

Research Computing New User Seminar

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 - Slides:

https://github.com/ResearchComputing/New_User_Seminar

Before We Begin

- Goals
 - Inform people about RC resources, expectations, etc.
 - Reduce user frustrations, questions
 - Avoid misunderstandings, lost time, lost work
 - Inform users about best practices
- Things to take particular note of
 - Confusing, ambiguous, highly nuanced concepts
 - Common mistakes or frustrations
 - Best Practices
- Good questions to ask
 - Why? Questions
 - If a question is said to be covered later feel free to re-ask if it's not answered to your satisfaction.





Outline

- What is RC?
 - Resources and services/support
 - Summit overview
- Steps to get access to our systems
 - Accounts
 - Two-factor authentication
 - Logging in
- Navigating our systems
 - Blanca
 - Petalibrary
- Summit (and other RC cluster) New user
 - Allocations
 - Storage spaces
 - Data transfer Globus
 - Software
- Running jobs





What is Research Computing?

- Provide services for researchers that include:
 - Large scale computing
 - Data storage
 - High speed data transfer
 - Data management support
 - Consulting
 - Training
- We are likely best known for:
 - Summit Supercomputer
 - PetaLibrary storage





What Would I Use Summit For?

- Research Computing is more than just Summit
- What would you use Summit For?
 - Solving large problems that require more:
 - Memory than you have on your personal computer
 - Cores/nodes/power than you have on your personal computer
 - High performance GPU computing
 - High memory jobs
 - Visualization rendering
- Not a place for:
 - Large data storage





Hardware - Summit Supercomputer

- 450+ compute nodes (mostly Intel Xeon Haswell)
- 24 cores per "shas" (general compute) node, different core counts for other node types
- 11,400 total cores
- Omni-Path network
- 1.2 PB scratch storage
- GPFS Scratch File system
- 67% CU, 23% CSU, 10% RMACC



Additional Types of Summit Compute Nodes

- 10 Graphics Processing Unit (GPU) Nodes
 - NVIDIA Tesla K80 (2/node)
- 5 High Memory Nodes
 - 2 TB of memory/node, 48 cores/node
- Phi Nodes
 - 20 nodes
 - Intel Xeon Phi





How To Access RC Resources?

- Get an account
- 2. Set up two factor authentication
- 3. Inform us of any specific needs
- 4. Log in
- 5. Create greatness
- After you login, you will need to do many additional things that we will discuss today





Getting an RC Account

- CU Boulder users and affiliates:
- Request an account through the RC Account request portal
 - https://portals.rc.colorado.edu/accounts/account-request/create
 - If you are requesting access to Summit make sure you indicate this

CSU Users:

- Request an CSU eID if you don't have one
- Fill out account application form
- Duo authentication
- Then get an RC user account
- https://www.acns.colostate.edu/hpc/summit-get-started/

RMACC Users:

- Email <u>rc-help@colorado.edu</u> and we'll guide you through the process
- Email your local RMACC representative also, if known.





Two-Factor Authentication

- Provides an extra level of authentication
 - We are outside the firewall!
 - Valuable resources
 - Inviting, high-profile target
 - Lost time investigating/fixing and damage to our reputation if compromised

• Duo

- Most users use the Duo smartphone app
- "Phone Call" or SMS (text) are alternatives
- Physical code generator "token" available for \$20





Duo Authentication

- Once UCB users get an account, we send a Duo invitation
- Once you get the invitation, you'll get a series of steps to complete Duo enrollment
- For UCB RC supports Duo "push", "SMS", "phone call" and the physical "Token" (code generator) for authentication
- Users greatly prefer "push" it's also the cheapest for UCB
- Once your Duo is set up UCB users can manage their Duo setup at duo.colorado.edu





Logging In

• It's important to note that you are NOT logging into any specific resource, Summit, etc.

- When you log in, you land on our login nodes
 - RC users have used the same hostname to log in and accessed the same non-scratch storage for over 9 years and 5 clusters
- From there, you can access our other resources



Accessing RC Resources

To login to an RC login node:

```
ssh <u>username@login.rc.colorado.edu</u>
Supply your IdentiKey password and your Duo
app will alert you to confirm the login
```

 If you need to use something other than the Push app your enter your password as:

```
identikey_password,phone or
identikey password,sms
```





Navigating our Systems

- Now that you've logged in, now what?
 - What are the different node types we have?
 - What are the different storage spaces?
 - What should I be putting in these storage spaces?
 - How do I transfer data around?
 - How do I deal with software?



Blanca

- If you are a new Blanca user you need an RC account, and we need to know what resources to connect you to.
- To run jobs as a Blanca user, once you've logged into a login node, load the Blanca slurm module

```
module load slurm/blanca
```

- Only certain users have access to Blanca paid service
- If you are unsure, you can ask your advisor or RC





PetaLibrary

- To access the PetaLibrary, you login in to one our RC's login nodes as normal
- Then you cd to either /work/<groupname> or /archive/<groupname>, depending on your PetaLibrary service
 - <groupname> is the name set for your group when you set up the PetaLibrary service
 - You do not include the <>
- Only certain users have access to PetaLibrary paid service
- If you are unsure, you can ask your advisor or RC
 - But likely if you are unsure you don't have access





Using Summit (and other RC clusters)



Allocations

- You will need a compute time allocation to use any of our compute resources.
- Blanca and Summit Condo allocations are part of the buy-in
- New users are offered a checkbox for Summit access when creating their account, this gets you added to the Summit General allocation.
- Otherwise, to request access please email <u>rc-help@colorado.edu</u> and ask for access to the General allocation
 - Need to provide a few sentences on your project
- Once you have some benchmarks, most users will want to get an allocation to get access to a lot larger "share" of Summit.





Why Do I Need to request access to Summit?

- Not all UCB users need or want to use Summit
- I have an account why do I need an allocation?
 - An account validates you are eligible to use RC compute resources
 - An allocation allows us to keep track of your use of the system
 - This is important because:
 - We need to make sure we have enough resources to accommodate all of our users
 - Helps for reporting to NSF and the CU Research & Innovation Office
 - Applying for an allocation beyond General:
 - Gives you higher priority in the system and access to more compute time





What is Fair Share?

- Fair share scheduling uses a complex formula to determine priority in queue
- Looks at load for each user and each QOS and balances utilization to fairly share resources
 - Involves historical use by user plus how long job has been in the queue
- System will first look at weighted average utilization of user mostly over the last 4 weeks
- Then compare it to the fair share target percentage of a user





Fair Share Target Percentage

- The target percentage depends on your priority based on your project proposal
- Everyone not associated with a project shares a target percentage of 13% (20% of the CU fraction)
 - No guaranteed level per user
- If you are under (over) your target percentage (based on a 4 week average) your priority is increased (decreased)
- Reminder: this all only impacts pending jobs
- If no other pending jobs and enough resources are available then your job will run regardless of your previous usage





Allocations

- Did not request Summit access when you signed up?
- Make a request now!
 - Include 2-3 sentences describing your proposed usage
 - Email <u>rc-help@colorado.edu</u>
- If you need access to a condo or research group allocation email us and cc the PI or Faculty who can authorize it.



Different Node Types

- Login nodes
 - Four virtual machines
 - This is where you are when you log in
 - No heavy computation, interactive jobs, or long running processes
 - Great for script or code editing
 - Also Job submission, checking on jobs, looking at output
- Compile nodes
 - Where you compile code, install packages, etc.
 - Explore the Summit software environment
- Compute/batch nodes
 - This is where jobs that are submitted through the scheduler run
 - Intended for heavy computation





Storage Spaces

Home Directories

- /home/\$USER
- Not for direct computation
- Small quota (2 GB)
- Backed up

• \$PROJECT Space

- /projects/\$USER
- Mid level quota (250 GB)
- Large file storage
- Backed up

Scratch Directory

- /scratch/summit/\$USER
- 10 TB
 - Can ask for more if needed
- Files purged around 90 days





What Belongs Where?

/home

- Scripts
- Code
- Small, important files/directories
- Inappropriate for sharing files with others
- Inappropriate for job output

/projects

- Code/files/libraries
- Software you are installing
- · Mid-level size input files
- Appropriate for sharing files with others
- Inappropriate for job output
- /scratch/summit
 - Output from running jobs
 - Large files/datasets
 - Appropriate for sharing files with others
 - THIS IS NOT APPROPRIATE FOR LONG TERM STORAGE





Transferring Data

- Globus is Research Computing's preferred method of data transfer for larger files or datasets
- Designed with researchers in mind
- End points between computers make for efficient data transfer with an easy to use interface
 - Endpoints are different locations that data can be moved to/from
 - Personal or multi-user
- rsync and sftp through the login nodes is good for small transfers – transfers that take a few minutes.





Setting Up Globus

- Create an account at Globus.org
- Make your personal computer an endpoint if you want
 - Not needed if you are transferring between two other endpoints, like a repository and RC
- Transfer data
 - www.globus.org

Software

- Common software is available to everyone on the systems
- Can install your own software
 - But you are responsible for support
 - We are happy to assist
- Research Computing uses modules to manage software
 - You load modules to prepare your environment for using software
 - Modules set any environment variables, paths, etc.
 - Set environment so application can find appropriate libraries, etc.





Important Things to Know About Modules

- You need to be on a compile node to browse the modules
- · Some modules might require a specific hierarchy to load
 - For some modules, you may need to specify a specific version
 - For example, module load R/3.3.0
 - For other modules, you may be able to be more generic
 - For example, module load matlab
- Some modules may require you to first load other modules that they depend on
- To find dependencies for a module, type module spider <package>
- To find out what software is available, you can type module avail
- To set up your environment to use a software package, type module load <package>/<version>





Job Submission



Running Jobs

- What is a "job"?
 - Jobs are scripted packages of work for the cluster to perform
 - Have a job number
- Interactive jobs
 - Work interactively at the command line of a compute node
- Batch jobs
 - Submit job script which will be executed when resources are available
 - Create a text file, shell script containing information about the job
 - Submit the job file to a queue





Job Scheduling

- On a supercomputer, jobs are scheduled rather than just run instantly at the command line
 - Shared system
 - Jobs are put in a queue until resources are available
- Need software that will distribute the jobs appropriately and manage the resources
 - Simple Linux Utility for Resource Management (Slurm)
 - Keeps track of what nodes are busy/available, and what jobs are queued or running
 - Tells the resource manager when to run which job on the available resources





Partitions and 'Quality of Services'

- Defining where and how your job will run
- Partitions (basically a queue):
 - Resources/hardware
- QoS:
 - Tells what the limits or characteristics of a job should be
 - Maximum wall time (timelimit)
 - Number of nodes
- One partition might have multiple QoS
- A QoS might exist on multiple partitions





Available Partitions

Partition	Description	# of nodes	cores/node	GPUs/node
shas	General Compute (Haswell)	380	24	0
sgpu	GPU-enabled nodes	10	24	effectively 4
smem	High-memory nodes	5	48	0
sknl	Phi (Knights Landing) nodes	20	68	0



Quality of Service

QoS	Description	Maxwall	Max jobs/user	Max nodes/user
normal	Default QoS	Derived from partition	n/a	256
testing	For quick turnaround when testing	30 Min	1	2/user; max 12 cores/node
interactive	For interactive jobs (command or GUI)	4 Hours	1	1 core
long	For jobs needing longer wall times	7 D	n/a	20
condo	For groups who have contributed to the Summit condo	7 D	n/a	n/a

Useful Slurm Commands - sbatch

- sbatch: submit a batch script to slurm
- You can use a bunch of flag options in a batch script or on the command line
- Useful to put in script so have for future use

• Example:

```
sbatch test.sh
```

OR

```
sbatch --partition=shas test.py
```

http://slurm.schedmd.com/sbatch.html





SBATCH Options

http://slurm.schedmd.com/sbatch.html

#SBATCH <options> sbatch <options>

Allocation: --account=<account no>

Partition: --partition=<partition name>

• Sending emails: --mail-type=<type>

Email address: --mail-user=<user>

Number of nodes: --nodes=<nodes>

Number of tasks: --ntasks=<processes>

Quality of service: --qos=<qos>

Reservation: --reservation=<name>

• Wall time: --time=<wall time>

• **Job Name**: --job-name=<jobname>

• FYI: You do NOT actually type <> above - this designates something specific you as a user must enter about your job



Working on Summit

 If you use both Summit and Blanca, make sure you load the appropriate slurm module – if on a Login Node:

```
$ module load slurm/blanca
```

- \$ module load slurm/summit
- \$ ml slurm/summit #shorthand
- If on a compile node, already connected to Summit, this is not needed and will return an error.





Practice Examples

Submit Your First Job!

- Submit a slurm job with the following instructions:
 - 1. The job should run the Unix "hostname" command
 - The job will be submitted from a bash script named hostname_summit.sh
 - 3. The job will run on 1 node
 - 4. We will request 1 minute wall time
 - 5. Run using "testing" QOS
 - 6. Run on the shas-testing partition





Hostname_summit.sh

```
#!/bin/bash
#SBATCH --nodes=1  # Number of requested nodes
#SBATCH --time=0:01:00  # Max wall time
#SBATCH --qos=testing  # Specify debug QOS
#SBATCH --partition=shas-testing  # Specify Summit haswell nodes
#SBATCH --output=hostname_%j.out  # Rename standard output file
```

```
# purge all existing modules
module purge
hostname
```



Running the script

Load up the slurm module (probably not needed)

module load slurm/summit

Submit the job:

sbatch hostname summit.sh

Check output



Another slurm command

squeue

View information about jobs located in the slurm scheduling queue

• OPTIONS:

- User: -u <user list>
- Queues: --qos=<qos list>

• EXAMPLE:

http://slurm.schedmd.com/squeue.html





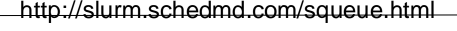
Your turn

- Submit a slurm job with the following instructions:
 - The job should run first the whoami command, then the Unix "sleep" command for 30 seconds, then the hostname command
 - Syntax for these Unix commands are below:

whoami

sleep 30

hostname







Your turn

- Submit a slurm job with the following instructions:
 - 1. The job will be submitted from a bash script named sleep.sh
 - 2. The job will run on 1 node
 - 3. Request a 1 minute wall time
 - 4. Run the job from the normal QOS
 - 5. Run the job from the Summit haswell partition
 - 6. Name your job sleep
 - 7. Email yourself the results at the end of the job run
 - Hint: This requires two SBATCH directives to do this see link at top of this slide

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Sleep.sh

```
#!/bin/bash
#SBATCH --nodes=1
#SBATCH --time=0:01:00
#SBATCH --gos=normal
#SBATCH --partition=shas
#SBATCH --output=sleep %j.out
#SBATCH --job-name=sleep
#SBATCH --mail-type=end
###SBATCH --mail-user=<user>@colorado.edu
# purge all existing modules
module purge
whoami
sleep 30
hostname
```

```
# Number of requested nodes
# Max walltime
# Specify normal QOS
# Specify Summit GPU nodes
# Rename standard output file
# Job submission name
# Email you when the job ends
# Email address to send to
```

Running an external script

- Let's run a Matlab program
- We will run the bash script matlab.sh
- This script calls and runs matlab_tic.m

Running the script

Submit the job:

sbatch matlab.sh

Check output

Matlab.sh

```
#!/bin/bash
#SBATCH --nodes=1
                                        # Number of requested nodes
#SBATCH --time=0:02:00
                                        # Max walltime
#SBATCH --qos=testing
                                        # Specify debug QOS
# #SBATCH --partition=shas
                                        # (disabled) specify Summit haswell nodes
#SBATCH --output=matlab %j.out
                                        # Output file name
# purge all existing modules
module purge
# Load a specific Matlab module
module load matlab/R2016b
# Run matlab without a GUI
matlab -nodisplay -nodesktop -r "clear; matlab tic;"
```



Your turn

- Submit a slurm job with the following instructions:
 - 1. Create an R program called R_program.R that creates a vector called "planets" and then list the planets in the vector
 - Syntax: planets -> planets <- c("Mercury", "Venus", "Earth", "Mars", "Jupiter", "Saturn", "Uranus", "Neptune", "Pluto")
 - Print off the vector
 - Syntax: planets
 - 3. Create a bash script called R code.sh that runs the R script
 - Syntax: Rscript R program.R
 - 4. The job will run on 1 node
 - 5. We will request a 1 minute wall time
 - 6. Specify the debug QOS
 - 7. Specify the shas partition
 - 8. The output will be put in a file called R_code_%j.out
 - 9. Don't forget to load the R module!





Solution - R_code.sh

```
#!/bin/bash
#SBATCH --nodes=1
#SBATCH --time=0:01:00
#SBATCH --gos=debug
#SBATCH --partition=shas
#SBATCH --output=R code %j.out
# purge all existing modules
module purge
# Load the R module
module load R/3.3.0
# Run R Script
Rscript R program.R
```

```
# Number of requested nodes
# Max walltime
# Specify debug QOS
# Specify Summit haswell nodes
# Output file name
```

Solution - R_program.R

```
#Simple R code example by Shelley Knuth
(shelley.knuth@colorado.edu)
# Create vector
planets <- c("Mercury", "Venus", "Earth", "Mars",</pre>
"Jupiter", "Saturn", "Uranus", "Neptune", "Pluto")
# Print off vector
planets
```



Interactive jobs!

- Sometimes we want our job to run in the background
- Sometimes we want to work in program in real time
- For example, Matlab
- Let's run an interactive Matlab job

Interactive job using X

- In this example we'll use X windows to access the Matlab GUI
- To do this, we are going to log out and log back in
 - Only necessary for demo
 - Need to add something to the sign in process
- For Mac Users:

```
ssh -X username@login.rc.colorado.edu
```

- For Windows Users, must set up X-forwarding through your SSH client program
- Also must have an X-server package on your laptop
 - Xming for Windows or XQuartz for Mac





Interactive job

- To work with Matlab interactively, we're going to request some time from the supercomputer
- When the resources become available then we will start up Matlab
- Commands to run:

```
module load slurm/summit
sinteractive --qos=debug --time=00:05:00
```

Once we receive a prompt, then:

```
module load matlab
matlab
```

Once we finish we must exit!



Questions?

- Email rc-help@colorado.edu
- Twitter: @CUBoulderRC

 Link to survey on this topic: http://tinyurl.com/curc-survey18

 Slides: https://github.com/ResearchComputing/New_User_Seminar



