Linux and the HPC Environment

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Executive Director of Research Computing
September 28th, 2023
Research Computing / HPC Training

1. Thursday – September 14th 5-6pm Zoom – Introduction to Linux command line
2. Thursday – September 21st 5-6pm Zoom – Introduction to HPC at W&M/VIMS
3. Thursday – September 28th 5-6pm Zoom – Linux in the HPC environment

- [https://www.wm.edu/it/rc](https://www.wm.edu/it/rc) - shortcut to RC/HPC web
- [hpc-help@wm.edu](mailto:hpc-help@wm.edu) - help desk email – best way to reach us
- [hpc-announce@wm.edu](mailto:hpc-announce@wm.edu) - list for announcements from RC (auto-subscribed)
- Office hours: by appointment – email [hpc-help@wm.edu](mailto:hpc-help@wm.edu)
SSH/SCP/SFTP

**ssh** – is the standard app for connecting to a remote computer with Linux
**scp** – is the standard app for copying files from one site to another
**sftp** – more convenient for multiple hops

Both are easy within Linux. Mac also supports both via **terminal**. Windows users can use **powershell** to do this or use ssh client program (**putty**) and a separate scp client (**WinSCP**).

**Example Commands**

- **ssh to bora**
  - `ssh ejwalt@bora.sciclone.wm.edu`

- **ssh to bora through bastion host**
  - `ssh -J ejwalt@bastion.wm.edu ejwalt@bora.sciclone.wm.edu`

- **scp to bora**
  - `scp file1 ejwalt@bora.sciclone.wm.edu:`

- **scp from bora**
  - `scp ejwalt@bora.sciclone.wm.edu:file1`

- **sftp double hop**
  - `sftp -J ejwalt@bastion.wm.edu ejwalt@bora.sciclone.wm.edu`

See [https://www.wm.edu/offices/it/services/researchcomputing/using/connecting/](https://www.wm.edu/offices/it/services/researchcomputing/using/connecting/) - connecting/logging in to HPC
[https://www.wm.edu/offices/it/services/researchcomputing/using/xfers/](https://www.wm.edu/offices/it/services/researchcomputing/using/xfers/) - transferring files to/from HPC
Globus for File Transfers

**Globus** is another possibility: [https://www.globus.org/data-transfer](https://www.globus.org/data-transfer)

**Globus connect personal**: [https://www.globus.org/globus-connect-personal](https://www.globus.org/globus-connect-personal)

- Can use Globus to transfer files to/from and within clusters (Main and VIMS campus).
- Using Globus Connect Personal - attach your own computer to the Globus site and use the Globus GUI
- Available for Windows, Mac and Linux

HPC guide: [https://www.wm.edu/offices/it/services/researchcomputing/using/xfers/globus/personal/](https://www.wm.edu/offices/it/services/researchcomputing/using/xfers/globus/personal/)

Recent talk on using Globus in HPC: [https://www.wm.edu/offices/it/services/researchcomputing/using/tutorials/](https://www.wm.edu/offices/it/services/researchcomputing/using/tutorials/)
File/Folder Permissions

**umask** – controls the permissions set for new files. The default umask on all HPC systems is:

1 [bora] umask 0077
2 [bora] umask -S
   u=rwx,g=,o=

So, all new files get **u=rw**, and folders get **u=rwx** and nothing else set.

Other reasonable umasks are (set in .cshrc/.bashrc at end)

027 (u=rwx, g=rx, o=) - members of group can read and enter folders
022 (u=rwx, g=rx, o=rx) - everyone can read and enter folders

Must have permissions correct the whole way through a path:
1 [bora] pwd
   /sciclone/home/ewalter
2 [bora] ls -ld /sciclone
   drwxr-xr-x 20 root root 279 Apr 22 15:49 /sciclone
3 [bora] ls -ld /sciclone/home
   drwxr-xr-x 89 root root 4096 May 11 19:04 /sciclone/home
4 [bora] ls -ld /sciclone/home/ewalter
   drwxr-x---, 565 ewalter hpcf 94208 Jun 21 18:43 /sciclone/home/ewalter
5 [bora] ls -ld /sciclone/home/ewalter/file1
   -rw-rw-rw- 1 ewalter hpcf 0 Jun 21 18:43 /sciclone/home/ewalter/file1

"other" blocked from 'ewalter' folder
Main reasons to change permissions:

• **I want to give another user a file or folder(s).**
   If it is a single file (tar file for multiple files or folders) and it is small enough (< ~1GB) ...
   ... Just change the permissions on the file to be "world readable" and copy to /tmp (be sure it won't fill up /tmp)

   ```
   cp <file> /tmp
   chmod ugo+r /tmp/<file>
   ```

• **I want another user to be able to enter a directory of mine and read/copy any file.**
   If you are members of the same group (groups), you could make the whole path readable and folders executable by group:

   ```
   chgrp hpcstaff <folder>
   chmod g+rX -R <folder>
   ```

Don't forget to check the folders above for correct permissions
Research teams can request secondary groups to be created for them to share
Process Control

Sometimes, you will run a command and it takes too long / you want to kill it for some reason: `^c`

Sometimes (interactive job session) you want to run a calculation in the "background": `&`, `jobs`, `kill`

```
1 [bora] find . -name hello &
   [1] 14449

2 [bora] jobs
   [1]+ Running find . -name hello &

3 [bora] kill %1

4 [bora]
   [1]+ Terminated find . -name hello
```

`&` - puts the calculation in the background
`jobs` – lists current running "jobs"
`kill` – kills the job (%1)
`^c` – cancel
`^z` – background

Sometimes, you must use "kill –9" (always try without –9 first!)
You can also move a bg job to the fg and vice/versa:

1 [bora] find . -name hello
   ^z
[1]+   Stopped          find . -name hello

2 [bora] bg
[1]+ find . -name hello &

3 [bora] jobs
[1]+   Running          find . -name hello &

4 [bora] fg
find . -name hello
   ^c
"job" control with `%n only works for the shell that launched the process`. Any other shell need to use `ps`.

- `ps` – lists the current processes
  - `ps -ef` (all processes)
  - `ps -fu ewalter` (processes for ewalter)

### Can kill using process id:

<table>
<thead>
<tr>
<th>UID</th>
<th>PID</th>
<th>PPID</th>
<th>C</th>
<th>STIME</th>
<th>TTY</th>
<th>TIME</th>
<th>CMD</th>
</tr>
</thead>
<tbody>
<tr>
<td>ewalter</td>
<td>20926</td>
<td>35637</td>
<td>23</td>
<td>19:46</td>
<td>pts/0</td>
<td>00:00:00</td>
<td>find . -name hello</td>
</tr>
<tr>
<td>ewalter</td>
<td>21014</td>
<td>35637</td>
<td>0</td>
<td>19:46</td>
<td>pts/0</td>
<td>00:00:00</td>
<td>ps -fu ewalter</td>
</tr>
<tr>
<td>ewalter</td>
<td>35636</td>
<td>35633</td>
<td>0</td>
<td>19:10</td>
<td>?</td>
<td>00:00:00</td>
<td>sshd: ewalter@pts/0</td>
</tr>
<tr>
<td>ewalter</td>
<td>35637</td>
<td>35636</td>
<td>0</td>
<td>19:10</td>
<td>pts/0</td>
<td>00:00:00</td>
<td>-bash</td>
</tr>
</tbody>
</table>

1 [bora] `ps -fu ewalter`

2 [bora] `kill 20926`

- `top` – shows you the current running processes on a computer interactively
Shell Scripting

- Shell scripting is essential to utilize Linux environment efficiently
- tcsh/csh and bash/sh are two different shell flavors
- All HPC users default to tcsh
- # - comment character
- #! - NOT comment if on first line
- Linux is case sensitive

```bash
#!/bin/tcsh
# comment : #! Is a script shell interpreter ("do this with tcsh syntax")
foreach i (1 2 3 4 5)
    echo $i
end
```

```
[68 ewalter@bora ~ ]$ chmod u+x test.csh Change to executable
[69 ewalter@bora ~ ]$./test.csh . Is not in your PATH, must add it explicitly
1
2
3
4
5
```
Say you have a file of all the inputs you want to run:

1 [bora] cat joblist
1.45
1.44
1.40
1.33
1.10

Say you run the job like this:

1 [bora] ./a.out -i 1.45

2 [bora] cat runjobs
#!/bin/tcsh
foreach i (`cat joblist`) ` (backtick) means "the result of this command"
    echo $i
    ./a.out -i $i
end

3 [bora] chmod u+x runjobs
4 [bora] ./runjobs ./runjobs will run all parameters in joblist file
Shell Scripting – cont. 2

Main tcsh constructs:

**foreach loop**

```bash
#!/bin/tcsh
foreach i (`cat joblist`
    echo $i
    ./a.out -i $i
end
```

**If-then-else**

```bash
#!/bin/tcsh
set scen = b
if ($scen == a) then
    echo $scen
else
    echo "wrong value"
endif
```

**while loop**

```bash
#!/bin/tcsh
set i = 1
while ($i < 5)
    echo "i is $i"
    @ i++
end
```

**switch/case statement**

```bash
switch ($myvar)
    case 'foo':
        ./foo.csh
        breaksw
    case 'bar':
        ./bar.csh
        breaksw
    default:
        echo $usage
        breaksw
endsw
```

Even though you have a tcsh environment, you can still use bash shell scripts
Bash is considered more powerful for scripting / sometimes easier
PBS/Torque batch jobs

Serial Gaussian16 job:

#!/bin/tcsh
#PBS -N Gtest
#PBS -l nodes=1:vortex:ppn=1
#PBS -l walltime=0:60:00
#PBS -j oe

cd $PBS_O_WORKDIR
setenv GAUSS_SCRDIR /sciclone/scr10/ewalter/GAUSS
module load gaussian/g16

g16 test0032.com

Shared Memory Parallel Gaussian16 job:

#!/bin/tcsh
#PBS -N Gtest
#PBS -l nodes=1:vortex:ppn=12
#PBS -l walltime=0:60:00
#PBS -j oe

cd $PBS_O_WORKDIR
setenv GAUSS_SCRDIR /sciclone/scr10/ewalter/GAUSS
module load gaussian/g16

g16 -p=12 test0032.com

Once you have identified/prepared/installed software:

How can it run, serial only?
Shared memory parallel, distributed parallel?

Two vortex nodes
vx01
vx02

Serial
Shared memory parallel
Distributed parallel

Every code/language/framework is different. How parallelism is achieved can vary!
SLURM batch jobs

Serial Gaussian16 job:

```bash
#!/bin/tcsh
#SBATCH -J Gtest
#SBATCH -N 1 -n 1
#SBATCH -t 0:60:00

setenv GAUSS_SCRDIR /sciclone/scr10/ewalter/GAUSS
module load gaussian/g16
g16 test0032.com
```

Once you have identified/prepared/installed software:

How can it run, serial only?
Shared memory parallel, distributed parallel?

Shared Memory Parallel Gaussian16 job:

```bash
#!/bin/tcsh
#SBATCH -J Gtest
#SBATCH -N 1 -n 12
#SBATCH -t 0:60:00

setenv GAUSS_SCRDIR /sciclone/scr10/ewalter/GAUSS
module load gaussian/g16
g16 -p=12 test0032.com
```

Every code/language/framework is different. How parallelism is achieved can vary!
Distributed Parallel job:

```
#!/bin/tcsh
#PBS -N cpitest
#PBS -l nodes=3:vortex:ppn=12
#PBS -l walltime=1:00:00
#PBS -j oe

cd $PBS_O_WORKDIR
mvp2run -v ./a.out
```

How to run a job over **multiple processors** varies
Most parallel HPC codes use some flavor of **MPI**
The PBS/Torque clusters (only) have **mvp2run**

**mvp2run** will take your inputs and convert to the flavor of MPI you are using.

Can still run with mpirun, mpiexec, etc. But you may need to construct your own nodefile for some MPI (mvapich2).

[https://www.wm.edu/offices/it/services/researchcomputing/using/jobs/mvp2run/index.php](https://www.wm.edu/offices/it/services/researchcomputing/using/jobs/mvp2run/index.php)
Distributed Parallel job:

```
#!/bin/tcsh
#SBATCH -J cpitest
#SBATCH -N 3 -n 12
#SBATCH -t 1:00:00
```

How to run a job over **multiple processors** varies

Most parallel HPC codes use some flavor of **MPI**

ALL parallel executables should be run using `srun` in SLURM

```
srun ./a.out
```
Help with Common Software

- [https://www.wm.edu/offices/it/services/researchcomputing/using/software/](https://www.wm.edu/offices/it/services/researchcomputing/using/software/)
  basic info about using: *Matlab, Stata, Gaussian, and Python*

- Try an **interactive job** via batch system to work out job scripts, and workflow

- One thing to remember – your home directory is shared amongst all clusters / front-ends. Some software likes to install some files in your home directory. **Be careful to not overwrite software installed for one cluster by another.**

- One common example is *R* local packages. Likes to install local packages in home directory. If these packages are compiled on one cluster, may not work on another cluster.

- All Python on cluster should use Anaconda environments, not Python virtual environments. See software guide for more info.
Using the ticket system is useful since it is monitored by 5 of us

Thank you!
PBS/Torque jobs

Interactive batch system job

27 [vortex] qsub -I -l walltime=30:00 -l nodes=1:vortex:ppn=12
qsub: waiting for job 1552781 to start
qsub: job 1552781 ready
1 [vx01] python prog.py

Vortex has 2 nodes/24 cores reserved for walltimes of 1hr max 8am-6pm M-F
– can be used for interactive tests

Typical workflow is to test code with interactive job and run a batch job
Other clusters may be hard to schedule an interactive job.

pbstop – bora/james will show open slots
showstart – will show when job should start if it were next

https://www.wm.edu/offices/it/services/researchcomputing/using/jobs/index.php - HPC torque documentation
SLURM jobs
Interactive batch system job

```
1 [gust] salloc -N1 -n1 -t 0:30:00
salloc: Granted job allocation 388
salloc: Waiting for resource configuration
salloc: Nodes gt01 are ready for job

2 [gt01] python prog.py
```

- `shownodes` – will show available slots
- `squeue --start` – will show when job should start

https://www.wm.edu/offices/it/services/researchcomputing/using/jobsslurm/index.php - HPC SLURM documentation