SCF Computing Cluster

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Introduction

Much of this material comes from the SCF documentation at the SCF homepage.

Outline

This training session will cover the following topics:

- System capabilities and hardware
 - SCF computing nodes
 - Disk space
- Logging in, data transfer, and software
 - Logging in
 - Data transfer: SCP/SFTP
 - Software modules
- Submitting and monitoring jobs
 - Accounts and partitions
 - Interactive jobs
 - Basic job submission
 - Parallel jobs
 - Monitoring jobs and cluster status
- Exercise

System Capabilities and Hardware

- The SCF Linux cluster contains 352 cores, separated into 12 nodes and two partitions. The low partition is mainly made up of older computers and is slower, while the high partition are newer machines. By default computations are done on the low partition nodes, which is more than sufficient for the work you will do in this class.
 - Low partition
 - * 8 nodes
 - * 256 total cores
 - $\ast\,$ each node contains 32 cores and 256 GB of RAM
 - high partition
 - * 4 nodes
 - * 96 total cores
 - \ast each node contains 24 cores and 128 GB of RAM

SCF Computing Nodes

The SCF cluster is has several compute nodes, which you can list using sitehosts compute. They are also listed on this page.

The nodes are divided into 3 partitions, the two listed above and a GPU partition, with restrictions associated with each. Any job you submit must be submitted to a partition to which you have access.

For the puposes of this class, we only have access to the *low* partition.

Disk Space

Every account is given 5 GB of disk space. (This can be queried using quota <account_name>) This includes your home directory and a /tmp directory tied to your account. Space in the /scratch partition is available by request only.

Logging in

There login servers include: arwen, beren, bilbo, gandalf, gimli, legolas, pooh, radagast, roo, shelob, springer, treebeard

To login, in your UNIX terminal type:

```
ssh SCF_USERNAME@LOGINSERVER.berkeley.edu
```

Then enter your password for your SCF account. One can then navigate around and get information using standard UNIX commands such as ls, cd, du,df, etc.

Logging out

To logout you type logout in the command line.

Data transfer: SCP/SFTP

We can use the scp and sftp protocols to transfer files from any login node on the SCF cluster.

For example, we show how to transfer the file data.csv.

To transfer that to your directory on the SCF cluster, the syntax for scp is scp <from-place> <to-place>

Below are some examples of transferring this file to various locations on the remote machine (your home directory, a folder called data in your home directory, and the tmp folder)

```
# to SCF, while on your local machine
scp data.csv zoe_vernon@arwen.berkeley.edu:~/
scp data.csv zoe_vernon@arwen.berkeley.edu:~/data/new_name.csv
scp data.csv zoe_vernon@arwen.berkeley.edu:/tmp/
```

Here is how to transfer data from the SCF to your local machine, while on your local machine. This will transfer the new_name.csv file to your Desktop with the name data_scf.csv

```
# from SCF, while on your local machine
scp zoe_vernon@arwen.berkeley.edu:~/data/new_name.csv ~/Desktop/data_scf.csv
```

One program you can use with Windows is WinSCP, and a multi-platform program for doing transfers via SFTP is FileZilla. After logging in, you'll see windows for the SCF filesystem and your local filesystem on your machine. You can drag files back and forth.

Submitting jobs: accounts and partitions

All computations are done by submitting jobs to the scheduling software that manages jobs on the cluster, called SLURM (Simple Linux Utility for Resource Management).

There is nothing you must specify to submit jobs to the SCF cluster, however, there are several options that are good form (more on this below).

Interactive jobs

You can do work interactively.

For this, you may want to have used the -Y flag to ssh if you are running software with a GUI such as MATLAB.

```
# ssh -Y SCF_USERNAME@arwen.berkeley.edu
srun --pty /bin/bash  # interactive bash job
srun --pty --x11=first matlab  # interactive matlab job
```

To display the graphical windows on your local machine, 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.

This is not something I have much experience with, but if you are interested instructions are available here.

Submitting a batch job

Or you can submit a job to run in the background.

Let's see how to submit a simple job. If your job will only use the resources on a single node, here's an example job submission script, exSub in the section folder:

#!/bin/bash

```
####################
# SBATCH OPTIONS
#####################
#SBATCH --job-name=example
                                        # job name fore queue, default may be u$
#SBATCH --partition=low
                                         # high/low/gpu, default if empty is low
                                         # error file, default if empty is slurm$
#SBATCH --error=ex.err
                                         # standard out file, no default
#SBATCH --output=ex.out
#SBATCH --time=00:01:00
                                         # hours:minutes:seconds
##SBATCH --nodes=1
                                         # only use 1 node, MPI option
##SBATCH --ntasks=1
                                         # how many tasks to start
                                         # number of cores to use, multi-core/mu$
##SBATCH --cpus-per-task=1
#####################
# What to run
#####################
./example.sh
```

This script runs the following bash script, example.sh in the section folder:

```
#!/bin/bash
# set file name
fileName="testFile.txt"
# create file
touch $fileName
# intro message
echo "I'm your new test script!" >> $fileName
echo >> $fileName
echo >> $fileName
# fill it with some things
for i in \{1...10\}
do
        echo $i >> $fileName
done
# finish it out
echo >> $fileName
echo >> $fileName
echo "All Finished!" >> $fileName
Now let's submit and monitor the job:
sbatch exSub
squeue -u <SCF_USERNAME>
```

Parallel job submission

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

- --nodes (or -N): indicates the number of nodes to use
- --ntasks-per-node: indicates the number of tasks (i.e., processes) one wants to run on each node
- --cpus-per-task (or -c): indicates the number of cpus to be used for each task

In addition, in some cases it can make sense to use the <code>--ntasks</code> (or <code>-n</code>) option to indicate the total number of tasks and let the scheduler determine how many nodes and tasks per node are needed. In general <code>--cpus-per-task</code> will be 1 except when running threaded code.

When setting up parallel R code, you can find out how many cores there are on the node assigned to you with:

```
ncores <- Sys.getenv("SLURM_CPUS_ON_NODE")</pre>
```

In addition to SLURM_CPUS_ON_NODE here are some of the variables that may be useful: SLURM NTASKS, SLURM CPUS PER TASK, SLURM NODELIST, SLURM NNODES.

Monitoring jobs and the job queue

The basic command for seeing what is running on the system is squeue:

```
squeue -u SCF_USERNAME
```

To see what nodes are available in a given partition:

```
sinfo -p low
sinfo -p high
sinfo -p gpu
```

You can cancel a job with scancel. The job ID can be found on the left hand side when looking at the queue.

```
scancel YOUR_JOB_ID
```

The SCF has some tips about monitoring your job.

Exercise

For the exercise in section we want you to get comfortable copying files from your local computer and running parallel computations on the SCF cluster. For this we will use an example from this webpage to compute bootstrap estimates of a coefficient in a logistic regression and compare the timing for a parallelized and non-parallelized version of the code. Your job is execute this code on the SCF.

To accomplish this you will need to complete the following tasks.

Running on SCF

- 1) Create a submission script bootSub that will execute the code in boot.R. To do this I suggest copying the text from exSubR and updating a couple lines:
 - Change job-name to whatever you want to call this job
 - Change rEx.err and rEx.out to boot.err and boot.out. Note these files can be useful for fixing bugs, particuarily when excecuting shell scripts. These files show errors and output printed to the terminal.
 - 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 nodes.
 - Change the last line R CMD BATCH --no-save boot.R boot.Rout. The boot.Rout file will contain all of code, messages, errors, etc. from the R session.
- 2) Transfer the files boot.R and bootSub to your SCF account. This can be done using scp before logging onto the SCF cluster (see examples above)
 - Note, there are other ways to transfer files. For example, one other option would be to clone the repo for this class onto your SCF account after logging in.
- 3) Call sbatch bootSub to excecute the R script boot.R.
- 4) Check that the excecution worked by examining the .Rout file.
- 5) If you have time:
 - Update parallel code to be execute with the future package to compare the results.