## O2 for Orchestra Users

# **HMS** Research Computing

Fall 2017

### Welcome to O2!

- HMS Research Computing's second High-Performance Compute cluster to enhance the compute capacity available to HMS Researchers
- Homogeneous environment of newer, faster cores with high memory allocation to facilitate multi-core and parallelized workflows
- SLURM scheduler to efficiently dispatch jobs

## O2 Tech Specs

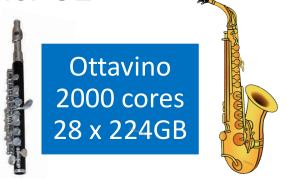


- 5000 homogeneous cores
- Dell Xeon E5-2683 processors: 2.1GHz Broadwell chips
- 16 x 2 (32) cores per node
- 256GB RAM per node (8GB/core)
- Login/load balancer 5 VM (8 cores/16GB memory)
- InfiniBand connectivity between nodes available
- CentOS 7



## Coming Soon to O2...

- Orchestra fastest compute nodes "ottavinos" reprovisioned for O2
- Orchestra High-Memory nodes "saxophone" reprovisioned for O2
- Orchestra GPU K80/M40 resources re-provisioned for O2



Saxophone 24 cores 12 X 1TB



GPU 24 cores



## Storage on O2

- Mounts all Orchestra storage
- Isilons: /home, /n/groups, /n/data1, /n/data2
- research.files: transfer partition, /n/files
- Lustre: /n/scratch2
- GPFS: /n/no\_backup









## Storage Mount Points

- /home, /n/data1, /n/data2 same
- /groups = /n/groups
- /files= /n/files on transfer partition (ask us)
- /n/no\_backup
- /n/scratch2

### slurm

- slurm is an open-source scheduler
- Rich, user-contributed community of questions and answers
- Simpler partition structure and management of fairshare (partition, user, group)
- Optimization for MPI and GPU
- Better resource allocation
- Advanced job accounting



## Login/transfer

- Login Nodes (5 Virtual Machines)
   \$ ssh eCommons@o2.hms.harvard.edu
- With X11:
  - \$ ssh -XY eCommons@o2.hms.harvard.edu
- Transfer Server (sFTP/port 22, 10 nodes):
   transfer.rc.hms.harvard.edu

### Class Practical

- Login to O2 via
  - \$ ssh eCommons@o2.hms.harvard.edu
- Copy the class files and scripts to your /home
  - \$ cp -r /n/groups/rc-training/o2

### .bashrc

- Orchestra and O2 both source the same ~/.bashrc
- Perl local::lib from Orchestra will not work on O2
- Module files are not synonymous between systems, and module loads will break
- Modify .bashrc as follows:

```
if [[ ! -z $SLURM_CONF ]]
then
    # put O2 specific commands here
else
    # put Orchestra specific commands here
fi
```



### Interactive Job:

- Interactive session: up to 20 cores, 12 hours
  - \$ srun -p interactive -t 0-12:00 --pty /bin/bash
- Can add cores (up to 20)n
- Can add total memory (default 1G, max 250G):
  - --mem 8G

### Jobs: bsub -> sbatch

- All in one line: --wrap="command here" #not recommended
- sbatch -p partition -t 0-1:00 --wrap="command"\$ sbatch -p short -t 0-1:00 --wrap="cp file.txt .."
- Complete shell script #recommended
- \$ sbatch completeSlurmJob.run

```
#!/bin/bash
#SBATCH -p short
#SBATCH -t 0-1:00
cp file.txt ..
```

# Partitions (queues): -q -> -p

Partition	Priority	Max Runtime	Max Cores	Limits
short	10	12 hours	20	
medium	8	5 days	20	
long	6	30 days	20	
interactive	14	12 hours	20	2 job limit
priority	14	30 days	20	2 job limit
mpi	12	5 days	640	20 core min
transfer		30 days	4	



### Walltime: -W -> -t

- -t days-hours:minutes
- -t hours:minutes:seconds
- Need to specify how long you estimate your job will run for
- Aim for 125% over
- Subject to maximum per partition
- Excessive runlimits (like partition max) take longer to dispatch, and affect fairshare

### $CPU: -n \rightarrow -n$

- -n X to designate CPU: max 20
- -N X to constrain all cores to X nodes
- CPU time: wall time (-t) \* (-n) CPUs used
- Unable to use CPU not requested (no overefficient jobs): cgroups constraint

## Memory: -Rusage -> --mem

- Only 1G is allocated by default
- --mem XG #total memory over all cores
- --mem-per-cpu XG #total memory per CPU requested, use for MPI
- No unit request (G) defaults to Megabytes
  - 8G ~= 8000

### **Job Construction**

```
#!/bin/bash
#SBATCH -p #partition
#SBATCH -t 0-01:00 #time days-hr:min
#SBATCH -n X #number of cores
#SBATCH -N 1 #confine cores to 1 node, default
#SBATCH --mem=XG #memory per job (all cores), in GB
#SBATCH -o %j.out #out file
#SBATCH -e %j.err #error file
#SBATCH --mail-type=BEGIN/END/FAIL/ALL
#SBATCH -mail-user=mfk8@med.harvard.edu
```



# Output/Error Files

- Can add jobid to filename with %j
- Sample:
  - -e %j.err
  - -o %j.out
- SLURM by default creates this outfile: slurm-jobid.out
- Additional Flags
- %a #job array id
- %A #master array job id
- %N #node name
- %u #user id

### Mail

- Mail is not auto-generated upon completion/failure
- #SBATCH --mail-type= NONE, BEGIN, END, FAIL, REQUEUE, ALL
- #SBATCH \_--mail-user=mfk8@med.harvard.edu
- Not recommended, not a verbose output like LSF
- Use sacct instead

## Practical: simple sbatch script

From your ~/o2 directory,

\$ sbatch submit.slurm

```
#!/bin/bash
```

#SBATCH -p short

#SBATCH -t 0-00:01

#SBATCH -n 1

#SBATCH -N 1

#SBATCH --mem=2G

#SBATCH -o hostname.%j.out

#SBATCH -e hostname.%j.err

srun hostname #command

# Partition to submit to

# Time in minutes

# Number of cores requested

# Ensure that all cores are on one machine

# Memory total in GB

# Standard out goes to this file

# Standard err goes to this file



## Practical: checking result

- From ~/o2 directory:
  - \$ less hostname.\*.out
- This reads the file "hostname.%j.err"
- The host the job ran on is reported as:

compute-a-16-X.o2.rc.hms.harvard.edu

# Job Master Allocations/Job Steps

- "sbatch" or "salloc" will create a master allocation of shared resources
- To run job steps or multiple commands in parallel with resources from the master allocation, use srun

```
#/bin/bash
#SBATCH -n 4
#SBATCH --mem 32000
srun -n 1 --mem=8000 COMMAND1 &
srun -n 1 --mem=8000 COMMAND2 &
srun -n 1 --mem=4000 COMMAND3 &
srun -n 1 --mem 12000 COMMAND4 &
wait #necessary
```

# starts 4 independent, single-threaded parallel tasks with memory constraints

```
#/bin/bash
#SBATCH -c 4
#SBATCH --mem 32000
```

```
srun -c 2 COMMAND1 & srun -c 2 COMMAND2
```

wait #necessary

# starts 2 independent, multicores tasks





### Practical: Job Parallelization

From ~/o2:

```
$ sbatch date_parallel.sh
```

#### Contents:

```
#SBATCH -n 3
srun -n 1 date &
srun -n 1 sleep 2m &
srun -n 1 -c 2 date &
srun -n 1 -c 3 date &
wait
```

- Output file can be read as
  - \$ less parallel.\*.out



## Job Arrays

- sbatch --array=1-30 submit.sh
- #SBATCH --array=1-30
- slurm creates:
- \$SLURM\_JOB\_ID #jobid of each job in array %j
- \$SLURM\_ARRAY\_JOB\_ID #jobid of entire array %A
- \${SLURM\_ARRAY\_TASK\_ID} #index of job %a

#### Example:

Files named: File1.txt File2.txt File3.txt ...File30.txt

Execution is:

myscript.sh File\${SLURM\_ARRAY\_TASKID}.txt



## Practical: Job Array

From ~/o2

```
$ sbatch --array=1-4 fastqc_job_array.sh
```

Relevant file pieces:

```
#SBATCH -o fastqc_%A_%a.out
module load fastqc/0.11.5
fastqc sample_"{SLURM_ARRAY_TASK_ID}" _R1.fastq
```

- Creates fastqc report for reach fastq file
- Creates output progress file named fastqc\_ArrayId\_ArrayIndex.out

## Job Dependencies

- sbatch --dependency=
- after:jobid #(asynchronous)
- afterany:jobid #after exit or done
- afterok:jobid #success, exit code 0
- afternotok:jobid #failure
- singleton #after jobs with same name have terminated
- --kill-on-invalid-dep=<yes|no> #kill on unmet dependency (on by default)

## Command Line Arguments

- slurm scripts can take command line arguments after them, unlike bsub
- Reference as \$1, \$2 etc
- sbatch submit.run 25 output.txt

```
#!/bin/bash
#SBATCH -p short
#SBATCH -t 0-1:00
python myscript.py $1 $2
```

### X11 on O2

- To visualize or initiate plot devices, an X11 device must be active
- Mac: Xquartz installed and running
- Windows: Xming installed and running
- Login: ssh –XY
- SSH keys:

Host login.rc.hms.harvard.edu

ForwardX11 yes

ForwardX11Trusted yes

- To sbatch jobs add: --x11=batch
- To interactives, srun add: --x11





## Fairshare -> Priority

- Dynamically assigned
- Factors contributing:
- Age, Fairshare, Partition, QOS, Nice
- Fairshare: 0-1 scale

## Job Dispatch Order in O2

#### Sum of

- AGE = how long a job has been pending
- FAIRSHARE = fairshare of the user
- PARTITION = priority of the partition used
- QOS = quality of services, normally not used
- NICE = additional custom priority, normally not used

#### Notes:

- Jobs submitted to the partitions interactive and priority are evaluated before jobs in any other partitions, independently of the priority values.
- A lower priority job can be dispatched before an higher priority job if it does not impact the expect start time of the higher priority job (job backfilling)

## Job Monitoring

- \$ squeue -u eCommons -t RUNNING/ PENDING
- \$ squeue -u eCommons -p partition
- \$ squeue -u eCommons --start
- Detailed job info:
   \$ scontrol show jobid <jobid>
- Completed job statistics:

```
$ sacct -j <jobid> --
format=JobID,JobName,MaxRSS,Elapsed
```

## Job information: sacct

- Advanced job accounting options
- \$ saact -j jobid

#### Options:

- --name
- -r/--partition
- -s/--state
- -o/--format
- \$ sacct -u eCommons
- \$ sacct --helpformat #get available accounting features

### sacct: basic information

- sacct --helpformat displays fields
- RC Recommends:

```
sacct -u eCommons --
format=jobid,Partition,state%22,MaxRSS,MaxVMSize,ReqTRES%20,Star
t%20,End%20,Timelimit%14,exitcode --units=G
```

 Gives jobid, partition, end state, Max Memory, Max Virtual Memory, Requested Cores/Mem, Start, Stop, Timelimit, Exit Code in GB format

### saact: state

- BF BOOT\_FAIL
- CA CANCELLED
- CD COMPLETED
- CF CONFIGURING
- CG COMPLETING
- DL DEADLINE
- F FAILED
- NF NODE\_FAIL

- PD PENDING
- PR PREEMPTED
- R RUNNING
- RS RESIZING
- S SUSPENDED
- TO TIMEOUT



## Cancelling/Pausing Jobs

- \$ scancel <jobid>
- \$ scancel -t PENDING
- \$ scancel --name JOBNAME
- \$ scancel jobid\_[indices] #array indices
- \$ scontrol hold <jobid> #pause
- \$ scontrol release <jobid> #resume
- \$ scontrol requeue <jobid> #cancel and rerun

### MPI on O2

- Message Passaging Interface
- Distribute work over multiple nodes, allowing for the utilization of more cores
- openMPI-2.0.1 compiled against GCC 6.2.0
- MATLAB, Python, R, Perl, Java, C++, Fortran implementations
- Needs wrapper function "mpirun" to dispatch to compute nodes with SLURM
- Run in "mpi" partition –p mpi after being added to partition
- Core cap: 640 processors, 5 day runtime

### LMOD: Software Modules

- LMOD system adds directory paths of software into \$PATH variable, and resolves software dependencies and conflicts
- Most software compiled against gcc-6.2.0: load first
- \$ module load gcc/6.2.0 #note, no directories anymore
- \$ module spider #software currently available
- \$ module load software/version #load software, no directories
- \$ module unload software/version #unload
- \$ module purge #dump all modules
- \$ module help <software> #displays run info



## Programming Languages

- Python: load module (2.7.12, conda2, 3.6.0)
  - use virtualenv to maintain packages (pip/easy install)
- R: load module (3.2.5, 3.3.3, 3.4.1)
  - Setup O2-specific personal R library, .Renviron (install.packages/biocLite)
- Perl: load module (5.24.0)
  - Setup O2-specific local::lib (cpan/cpanm) in .bashrc
- MATLAB: load module (2016b, 2017a)
  - Setup cluster profile, only use on 1 HPC at a time



### For more direction

- http://hmsrc.me/O2docs
- http://rc.hms.harvard.edu
- RC Office Hours: Wed 1-3p Gordon Hall 500
- rchelp@hms.harvard.edu