrol show job -dd <job id=""></job></job>
Job's resource usage	sacct-gacrc
View job batch script	scontrol write batch_script <job id=""> [filename]</job>



## Slurm headers for running a Serial (single-core) job

https://wiki.gacrc.uga.edu/wiki/Sample batch job submission scripts on Sapelo2

```
#!/bin/bash
#SBATCH --job-name=testBowtie2
                                         # Job name (testBowtie2)
#SBATCH --partition=batch
                                         # Queue name (batch)
#SBATCH --ntasks=1
                                         # Run in a single task using one CPU core on a single node
#SBATCH --mem=4G
                                         # Job memory limit (4 GB)
#SBATCH --time=1:00:00
                                         # Time limit hrs:min:sec or days-hours:minutes:seconds
#SBATCH --export=NONE
                                         # Do not load any users' explicit environment variables
#SBATCH --output=%x %j.out
                                         # Standard output log, e.g., testBowtie2_1234.out
#SBATCH --error=%x_%j.err
                                         # Standard error log, e.g., testBowtie2 1234.err
#SBATCH --mail-type=END,FAIL
                                         # Mail events (BEGIN, END, FAIL, ALL)
#SBATCH --mail-user=username@uga.edu
                                         # Where to send mail
cd $SLURM SUBMIT DIR
                                         # Change directory to job submission directory
ml Bowtie2/2.4.1-GCC-8.3.0
                                         # Load software module and run bowtie2 below
bowtie2 -x ./index/lambda_virus -U ./myreads.fq -S output.sam
```





```
#!/bin/bash
                                         # Job name (testBowtie2)
#SBATCH --job-name=testBowtie2
#SBATCH --partition=batch
                                         # Queue name (batch)
#SBATCH --nodes=1
                                         # Run all processes on a single node
#SBATCH --ntasks=1
                                         # Run in a single task on a single node
#SBATCH --cpus-per-task=8
                                         # Number of CPU cores per task (8)
#SBATCH --mem=10G
                                         # Job memory limit (10 GB)
#SBATCH --time=1:00:00
                                         # Time limit hrs:min:sec or days-hours:minutes:seconds
#SBATCH --export=NONE
                                         # Do not load any users' explicit environment variables
#SBATCH --output=%x %j.out
                                         # Standard output log, e.g., testBowtie2 1234.out
#SBATCH --error=%x %j.err
                                         # Standard error log, e.g., testBowtie2 1234.err
#SBATCH --mail-type=END,FAIL
                                         # Mail events (BEGIN, END, FAIL, ALL)
#SBATCH --mail-user=username@uga.edu # Where to send mail
cd $SLURM_SUBMIT_DIR
ml Bowtie2/2.4.1-GCC-8.3.0
bowtie2 -p 8 -x ./index/lambda virus -U ./myreads.fq -S output.sam
```

## Slurm headers for running an Array job



```
#!/bin/bash
#SBATCH --job-name=testBowtie2Array
                                          # Job name (testBowtie2Array)
#SBATCH --partition=batch
                                          # Queue name (batch)
#SBATCH --ntasks=1
                                          # Run in a single task using one CPU core on a single node
#SBATCH --mem=4G
                                          # Job memory limit (4 GB)
#SBATCH --time=1:00:00
                                          # Time limit hrs:min:sec or days-hours:minutes:seconds
#SBATCH --export=NONE
                                          # Do not load any users' explicit environment variables
#SBATCH --output=%x_%j.out
                                          # Standard output log, e.g., testBowtie2Array 1234.out
#SBATCH --error=%x %j.err
                                          # Standard error log, e.g., testBowtie2Array 1234.err
#SBATCH --array=0-9
                                          # Array element range from 0 to 9, i.e. 10 element jobs
cd $SLURM_SUBMIT_DIR
ml Bowtie2/2.4.1-GCC-8.3.0
                                         # Original data is split into 10 pieces and run in each element job
bowtie2 -x ./index/lambda_virus -U ./myreads_$SLURM_ARRAY_TASK_ID.fq \
-S output_$SLURM_ARRAY_TASK_ID.sam
```



## Slurm headers for running a Threaded (OpenMP) job

```
#!/bin/bash
#SBATCH --job-name=testOpenMP
                                             # Job name (testOpenMP)
#SBATCH --partition=batch
                                             # Queue name (batch)
#SBATCH --nodes=1
                                             # Run all processes on a single node
#SBATCH --ntasks=1
                                             # Run in a single task on a single node
#SBATCH --cpus-per-task=12
                                             # Number of CPU cores per task (12)
                                             # Job memory limit (10 GB)
#SBATCH --mem=10G
#SBATCH --export=NONE
                                             # Do not load any users' explicit environment variables
#SBATCH --time=24:00:00
                                             # Time limit hrs:min:sec or days-hours:minutes:seconds
#SBATCH --output=%x_%j.log
                                             # Standard output and error log, e.g., testOpenMP_1234.log
#SBATCH --mail-type=END,FAIL
                                             # Mail events (BEGIN, END, FAIL, ALL)
#SBATCH --mail-user=username@uga.edu
                                             # Where to send mail
cd $SLURM_SUBMIT_DIR
export OMP_NUM_THREADS=$SLURM_CPUS_PER_TASK # Sets the number of threads to use for OpenMP parallel regions
ml foss/2019b
                                             # Load toolchain module
./myProgram.x
                                             # Run your program binary compiled with OpenMP
```

## Slurm headers for running an MPI job



```
#!/bin/bash
#SBATCH --job-name=testMPI
                                         # Job name (tesMPI)
#SBATCH --partition=batch
                                         # Queue name (batch)
#SBATCH --nodes=2
                                         # Run on two nodes
                                         # How many tasks on each node; Number of tasks=32=MPI ranks
#SBATCH --ntasks-per-node=16
#SBATCH --cpus-per-task=1
                                         # Number of CPU cores per task; 16 CPU cores per node
#SBATCH --mem-per-cpu=500M
                                         # Memory per allocated CPU; 8GB (500MB*16) memory per node
                                         # Time limit hrs:min:sec or days-hours:minutes:seconds
#SBATCH --time=24:00:00
#SBATCH --export=NONE
                                         # Do not load any users' explicit environment variables
#SBATCH --output=%x %j.log
                                         # Standard output and error log, e.g., testMPI 1234.log
#SBATCH --mail-type=END,FAIL
                                         # Mail events (BEGIN, END, FAIL, ALL)
#SBATCH --mail-user=username@uga.edu
                                        # Where to send mail
cd $SLURM_SUBMIT_DIR
ml foss/2019b
                                         # Load toolchain module
mpirun -n 32 ./myProgram.x
                                         # Run your program binary compiled with OpenMPI with 32 ranks
```



# Slurm headers for running a Hybrid MPI/OpenMP job

```
#!/bin/bash
#SBATCH --job-name=testHybrid
                                         # Job name (testHybrid)
#SBATCH --partition=batch
                                         # Queue name (batch)
#SBATCH --nodes=2
                                         # Run on two nodes
#SBATCH --ntasks-per-node=8
                                         # How many tasks on each node; Number of tasks=16=MPI ranks
#SBATCH --cpus-per-task=4
                                         # Number of CPU cores per task; 32 CPU cores per node
                                         # Memory per allocated CPU; 16GB (500MB*32) memory per node
#SBATCH --mem-per-cpu=500M
#SBATCH --time=24:00:00
                                         # Time limit hrs:min:sec or days-hours:minutes:seconds
                                         # Do not load any users' explicit environment variables
#SBATCH --export=NONE
#SBATCH --output=%x_%j.log
                                         # Standard output and error log
#SBATCH --mail-type=END,FAIL
                                         # Mail events (BEGIN, END, FAIL, ALL)
                                         # Where to send mail
#SBATCH --mail-user=username@uga.edu
cd $SLURM_SUBMIT_DIR
                                         # Change directory to job submission directory
export OMP_NUM_THREADS=$SLURM_CPUS_PER_TASK # Sets the number of threads to use for OpenMP parallel regions
ml foss/2019b
                                         # Load toolchain module
mpirun -n 16 ./myProgram.x
                                         # Run your program binary compiled with OpenMPI with 16 ranks
```

## Slurm headers for running a GPU job



```
#!/bin/bash
#SBATCH --job-name=amber
                                              # Job name
                                              # Partition (queue) name, i.e., gpu p
#SBATCH --partition=gpu p
#SBATCH --gres=gpu:1
                                              # Requests one GPU device; --gres=gpu:P100:1, --gres=gpu:K40:1
#SBATCH --ntasks=1
                                              # Run a single task
#SBATCH --cpus-per-task=2
                                              # Number of CPU cores per task
#SBATCH --mem=40gb
                                              # Job memory request
#SBATCH --time=10:00:00
                                              # Time limit hrs:min:sec
#SBATCH --export=NONE
                                              # Do not load any users' explicit environment variables
#SBATCH --output=%x %j.out
                                              # Standard output and error log, e.g., amber_1234.out
#SBATCH --mail-type=END,FAIL
                                              # Mail events (BEGIN, END, FAIL, ALL)
#SBATCH --mail-user=username@uga.edu
                                              # Where to send mail
cd $SLURM_SUBMIT_DIR
ml Amber/18-fosscuda-2018b-AmberTools-18-patchlevel-10-8
mpiexec $AMBERHOME/bin/pmemd.cuda -O -i ./prod.in -o prod_c4-23.out -p ./dimerFBP_GOL.prmtop -c ./restart.rst \
-r prod.rst -x prod.mdcrd
```





```
#!/bin/bash
#SBATCH --job-name=test sortmerna
                                               # Job name
#SBATCH --partition=batch
                                               # Partition (queue) name
#SBATCH --ntasks=1
                                               # Run on a single CPU
#SBATCH --mem=8gb
                                               # Job memory request
#SBATCH --time=02:00:00
                                               # Time limit hrs:min:sec
#SBATCH --export=NONE
                                               # Do not load any users' explicit environment variables
#SBATCH --output=sortmerna.%j.out
                                               # Standard output log, e.g., sortmerna.1234.out
#SBATCH --error=sortmerna.%j.err
                                               # Standard error log, e.g., sortmerna.1234.err
                                               # Number of CPU cores per task
#SBATCH --cpus-per-task=4
#SBATCH --mail-type=END,FAIL
                                               # Mail events (NONE, BEGIN, END, FAIL, ALL)
#SBATCH --mail-user=username@uga.edu
                                               # Where to send mail
cd $SLURM SUBMIT DIR
singularity exec /apps/singularity-images/sortmerna-3.0.3.simg sortmerna --threads 4 --ref db.fasta,db.idx --reads file.fa \
--aligned base name output
```



# General guidelines

- Do NOT use Login nodes to run CPU/memory intensive tasks directly → submit jobs to Compute nodes!
- Do NOT use Login nodes to transfer data between your local computer and cluster → use Transfer nodes!
- Do NOT use Home for storing job data → use /scratch/MyID
- Do NOT park data in Scratch or Local Scratch → clean up when job finishes or exits from node
- Do NOT park data permanently in Project → download data to your local drive
- NO large memory job running on batch partition → use highmem\_p
- NO small memory job running on highmem\_p partition → use batch
- In general, <u>number of threads you want to run with a parallel job = number of cores requested</u>
- When you archive data using tar on /scratch, please do not use the z option (compression option). After you archived data with tar, you can use gzip to compress it.



## General guidelines

No directory should not have too many files inside! A rule of thumb would be
to try to keep no more than a few tens of thousands of files (<10000 would be
even better) in any single directory which is accessed frequently</li>







All files are in ONE single dir!



Files are organized in subdirs!





# Thank You!

#### **Telephone Support**

EITS Help Desk: 706-542-3106

Monday – Thursday: 7:30 a.m. – 7:30 p.m.

Friday: 7:30 a.m. – 6 p.m.

Saturday – Sunday: 1 p.m. – 7 p.m.

Georgia Advanced Computing Resource Center

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University of Georgia

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