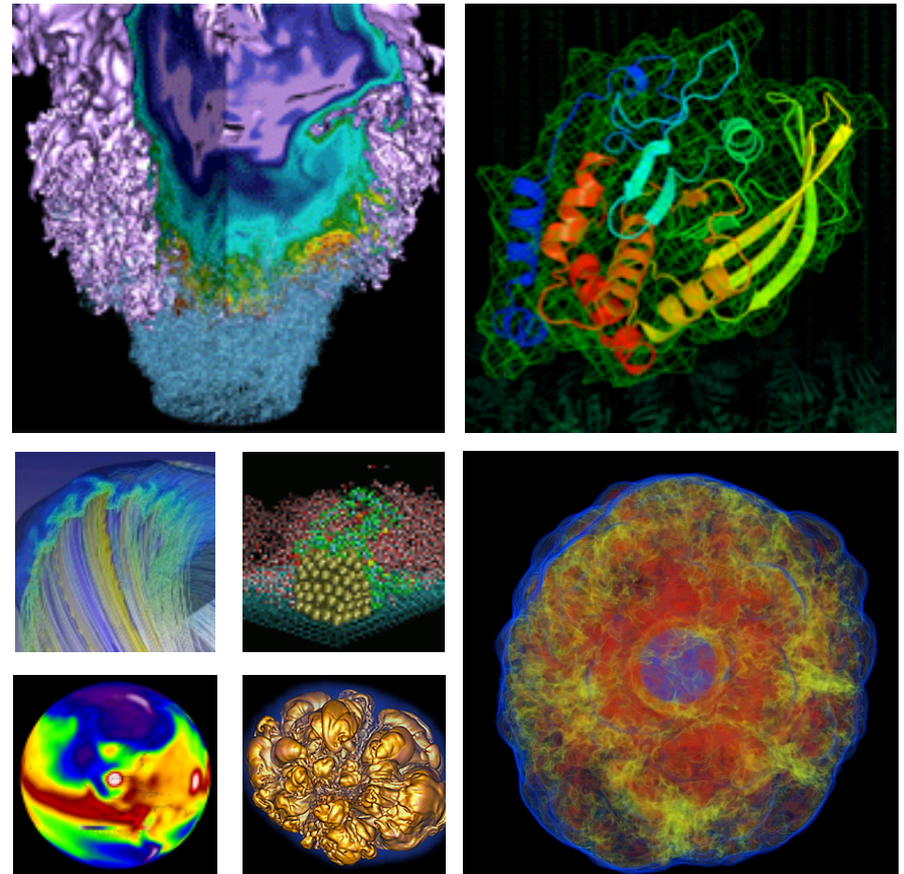


# Computing Environment



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# Node Types

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- **Login nodes**
  - Shared
  - Job preparation and submission
- **MOM nodes**
  - Shared
  - Where batch script executes
    - Parallel job launcher
      - Hopper/Edison: aprun
      - Carver: mpirun
- **Compute nodes**
  - Not shared
    - Except serial queue

# Login Nodes

- **Edison**
  - Six nodes
    - 16 cores, 2.0GHz Intel Sandy Bridge
    - 512GB
- **Hopper**
  - Eight nodes
    - 16 cores, 2.4GHz AMD Opteron
  - Four nodes
    - 32 cores, 2.0GHz AMD Opteron
  - 128GB
- **Carver**
  - Four nodes
    - 8 cores, 2.66GHz Intel Nehalem
    - 48GB

# Login Node Access



- **Connect (via ssh) to *load balancer***
  - edison.nersc.gov
  - hopper.nersc.gov
  - carver.nersc.gov
- **Load balancer selects login node based on:**
  - Number of connections
  - Memory of previous connections from same IP
    - If you login everyday, you'll probably end up on the same login node every time.

# Login Node Usage



- **Login nodes are shared by many users, all the time**
- **Edit files, compile programs, submit batch jobs**
- ***Some* post-processing/data analysis**
  - IDL
  - MATLAB
  - NCL
  - python
- ***Some* file transfers**
  - Use data transfer nodes for large/long-running transfers
- **Please use discretion**
  - *All* users get frustrated by sluggish interactive response

# Login Node Monitoring



- **Determine number of available cores**  
% grep processor /proc/cpuinfo | wc -l
- **Determine amount of physical memory**  
% grep MemTotal /proc/meminfo
- **Use “top” command to view process activity**

# Login Node Guidelines



- **Limit use of parallel “make”**
  - % make -j 4 all
- **Use *no more* than 50% of available cores**
- **Use *no more* than 25% of available memory**
- **NERSC will kill user processes if response becomes unacceptable**
- **Terminate idle sessions of licensed software**
  - IDL
  - MATLAB
  - Mathematica

# Shell Initialization Files

- **Standard dot files**
  - .bashrc, .profile, .cshrc, .login, etc.
  - Symbolic links to read-only files
    - Allows NERSC to provide common environment
- **Personal dot files**
  - Aliases, environment variables, modules, etc.
  - Use “.ext” files
  - .bashrc.ext, .profile.ext, .cshrc.ext, .login.ext, etc.
- **Use “fixdots” to start over**
  - Creates \$HOME/KeepDots.<timestamp>
  - Restores all dot files to current default state
- **Use NIM to change default login shell**

# NERSC Supported Software



- **NERSC provides a wide range of software**
  - Scientific Applications
    - VASP, Amber, NAMD, ABySS, ...
  - Compilers
    - PGI, Intel, GCC, Cray
  - Scripting Languages
    - perl, python, R
      - and packages for each!
  - Software Libraries
    - blas/lapack (MKL), boost, hdf5, netcdf, ...
  - Utilities
    - gnuplot, git, mercurial, cmake, ...
  - Debuggers and Profilers
    - CrayPat, DDT, TotalView, gdb, MAP, darshan
  - Visualization
    - Visit, ParaView, VMD, ...
- **See complete list**  
<http://www.nersc.gov/users/software/>

# Software is Managed by Modules



- **NERSC provides many versions of many software packages**
  - To support diverse workload on systems
- **Maintaining all these separate software installations on heterogeneous systems is a major challenge!**
  - Software can't just be installed in the base operating system
    - How many copies of `/usr/bin/vasp` could be supported?
  - Each software package installed in its own directory  
`/usr/common/usg/blast+/2.2.26`

**Modules is the user interface  
to software at NERSC**

# How to Access NERSC Software



- **Identify the software you need**

- Use the NERSC website

- <http://www.nersc.gov/users/software/>

- Use “module avail”

- *Lots* of output
- Each system has different modules!

- **Load the module**

```
% idl
```

```
idl: Command not found.
```

```
% module load idl
```

```
% which idl
```

```
/usr/common/usg/idl/idl82/bin/idl
```

# Loading Modules

- **Separate modules exist for each version of software**
    - Syntax: <name>/<version>
    - Default provided if no <version> supplied
- ```
% module avail idl  
idl/7.1    idl/8.0    idl/8.2 (default)  
% module load idl/7.1
```
- **Load modules in every batch script**
    - Ensure correct run-time environment
    - Self-documenting for troubleshooting and reproducibility

# Other Useful Module Commands

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## **module unload <modulename>**

- Remove the module from your environment

## **module swap <module1> <module2>**

- Unload one module and replace it with another

```
module swap pgi gcc
```

## **module list**

- See what modules you have loaded right now

## **module show <modulename>**

- See what the module actually does

## **module help <modulename>**

- Get more information about the software

# Default Modules



- **When you login, many *default* modules are loaded automatically**
  - Usually foundational modules which are required to get proper function from the system
    - Build environment, required libraries and applications, batch environment
  - Use caution in unloading these
- **Swapping to functionally equivalent module may be OK**
  - `carver% module swap pgi gcc`
  - `hopper% module swap PrgEnv-pgi PrgEnv-gnu`
- **Each NERSC system has different default modules**

# Types of Modules

- **Applications**
  - VASP, amber, blast, ...
  - Usually only set PATH, LD\_LIBRARY\_PATH
- **Libraries**
  - Set LD\_LIBRARY\_PATH
    - but probably not on Crays
  - Set “helper” environment variable for building software
    - Header/include file search paths
    - Library search paths
    - Library names

```

% module load hdf5
% mpicc mycode.f $HDF5
          
```

# Cray Programming Environment



- **Compiler specific**

PrgEnv-pgi, PrgEnv-intel, PrgEnv-cray, PrgEnv-gnu

– Intel is default on Edison, PGI is default on Hopper

- ***Meta-modules***

– Organize a set of modules

- Compiler (intel, pgi, cray, gnu)

- Libraries tuned for compiler

- **Swapping Programming Environments**

`module swap PrgEnv-pgi PrgEnv-intel`

– swaps compiler

– ***no need to swap libraries!***

# Carver “Programming Environment”

- Not as sophisticated as Cray PrgEnv
- Separate compiler and OpenMPI modules

| Compiler module | OpenMPI module |
|-----------------|----------------|
| pgi             | openmpi        |
| intel           | openmpi-intel  |
| gcc             | openmpi-gcc    |

- ***Must keep libraries consistent with compiler!***

# Compiler Wrappers



- **Edison/Hopper**
  - Defined by PrgEnv modules
  - **ftn, cc, CC**
  - Provides include and library search paths for MPI and some common math libraries
  - Provides consistent level of optimization across compilers
- **Carver**
  - Defined by openmpi modules
  - **mpif90, mpiCC, mpiCC**
  - Provides include and library search paths for OpenMPI
- **Seldom need native compilers!**

# Resources

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<http://www.nersc.gov/users/software/nersc-user-environment/>

<http://www.nersc.gov/users/software/nersc-user-environment/modules/>

<http://www.nersc.gov/users/computational-systems/edison/programming>

<http://www.nersc.gov/users/computational-systems/hopper/programming/>

<http://www.nersc.gov/users/computational-systems/carver/programming/>



**Thank you.**