



High Performance Computing

# Introduction to Wulver: Job Scheduler & Submitting Jobs

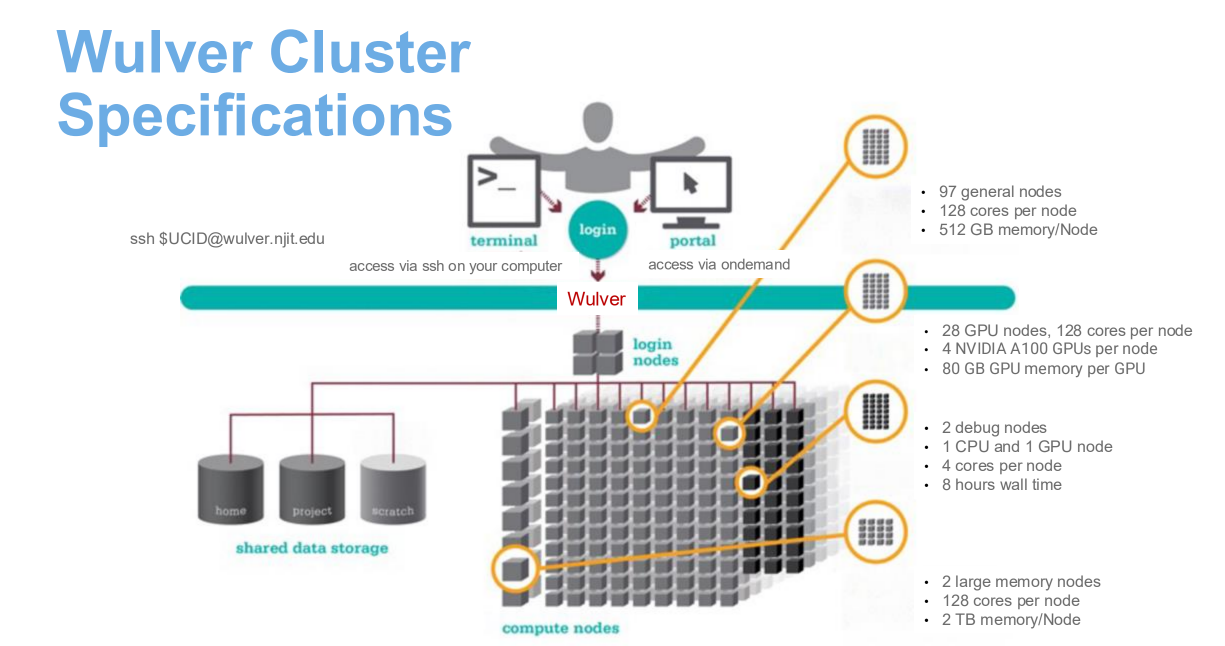
Oct 1, 2025



#### **Outline**

- Wulver Specifications
- Access the software on Wulver
- Batch Processing
- Example of Slurm Jobs
- Manage Slurm Jobs
- Troubleshooting Common Issues
- Slurm Interactive Jobs and Use GUI Apps
- Contact Us





#### **Environment Modules**

- Environment Modules allows for dynamic modification and management of a user's environment via modulefiles.
- Manages multiple versions of software that require unique environments.
- Allows the user to load only the environment variables important to their applications, from within their job.

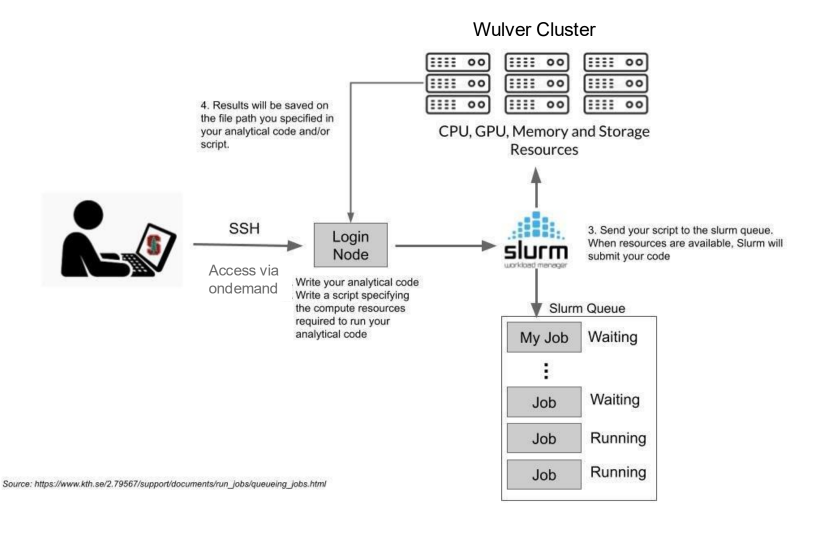
T	What modules do you have loaded?	module list
<b>*</b>	What modules are available?	module spider
	Multiple versions of the same compiler	module avail intel
	Add a software module to your environment	module load CUDA
~	Remove a software package from your environment	module unload intel



#### **Batch Processing**



# Why do supercomputers use queuing?



#### What is Slurm?

- Slurm is the predominant Open-Source scheduler for HPC compute
- Historically Slurm was an acronym standing for
  - Simple Linux Utility for Resource Management
- The Slurm scheduler provides three key functions:
  - it allocates access to resources (compute nodes) to users for some duration of time so they can perform work.
  - it provides a framework for starting, executing, and monitoring work (typically a parallel job such as MPI) on a set of allocated nodes.
  - it arbitrates contention for resources by managing a queue of pending jobs.



#### Manage Jobs – Options

#### **Mandatory Options**

Directive	Options	Description
account=	account	PI's UCID
partition=	<u>Partition</u>	Request a partition of resources for job allocation (queue)
time=	Time [[d-]h:]m[:s]	Maximum time limit on job allocation
qos=	<u>Job</u> <u>Priorities</u>	Define the job priority

#### **Partitions**

Partition	Nodes		CPU	GPU	Memory
partition=general	97	128	2.5G GHz AMD EPYC 7763 (2)	NA	512 GB
partition=debug	1	4	2.5G GHz AMD EPYC 7763 (2)	NA	16 GB
partition=debug_gpu	1	4	2.0 GHz AMD EPYC 7713 (2)	<ul><li>Eight 10G MIG</li><li>Four 20G MIG</li><li>Four 40G MIG</li></ul>	16 GB
partition=gpu	28	128	2.0 GHz AMD EPYC 7713 (2)	<ul><li>NVIDIA A100 GPUs (4)</li><li>MIG</li></ul>	512 GB
partition=bigmem	2	128	2.5G GHz AMD EPYC 7763 (2)	NA	2 TB

# Job Submission Time Interval Formats

Valid time formats (with a few exceptions) for -t / - time= option

Minutes	(-t 10 is 10 minutes)
Minutes:Seconds	(10:30 is 10 minutes & 30 secs)
Hours:Minutes:Seconds	(1:0:0 is 1 hr + 0mins + 0secs)
Days-Hours:Minutes:Seconds	(7-1:10:30 is 7days + 1hr + 10mins + 30secs)
Days-Hours	(7-0 is 7days + 0hrs i.e. 7 days)
Days-Hours:Minutes	(7-4:10 is 7days + 4hrs + 10mins)

#### QoS

- Standard Priority (--qos=standard)
  - Faculty PIs are allocated 300,000 Service Units (SU) per year on request at no cost
  - Additional SUs may be purchased at a cost of \$0.005/SU.
  - The minimum purchase is 50,000 SU (\$250)
  - Wall time maximum 72 hours
  - SUs will reset every year in mid-January with no carryover.
- Low Priority (--qos=low)
  - Not charged against SU allocation
  - Wall time maximum 72 hours
  - Jobs can be preempted by those with higher and standard priority jobs when they are in the queue
- Debug Priority (--qos=debug)
  - Not charged against SU allocation
  - Wall time maximum 8 hours
  - Must be used with --partition=debug or --partition=debug gpu
  - · Only one job per user is allowed at a time
- High Priority (--qos=high \$PI UCID)
  - Not charged against SU allocation
  - Wall time maximum 72 hours can be increased based on PI's request
  - Only available to contributors
  - Use listgos command

#### MIG

**Multi-Instance GPU (MIG)**: A feature that partitions a single NVIDIA A100 80GB GPU into multiple isolated GPU instances

Each instance has dedicated GPU memory, compute cores, cache, and bandwidth

#### Why Use MIG?

- Reduces queue times: smaller GPU slices fit into the scheduler more easily
- Improve overall cluster utilization.

#### MIG Implementation in Wulver

- 10gb
- 20gb
- 40gb



# Jobs - Options

#### **Additional Options**

Directives	Options	Description	
ntasks=	Number of cpus	Number of CPUs (tasks) to be allocated	
nodes= Node		Number of Nodes	
ntasks-per- Numbers of node= cpus per node		Number of CPUs (tasks) per each node to be allocated	
mem=	Memory	Total memory of the job	
mem-per- cpu=	Memory per cpu	Memory to be allocated per each cpu	
gres=	Generic resources	Set the Number of gpus	
cpus-per- task=	Cpus per task	Number of CPUs per task	
requeue	Requeue	This is required when you want to continue the job after 72h walltime.	

Directives	Options	Description
error=	File	Define standard error file
out=	File	Define standard output file
input=	File	File used for standard input
job	Name	Define job name
mail-type=	ALL, BEGIN, END, FAIL, REQUEUE	Notify user by email when <type> event occurs</type>
mail-user=	Email address	Send email to this address for events specified with mail-type option (default is submitting user).
dependency=	Job dependency	Set the job dependency when submitting multiple jobs

# Service Units (SU)

- Service Units (SUs) are the core accounting mechanism used to track and allocate compute usage on cluster.
- SUs are charged only in --qos=standard

Partition	SU Charges
partition=general	MAX(CPUs, RAM/4G) SU
partition=gpu	MAX(CPUs, RAM/4G) SU + 16 * (GPU Memory requested)/80G
partition=bigmem	MAX(1.5 * CPUs, 1.5 * RAM/16G) SU
partition=debug	No charges
partition=debug_gpu	No charges

## **Example of SU Charges**

4 CPUs + --mem=128G: MAX(4\*1.5, 1.5\*128G/16G) = 12

```
general
4 \text{ CPUs: MAX}(4, 4*4G/4G) = 4
4 \text{ CPUs} + --\text{mem} = 64\text{G} : \text{MAX}(4, 64\text{G}/4\text{G}) = 16
gpu
4 \text{ CPUs} + 40 \text{MIG} : \text{MAX}(4, 4*4 \text{G}/4 \text{G}) + 16* (40 \text{G}/80 \text{G}) = 12
4 CPUs + Full GPU: MAX(4, 4*4G/4G) + 16 * (80G/80G) = 20
4 CPUs + --mem=64G + Full GPU: MAX(4, 64G/4G) + 16 * (80G/80G) = 32
bigmem
4 CPUs: MAX(4*1.5, 1.5*4*16G/16G) = 6
```

#### **General Application Workflow**

- Log into cluster with ucid and password
- Copy input files to new directory
- Change to copied directory via command line cd /path/to/copied\_directory
- Copy job a template to the directory cp /path/to/templates/jobtemplate.job jobfile.job
- Modify the job file:
  - Change the number of resources to desired number
  - Change the module load command based on the application name and version
  - Update command line with commands required for job
  - Update the software modules
- Submit the job file using sbatch



#### **Examples of Slurm Jobs**

# Sample Simple Job Script

```
#!/bin/bash
#SBATCH --job-name=my_job
#SBATCH --partition=general
#SBATCH --output=%x.%j.out
                                                        Job setup information
         --error=%x.%j.err
                                                        for SLURM
         --account=PI UCID
         --gos=low
         --time=00:20:00
 SBATCH --nodes=1
 SBATCH --ntasks-per-node=1
#SBATCH --mail-type=ALL
#SBATCH -mail-user=ab1234@njit.edu
date
                                                       Commands to be run
sleep 60
date
```

- This runs a batch job called "my\_job" to the "general" partition, with 1 task, a wall time limit of 20 minutes.
- QOS is required. Account is recommended.
- %x.% j expands to JobName.JobID and prints this into a text file

## Sample MPI Job script

```
#!/bin/bash

#SBATCH --job-name=mpi_test_job
#SBATCH --partition=general
#SBATCH --output=%x.%j.out
#SBATCH --error=%x.%j.err
#SBATCH --account=PI_UCID
#SBATCH --qos=low
#SBATCH --time=00:10:00
#SBATCH --ntasks=256
#SBATCH --ntasks-per-node=128
#SBATCH --mem-per-cpu=2G

# Run application commands
srun /apps/testjobs/bin/mpihello
```

• This runs an MPI job named "mpi\_test\_job", with 256 processes total, spread over 2 nodes. Default setting is 1 core per process/task, so this also allocates 512Gb memory total. Wall time is 10 minutes.

# Sample Multi GPU Job script

```
#!/bin/bash

#SBATCH --job-name=test_gpu_job
#SBATCH --output=%x.%j.out
#SBATCH --error=%x.%j.err
#SBATCH --partition=gpu
#SBATCH --account=PI_UCID
#SBATCH --qos=low
#SBATCH --time=00:20:00
#SBATCH --time=01:20:00
#SBATCH --opus-per-task=32
#SBATCH --cpus-per-task=32
#SBATCH --gres=gpu:2

# Load application environment
module load CUDA

# Run application commands
nyidia-smi
```

 This runs a GPU job named "test\_gpu\_job", with 64 cpus and full access to 2 GPUs. Wall time is 20 minutes.

## Sample Multi GPU Job script

```
#!/bin/bash
#SBATCH --job-name=test_gpu_job
#SBATCH --output=%x.%j.out
#SBATCH --error=%x.%j.err
#SBATCH --partition=gpu
#SBATCH --account= PI UCID
#SBATCH --qos=low
#SBATCH --time=00:20:00
\#SBATCH --ntasks=1
#SBATCH --cpus-per-task=2
#SBATCH --qpus-per-task=2
# Load application environment
module load CUDA
# Run application commands
nvidia-smi
```

This runs a GPU job named "test\_gpu\_job", with 2 cpus and 2 GPUs

# Sample GPU Job script (MIG)

```
#!/bin/bash
#SBATCH --job-name=test mig job
#SBATCH --output=%x.%j.out
#SBATCH --error=%x.%j.err
#SBATCH --partition=qpu
#SBATCH --account= PI UCID
#SBATCH --qos=low
#SBATCH --time=00:20:00
#SBATCH --ntasks=2
#SBATCH --gres=gpu:a100 20g:1
# Load application environment
module load CUDA
# Run application commands
nvidia-smi
```

This runs a GPU job named "test\_mig\_job", with 2 CPUs and one 20G MIG



#### Manage Slurm Jobs

#### Manage Jobs – Submit via CLI

Submit a job script

- sbatch my script
- Submitted batch job 1234

#### Listing jobs

For current user in Pending, Running, Suspended states:
squeue -u \$LOGNAME

JOBID PARTITION NAME USER ST TIME NODES NODELIST (REASON)
1234 general uname.sh test PD 0:00 2 (Priority)

For a more detailed query on active job:

•scontrol show jobid=1234

JobId=2 JobName=simple.job
UserId=test(1001) GroupId=test(1001) MCS\_label=N/A
Priority=4294901759 Nice=0 Account=(null) QOS=standard
JobState=COMPLETED Reason=None Dependency=(null)

Canceling jobs

- scancel 1234 Cancel job ID 1234
- scancel --me Cancel all your jobs

Show information about an active or completed job

slurm\_jobid 1234

#### **Job States**

- CA CANCELLED Job was explicitly cancelled by the user or system administrator. The job may or may not have been initiated.
- CD COMPLETED Job has terminated all processes on all nodes with an exit code of zero.
- CF CONFIGURING Job has been allocated resources but are waiting for them to become ready for use (e.g. booting).
- CG COMPLETING Job is in the process of completing. Some processes on some nodes may still be active.
- F FAILED Job terminated with non-zero exit code or other failure condition.
- NF NODE\_FAIL Job terminated due to failure of one or more allocated nodes.
- PD PENDING Job is awaiting resource allocation.
- R RUNNING Job currently has an allocation.
- RQ REQUEUED Completing job is being requeued.
- **PR PREEMPTED** The job was terminated because of preemption by high priority job.
- ST STOPPED Job has an allocation, but execution has been stopped with SIGSTOP signal. CPUS have been retained by this job.
- S SUSPENDED Job has an allocation, but execution has been suspended and CPUs have been released for other jobs.
- TO TIMEOUT Job terminated upon reaching its time limit.

#### Waiting for Your Job To Run

- Queue wait time depends on many factors
  - System load
  - Resources requested
    - nodes, cores, large memory, gpus
    - reduced priority for users or groups using a lot of resources
- Check the running jobs in QoS
  - squeue -q [QoS]



# **Troubleshooting Common Issues**

# **Common inquiries**

#### checkload

• sinfo but more details

#### checkq

• squeue but more details

#### slurm\_jobid

Show information about a running or queued job

#### sq

 $\bullet$  Display pending job/queue info in a helpful way, You can also check the last job details with  ${\tt sq}$ 

#### squeue --start

- Jobs will be listed in order expected start time
- Times are only estimates and subject to change

#### quota\_info

Show storage and SU quotas for self or others

#### listqos

Show all QOSes or members of QOSes

#### Some Common Problems

After using sbatch, the job disappears in 30 seconds and there's no result output.

- Check the details with slurm\_jobid [JOBID], use --err and --out
- Use sq if you are unsure about the job id.

Invalid account or account/partition combination specified.

- Check --account
- Use quota info \$LOGNAME

```
JOBID PARTITION NAME USER ST TIME NODES NODELIST(REASON)

1234 general uname.sh xiss PD 0:00 2 (ReqNodeNotAvail, Reserved for maintenance)
```

 Jobs that do not end before the maintenance window begins will be held until the maintenance is complete

```
JOBID PARTITION NAME USER ST TIME NODES NODELIST(REASON)

1234 general uname.sh xiss PD 0:00 2 (MaxCpuPerAccount)
```

- listqos high \$PI
- squeue -q high\_\$PI

# Some Common Problems(contd)

```
JOBID PARTITION NAME USER ST TIME NODES
NODELIST(REASON)
1234 general uname.sh xiss PD 0:00 2
(AssocGrpBillingMinutes)
```

- Your PI group have reached the limit of SU in standard
- scontrol update JobId=Job\_ID QOS=low

#### Error message: cggroup out of memory handler

- Check the memory requirement
- You probably need to increase memory on --mem or the number of cpus from --mem-per-cpu
- If nothing works, then the problem is likely due to incorrect setup of the problem.

```
JOBID PARTITIONNAMEUSER ST TIMENODES NODELIST(REASON)1234general uname.shxiss PD 0:002 (MaxMemPerAccount)
```

- listqos high \$PI
- squeue -q high \$PI



# Slurm Interactive Jobs and Use GUI Apps

#### **Interactive Batch Jobs**



Interactive, but handled through batch system

Resource limits same as standard batch limits



Useful for tasks forbidden on login nodes

Debug parallel programs

Quickly test your code



May not be practical when system load is high

Long wait, same as standard batch job



To submit an interactive batch job (example)

- interactive -a ACCOUNT -q QOS -j JOB\_TYPE
- interactive -h

#### Using Applications with GUI on OnDemand

Login to ondemand.njit.edu

Go to "Interactive Apps" and select the application from the list.

If you don't find the app, select Linux Desktop

Once are you connected, select "Terminal Emulator" from "Applications" option from top left

#### Reminder



- Wulver will be temporarily out of service for maintenance once a month, specifically on the 2nd Tuesday, to perform updates, repairs, and upgrades.
- During the maintenance period, the logins will be disabled
- Jobs that do not end before the maintenance window begins will be held until the maintenance is complete
- Reduce the walltime in the job script to run the job



- Date: Every Tuesday and Friday
- Time: 2:00-4:00 p.m.
- Location: GITC 2404
- Meet with our student consultants and ask any questions you have about using HPC resources.
- There's no need to create a ticket in advance.

# Resources to get your questions answered

Getting Started: Access to Wulver

List of Software: Wulver Software

**HOW TOs:** Conda Documentation

Installing Python packages via Conda

Running Jobs: Jobs

Access to OnDemand Open OnDemand

MIG Information MIG

Contact: Please visit HPC Contact

Open a ticket: email to <a href="mailto:hpc@njit.edu">hpc@njit.edu</a>

Consult with Research Computing Facilitator: Facilitator Calendar Appointment

System updates

- · Read Message of the Day on login
- Visit NJIT HPC News



