Lab 6: SCF and Parallel Computing

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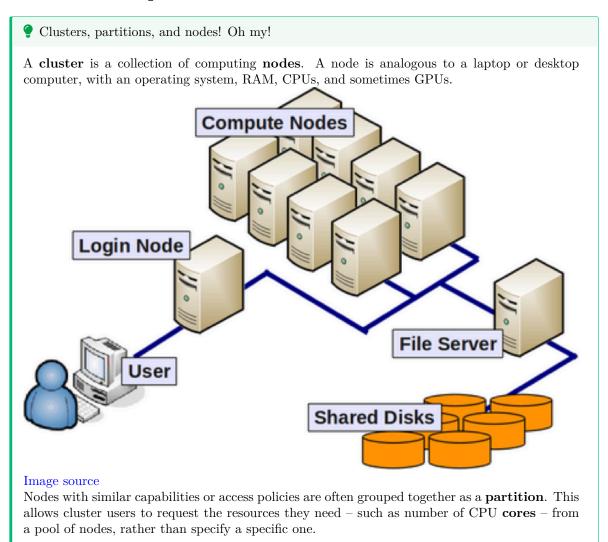
Overview

Today we'll discuss the Statistical Computing Facility's (SCF) cluster, which is administrated by the Department of Statistics. Much of this material comes directly from the SCF's documentation; see the links below for more details:

- SCF homepage
- SCF Linux cluster
- SCF FAQs and How-Tos
- SCF Parallel processing in Python tutorial
- How to get help on the SCF

• SCF JupyterHub

SCF cluster capabilities and hardware



The SCF cluster has 1064 cores, across 7 partitions with more than 26 nodes, but for this class we'll use the low partition. It's the default, so you won't need to do anything special to use it for your jobs (we'll explain how to submit jobs later on).

The ${\tt low}$ partition has the following resources:

- 8 nodes
- 32 cores per node

• 256 GB RAM per node

The SCF Linux cluster uses a batch job scheduler called Slurm, commonly used by large computer centers in both academia and industry. Although the SCF cluster has much greater computational capacity than our personal computers, it is a shared resource with many users. With Slurm, the cluster is able to schedule jobs asynchronously, balancing resource allocations so that everyone gets a turn. This also means that when you submit a job, it may not run until many minutes or hours later. Be sure to give yourself enough time!

Your ~/ on the SCF

Nodes in the SCF cluster use the Linux distribution Ubuntu. Every user has a private home directory which, as usual on Linux-based OSes, has the shortcut ~/. In this section, we'll give an overview of access, disk space, and remote file transfers to and from your SCF home directory.

Logging in

The SCF has a number of login nodes which you can access via ssh.

Note

For info on using ssh (including on Windows), see here.

For example, I'll use ssh to connect to the gandalf node:

```
james@pop-os:~$ ssh <scf-username>@gandalf.berkeley.edu
The authenticity of host 'gandalf.berkeley.edu (128.32.135.47)' can't be established.
ED25519 key fingerprint is SHA256:io5uUQGbCZie78mF+UUZ5guDK29JXQGQ6LVB129UoUo.
This key is not known by any other names
Are you sure you want to continue connecting (yes/no/[fingerprint])? yes
Warning: Permanently added 'gandalf.berkeley.edu' (ED25519) to the list of known hosts.
```

Notice that upon first connecting to a server you haven't visited there is a warning that the "authenticity of the host ... can't be established". So long as you have typed in the hostname correctly (gandalf.berkeley.edu, in this case), and trust the host (we trust the SCF!) then you can type yes to add the host to your known hosts file (found on your local machine at ~/.ssh/known_hosts).

You'll then be asked to enter your password for the SCF cluster. For privacy, you won't see anything happen in your terminal when you type it in, so type carefully (you can use Backspace if you make a mistake) and press Enter when you're done. If you were successful, you should see a welcome message and your shell prompt, like:

```
<scf-username>@gandalf:~$
```

To get your bearings, you can type pwd to see where your home directory is located on the SCF cluster filesystem:

```
<scf-username>@gandalf:~$ pwd
/accounts/grad/<scf-username>
```

Your home directory is likely also in the /accounts/grad/ directory, as mine is.

Other login nodes

Important

Don't run computationally intensive tasks on the login nodes!

They are shared by all the SCF users, and should only be used for non-intensive interactive work such as job submission and monitoring, basic compilation, managing your disk space, and transferring data to/from the server.

If for some reason gandalf is not working for you, the SCF has a number of nodes which can be accessed from your local machine with commands of the form ssh <scf-username>@<hostname>.berkeley.edu. Currently, these are:

- aragorn
- arwen
- dorothy
- gandalf
- gollum
- hermione
- quidditch
- radagast
- shelob

Note that all those machines are stand-alone login nodes. After logging in and launching a job, the job will be run on a cluster node. Usually, users cannot ssh into cluster nodes and the process of launching a job is via slurm from one of the login machines. However, if one has a job running on a cluster node, that specific node becomes accessible via ssh. This can be helpful with troubleshooting.

Disk space

Your home directory has a limited amount of disk space; you can check how much you have used and available using the quota command, which will show your home directory usage and quota on the line for the accounts filesystem.

```
<scf-username>@gandalf:~$ quota
FILESYSTEM USE QUOTA %
accounts 5.29 G 20 G 26.5
```

The accounts filesystem is accessible from all the nodes on the SCF cluster. This means that regardless of which login or compute node you use, you will have access to the files in your home directory. Moreover, your home directory is backed up regularly, so you are protected from accidental data loss.

If you're running out of space, you should try to selectively delete large files that you no longer need. See here for some tips on finding large files. If all else fails, you can request additional space or request access the /scratch filesystem. The latter is a good place for large datasets, but note that /scratch is not backed up, unlike your home directory. See the previous link for more info. Note though that disk space should not be a concern for Stat 243, but being aware of /scratch can be relevant for Stat graduate students (MA and PhD) working on projects in the future.

For temporary files (e.g., intermediate results of a computation that you don't need to store for later), every machine has a /tmp filesystem. However, /tmp is always linked to the specific machine you are on, meaning that if you put something in /tmp on gandalf and then later go to aragorn, you won't find your files in the /tmp directory there. If you go back to gandalf, you will likely find them again, but /tmp is automatically wiped when a machine reboots, so only use it for files you don't care about preserving!

Data transfer: SCP / SFTP

We can use the scp and sftp protocols to transfer files to and from any login node on the SCF cluster. scp is a shell program that should be available by default on macOS and Linux, while on Windows you can use WinSCP. WinSCP can also do sftp transfers, or you can use FileZilla on any platform for sftp. Both WinSCP and FileZilla have a GUI that allows you to drag and drop files between your local machine and a remote host.

The syntax for scp is scp <from-place> <to-place>, and you'll typically use scp on your local machine. For example, we'll show how to transfer the file data.csv (you can find it here):

To SCF while on your local machine

```
# transfer to my home directory without renaming the file
scp data.csv <scf-username>@gandalf.berkeley.edu:~/

# transfer to the data/ subdirectory and rename the file
scp data.csv <scf-username>@gandalf.berkeley.edu:~/data/new_name.csv

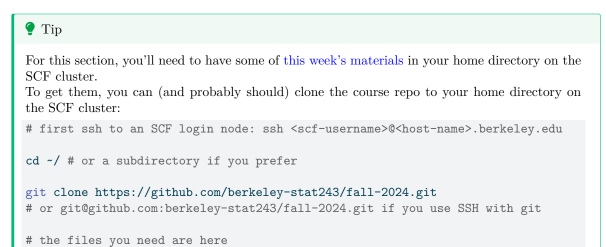
# transfer to the gandalf-specific /tmp/ directory
scp data.csv <scf-username>@gandalf.berkeley.edu:/tmp/
```

From SCF while on your local machine

```
# now <from-place> is a path in my home directory on the SCF
scp <scf-username>@gandalf.berkeley.edu:~/data/new_name.csv ~/Desktop/data_scf.csv
```

For more information, see here. In particular, if you have a very large dataset to transfer, Globus is a better option than either scp or sftp.

Running jobs on the SCF



As mentioned before, the SCF cluster uses Slurm to schedule jobs. We'll learn some Slurm commands now.

Non-interactive jobs: sbatch

cd fall-2024/labs/lab6-scfparallel/

```
This is the best option when you have a long-running job where no
```

The first command is sbatch. To use sbatch, you will first create a bash script (you can do this on the SCF cluster if you know how to use a shell text editor like nano, or just do it locally and transfer the file to the SCF via one of the methods discussed before).

In the course repo, you will find an example bash script called submit.sh. Here's what it contains:

```
#!/bin/bash

#################

# SBATCH OPTIONS
#############

#SBATCH --job-name=example # job name for queue (optional)
#SBATCH --partition=low # partition (optional, default=low)
#SBATCH --error=ex.err # file for stderr (optional)
#SBATCH --output=ex.out # file for stdout (optional)
#SBATCH --time=00:01:00 # max runtime of job hours:minutes:seconds
#SBATCH --nodes=1 # use 1 node
```

The example.sh script creates a file called testFile.txt and appends some output to that file. Note that since submit.sh is itself a bash script, you could have just put the contents of example.sh directly in submit.sh, but it is good practice to keep your computational code separate from your Slurm batch submission scripts.

Now we can submit the job using the Slurm command sbatch:

```
sbatch submit.sh
```

Once you run that, your job will be queued. To check it's status, you can use squeue:

```
squeue -u <scf-username> # use your username, not mine
```

And now you wait. Go get a cup of coffee or get some rest! Then check back later.

Interactive jobs



Interactive jobs are the best option when you need to do many short but computationally intensive tasks which require your hands at the keyboard.

srun



This is the best option when you want to work at the Linux command line and don't need a GUI, e.g. when debugging components of larger non-interactive job.

To run an interactive job via Slurm, you can use the srun command:

```
# ssh -Y <scf-username>@gandalf.berkeley.edu
# 10 minute time limit
srun -t 00:10:00 -N 1 -n 1 -c 1 --pty /bin/bash
# uses resources already made available from the previous srun command
srun --pty --x11=first matlab # interactive matlab job
```

Here we used the -Y flag to ssh to run software with a GUI, such as MATLAB, which allows the GUI to open in our local machine while still running computations on the SCF cluster. For this to work, you'll need X server software on your own machine to manage the graphical windows. For Windows, your options include eXceed or Xming and for Mac, there is XQuartz.

JupyterHub for RStudio or Jupyter Notebooks



Tip

This is the best option when you want to interact with the RStudio or Jupyter GUI while doing your computations or debugging.

The SCF's JupyterHub is another resource for interactive computing on the SCF, allowing you to use RStudio or Jupyter on one of the cluster's computing nodes.

Specifying resources

If you are submitting a job that uses multiple cores or nodes, you may need to carefully specify the resources you need. The main key flag for use in your job script is:

• --ntasks-per-node: indicates the number of tasks (i.e., processes) one wants to run on each node

The value passed to --ntasks-per-node becomes available through the SLURM_NTASKS environment variable, which can be retrieved in python via:

```
import os
ntasks = os.getenv("SLURM_NTASKS")
```

Specifying resources: alternatives

The --ntasks-per-node flag is convenient as it does not require the user to know the actual number of CPUs on each node. It is generally a good choice if one is simply running parallel code via multiple processes on one node. However, there are alternatives flags that can be used in conjunction for finer-grained control:

- --nodes (or -N): indicates the number of nodes to use
- --ntasks (or -n): indicates the number of tasks to run, not necessarily on the same node
- --cpus-per-task (or -c): indicates the number of CPUs to be used for each task

Since specifying --ntasks does not guarantee that all the cores will be on a single node, this can cause problems in the common case of code that works on one node but not multiple nodes. If one wants to do parallelization at multiple levels (e.g., multiple processes, each process using multiple threads, such as for linear algebra), then one would use both --ntasks-per-node and --cpus-per-task. In general --cpus-per-task will be 1 except when running threaded code.

The SLURM_NTASKS environment variable is set by Slurm when the job starts running, and therefore can be accessed within your jobs. In addition to SLURM_NTASKS here are some other variables that

may be useful: SLURM_CPUS_ON_NODE, SLURM_CPUS_PER_TASK, SLURM_NODELIST, SLURM_NNODES. An explanation of each one of those can be found on slurm's manual (man sbatch).

Monitoring your jobs

As we saw, the basic command for seeing what is running on the system is squeue:

```
squeue # this shows all the queued jobs!
squeue -u SCF_USERNAME # this shows your jobs
```

To see what nodes are available in a given partition, you can use sinfo:

```
sinfo -p low
```

Finally, you can cancel a job with scancel.

```
# you can find your job ID using `squeue`
scancel YOUR_JOB_ID
```

See the "How to Monitor Jobs" section of the SCF's documentation on this page for some more info.

Exercise

You can either do 1. on your local machine and transfer the file to the SCF cluster, or you can do it remotely. 2 - 4 should be done on the SCF.

- 1. Create a submission submitBoot.sh that will execute the code in boot_proc.py. To do this I suggest copying the text from submit_py.sh and updating a couple lines:
- Change job-name to whatever you want to call this job.
- Change time to 00:10:00 just in case your code runs longer than one minute.
- Change the --cpus-per-task to 3. This is where you are telling Slurm to execute in parallel on 3 CPUs.
- Change the last line python boot_serial.py.
- 2. Call sbatch submitBoot.sh to execute the python script boot_proc.py.
- 3. Check that the execution worked by examining the py.out file.
- 4. Note that the number of Dask workers in boot_proc.py and boot_dist.py is hard-coded. In many cases, it would be better for this value to be set dynamically based on an appropriate Slurm environment variable. Make this small adjustment to those files and verify it works as intended.

Acknowledgements

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