# Research Computing New User Seminar

Shelley Knuth <a href="mailto:shelley.knuth@colorado.edu">shelley.knuth@colorado.edu</a>

www.rc.colorado.edu

#### Slides:

https://github.com/ResearchComputing/Final\_Tutorials/tree/master/RC\_Access\_Specific\_Topics/New\_User\_Seminar

## Outline

- What is RC?
- Steps to get access to our systems
  - Accounts
  - Two-factor authentication
  - Allocations
  - Logging in
- Navigating our systems
  - 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:
  - Janus/Summit
  - PetaLibrary

## What Would I Use Summit For?

- Research Computing is more than just Summit
- But it is what we are most known for
- So 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
  - Large visualization jobs
  - High memory jobs
- Not a place for:
  - Large data storage

# Hardware - Summit Supercomputer

- 475 compute nodes (Intel Xeon Haswell)
- 24 cores per node
- 11,400 total cores
- Omni-Path network
- 1.2 PB scratch storage
- GPFS File system



# 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 (planned summer 2017)
  - 20 nodes
  - Intel Xeon Phi

## How To Access RC Resources?

- 1. Get an account
- 2. Set up two factor authentication
- 3. Set up an allocation
  - Don't need for Blanca or PetaLibrary
- 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
- CSU Users:
  - Request an account through CSU
  - Talk to Richard Casey
  - Then get an RC user account
- RMACC Users:
  - We are working on a login system through XSEDE
  - Not operational yet

## Setting up Two-Factor Authentication

- Two factor authentication is required to access our system
- Require this to provide an extra level of authentication
- Two methods for achieving this:
  - Duo
    - Access through a smart phone app
  - Vasco OTP (one time password)
    - Physical device

## **Duo Authentication**

- Once you get an account, contact <u>rc-help@colorado.edu</u> to request a Duo invitation
- Once you get the invitation, you'll get a series of steps to complete Duo enrollment
- RC supports Duo "push" and "phone call" for authentication
- Greatly prefer "push"

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## Vasco Authentication

- Physical device that generates a new password every 30 seconds
- To get the device, go to the IT Service Center or request a time to pick one up from Research Computing
- You will need to show an ID to get a device
- First, register the device
  - Otp.colorado.edu
- You will set a four digit password

## Allocations

- You will need an allocation to use any of our resources
  - The exception is if you will ONLY be using the PetaLibrary or Blanca
- Currently, to request an allocation please email <u>rc-help@colorado.edu</u> and ask for a General account
  - Need to provide a few sentences on your project
- In the future, we will have a place on our website to submit a more formal request
- Once you have some benchmarks, you will want to move to a project allocation

## Why Do I Need An Allocation?

- I have an account why do I need an allocation?
  - An account validates you are eligible to use RC 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 a general allocation:
    - Gives you higher priority in the system

## 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 over last 3-4 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 3-4 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

- Need an allocation? Plan to run on Summit?
- Make a request now!
- Include 2-3 sentences describing your proposed usage
- Email <u>rc-help@colorado.edu</u>

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# 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
- From there, you can access our other resources

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## RC Resource Access

- To login to an RC login node: ssh username@login.rc.colorado.edu
- If logging in with Duo, your enter your password as: duo:identikey password
- If logging in with Vasco, your enter your password as:

  Pin+six-digit number on Vasco

## 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?

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# 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
  - Script or code editing, minor compiling
  - Job submission
- Compile nodes
  - Where you compile code
- Compute/batch nodes
  - This is where jobs that are submitted through the scheduler run
  - Intended for heavy computation

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## 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
- Much larger depends on system
- Files purged around 90 days

# What Belongs Where?

- /home
  - Scripts
  - Code
  - Very small files
  - Inappropriate for sharing files with others
  - Inappropriate for job output
- /projects
  - Code/files/libraries relevant for any software you are installing (if you want to share files with others)
  - Mid-level size input files
  - Appropriate for sharing files with others
  - Inappropriate for job output
- /scratch/summit
  - Output from running jobs
  - Large files
  - 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
- 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

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# Setting Up Globus

- Create an account at Globus.org
- Make your personal computer an endpoint
- Transfer data
- www.globus.org

We will come back to this at the end if we have time

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## 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 can load modules to prepare your environment for using software
    - Set any environment variables
    - Set environment so application can find appropriate libraries, etc.

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## Important Things to Know About Modules

- Some modules might require a specific hierarchy to load
  - For some modules, you may need to specify a specific version
    - For example, ml R/3.3.0
  - For other modules, you may be able to be more generic
    - For example, ml 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 ml avail
- To set up your environment to use a software package, type ml
   <package>/<version>

## Job Submission

# Running Jobs

- What is a "job"?
- Interactive jobs
  - Work interactively at the command line of a compute node
- Batch jobs
  - Submit job that will be executed when resources are available
  - Create a text file containing information about the job
  - Submit the job file to a queue

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## 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'

- There are several ways to define where 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
    - Number of nodes
- One partition might have multiple QoS
- A QoS might exist on multiple partitions

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## **Partitions**

- By default, jobs will run on the general compute (Haswell) nodes
- Recommend specifying a partition
- Do this using the -p partition flag

## **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 - [not currently available]	20	68	0

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# Quality of Service

QoS	Description	Maxwall	Max jobs/user	Max nodes/user
normal	Default QoS	Derived from partition	n/a	256
debug	For quick turnaround when testing	1 H	1	32
long	For jobs needing longer wall times	7 D	n/a	20
condo	For groups who have purchased Summit nodes	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 -p=shas test.sh
```

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

## **SBATCH Options**

```
#SBATCH <options> sbatch <options>
```

- Allocation: -A=<account\_no>
- Partition: -p=<partition name>
- Sending emails: --mail-type=<type>
- Email address: --mail-user=<user>
- Number of nodes: -N <nodes> or --nodes=<nodes>
- Quality of service: --qos=<qos>
- Reservation: --reservation=<name>
- Wall time: --time=<wall time>
- Job Name: -J <jobname> or -job-name=<jobname>
- FYI: You do NOT actually type <> above this designates something specific you as a user must enter about your job

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# Working on Summit

- Right now we are in a transition phase between Summit and Janus
- You should be running all your jobs on Summit
- To ensure this, until Janus goes offline entirely, make sure you load the slurm module on Summit

ml slurm/summit

After you run this command you can run sbatch to submit jobs

### Blanca

- If you are a Blanca user, you need an RC account, but not an allocation
- To run jobs as a Blanca user, once you've logged into a login node, load the Blanca slurm module

ml slurm/blanca

- Only certain users have access to Blanca paid service
- If you are unsure, you can ask your advisor or RC
  - But likely if you are unsure you don't have access

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# 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

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# Practice Examples

### Submit Your First Job!

- Submit a slurm job with the following instructions:
- The job should run the Unix "hostname" command
- The job will be submitted from a bash script named hostname\_summit.sh
- The job will run on 1 node
- We will request 1 minute wall time
- 5. Run from the debug QOS
- 6. Run on the shas partition
- 7. Send the output to a file called hostname.out
- 8. User the new\_user reservation
  - This is only for this workshop

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# Contents of Batch Script

Bash Script hostname\_summit.sh:

```
#!/bin/bash
#SBATCH -N 1
                                 # Number of requested nodes
#SBATCH --time=0:01:00
                                 # Max walltime
#SBATCH --qos=debug
                                 # Specify debug QOS
#SBATCH --partition=shas
                                 # Specify Summit haswell nodes
#SBATCH --output=sleep.out
#SBATCH --reservation=new user
                                 # Reservation name
```

hostname

# Running the script

Load up the slurm module
 ml 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: -q <qos\_list> or --qos=<qos\_list>

#### • EXAMPLE:

squeue -q debug

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

### Your turn

- Submit a slurm job with the following instructions:
- 1. 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

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### 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. We will request a 1 minute wall time
- 4. The output will be put in a file called sleep.out
- 5. Email yourself the results at the end of the job run
  - Hint: Requires two SBATCH options to do this
- 6. Use the new\_user reservation
- 7. Specify the shas partition
- 8. Specify the debug QOS

# Solution – contents of script

```
#!/bin/bash
#SBATCH -N 1
#SBATCH --time=0:01:00
#SBATCH --output=sleep.out
#SBATCH --mail-type=end
#SBATCH --mail-user=ralphie@colorado.edu
#SBATCH --reservation=new user
whoami
sleep 30
hostname
```

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# Running an external script

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

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# Running the script

Submit the job:

sbatch matlab.sh

Check output:

nano Matlab.out

# Contents of Batch Script

#### Bash Script matlab.sh:

```
#!/bin/bash
#SBATCH -N 1  # Number of requested nodes
#SBATCH --time=0:02:00  # Max walltime
#SBATCH --job-name=Matlab  # Job submission name
#SBATCH --output=Matlab.out  # Output file name
#SBATCH --qos=debug  # Specify debug QOS
#SBATCH --partition=shas  # Specify Summit haswell nodes
#SBATCH --reservation=new_user  # Reservation name
```

```
# Run matlab without a GUI matlab -nojvm -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")
- 2. Print off the vector
  - Syntax: planets
- 3. Create a bash script called R code.sh that runs the R script
  - Syntax: Rscript R\_code.sh
- 4. The job will run on 1 node
- 5. We will request a 1 minute wall time
- 6. The output will be put in a file called R.out
- 7. Name your job R\_code
- 8. Specify the debug QOS
- 9. Specify the shas partition
- 10. Use the new user reservation
- 11. Don't forget to load the R module! (This is going to be a bit tricky follow the instructions!

## Solution – R\_code.sh

```
#!/bin/bash
#SBATCH -N 1
                                  # Number of requested nodes
#SBATCH --time=0:01:00
                                  # Max walltime
#SBATCH --job-name=R code
                                  # Job submission name
#SBATCH --output=R code.out
                                  # Output file name
#SBATCH --gos=debug
                                  # Specify debug QOS
#SBATCH --partition=shas
                                  # Specify Summit haswell nodes
#SBATCH --reservation=new user
                                  # Reservation name
Rscript R program.R
```

Before all of this, you must do:

```
ml intel
ml R/3.2.0
```

# Solution – R\_program.R

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

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# 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

- NEVER EVER EVER EVER EVER
- EVER EVER EVER EVER EVER EVER EVER
- Run Matlab from the command line without running an interactive job first
- BAD USER!

# Interactive job

- 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:

```
ml slurm/summit
sinteractive --reservation=new_user
```

• Once we receive a prompt, then:

```
ml matlab
matlab
```

Once we finish we must exit!

# Setting Up Globus

- Create an account at Globus.org
- Make your personal computer an endpoint
- Transfer data
- www.globus.org

### Questions?

- Email rc-help@colorado.edu
- Twitter: @CUBoulderRC
- Link to survey on this topic:

http://tinyurl.com/curc-survey16

Slides:

https://github.com/ResearchComputing/Final\_Tutorials/tree/master/RC\_Access\_Specific\_Topics/New\_User\_Seminar