

Programming Environments & Compilation on Perlmutter



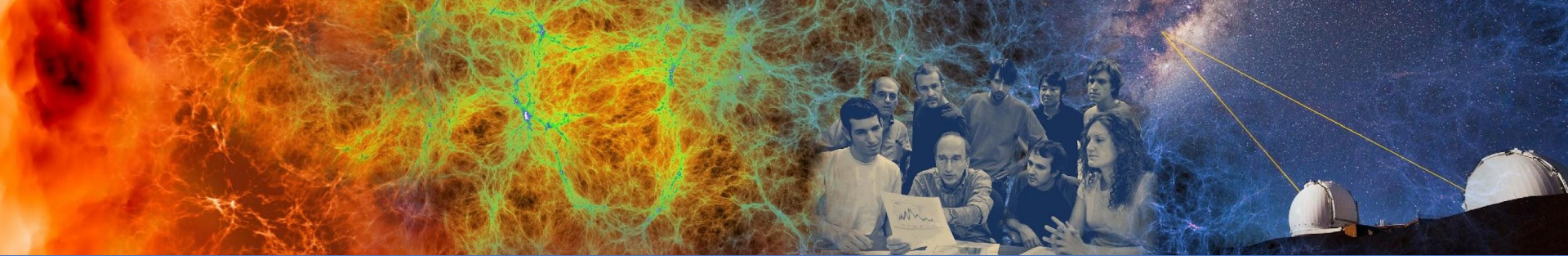
New User Training
September 7, 2023

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User Engagement Group

Outline: Four Main Ways of Getting Software

- **Loading Modules**
- Containers
- **Compiling from source**
- **Spack & E4S**

```
#####  
perlmutter  
#####  
Welcome to perlmutter!  
#####  
For all planned outages, see: https://www.nersc.gov/live  
For past outages, see: https://my.nersc.gov/outagelog-cs  
epalmer@perlmutter:login30:~> $  
epalmer@perlmutter:login30:~> $ Now what?  ^\_(\u0329)\_/^
```



Modules: Loading Preinstalled Software

Modules For Preinstalled Software: Ex. Python3

```
epalmer@perlmutter:login36:~> $ python --version
```

```
Python 2.7.18
```

```
epalmer@perlmutter:login36:~> $ module list
```

```
Currently Loaded Modules:
```

- | | | |
|---------------------|-----------------------|--------------------------|
| 1) gcc/11.2.0 | 4) libfabric/1.15.0.0 | 7) cray-libsci/21.08.1.2 |
| 2) craype/2.7.16 | 5) craype-network-ofi | 8) PrgEnv-gnu/8.3.3 |
| 3) cray-dsmml/0.2.2 | 6) cray-mpich/8.1.17 | |

```
epalmer@perlmutter:login36:~> $ module load python
```

```
epalmer@perlmutter:login36:~> $ module list
```

```
Currently Loaded Modules:
```

- | | | |
|---------------------|-----------------------|---------------------------------------|
| 1) gcc/11.2.0 | 4) libfabric/1.15.0.0 | 7) cray-libsci/21.08.1.2 |
| 2) craype/2.7.16 | 5) craype-network-ofi | 8) PrgEnv-gnu/8.3.3 |
| 3) cray-dsmml/0.2.2 | 6) cray-mpich/8.1.17 | 9) python/3.9-anaconda-2021.11 |

```
epalmer@perlmutter:login36:~> $ python --version
```

```
Python 3.9.7
```

172 Modules Currently Available on Perlmutter

Nsight-Compute: Nsight-Compute/2022.1.1
Nsight-Systems: Nsight-Systems/2022.2.1
OpenCoarrays: OpenCoarrays/2.10.1
PrgEnv-aocc: PrgEnv-aocc/8.3.3
PrgEnv-cray: PrgEnv-cray/8.3.3
PrgEnv-gnu: PrgEnv-gnu/8.3.3
PrgEnv-intel: PrgEnv-intel/8.3.3
PrgEnv-llvm: PrgEnv-llvm/0.1
PrgEnv-vidia: PrgEnv-vidia/8.3.3
R: R/4.2.3
amber:
aocc: aocc/3.2.0
aocc-mixed: aocc-mixed/3.2.0
arm-forge:
atp:
berkeleygw:
bupc: bupc/2022.10.0
bupc-narrow: bupc-narrow/2022.10.0
cce:
cce-mixed:
chapel: chapel/1.30.0
climate-utils:
cmake:
codee:
common-utils: common-utils/2023q2
conda:
contrib: contrib/1.0
cpe:
cpe-cuda:
cpu: cpu/1.0
cray-R: cray-R/4.2.1.1
cray-ccdb: cray-ccdb/4.12.13
cray-cti:
cray-dsmml: cray-dsmml/0.2.2
cray-dyninst: cray-dyninst/12.1.1
cray-fftw:
cray-hdf5:
cray-hdf5-parallel:
cray-libpals:
cray-libsci:
cray-lustre-client-ofd:
cray-lustre-client-ofd/2.15.0.4_rc2_cray_178_gf28cb6f-2...
cray-mpich:
cray-mpich-abi:

cray-mpich-abi-pre-intel-5.0:
cray-mpich-ucx:
cray-mpich-ucx-abi:
cray-mpixlate:
cray-mnnet: cray-mnnet/5.0.4
cray-netcdf:
cray-netcdf-hdf5parallel:
cray-openshmemx:
cray-pals:
cray-parallel-netcdf:
cray-pmi:
cray-python: cray-python/3.9.13.1
cray-stat: cray-stat/4.11.13
cray-ucx:
craype:
craype-accel-host: craype-accel-host
craype-accel-vidia70: craype-accel-vidia70
craype-accel-vidia80: craype-accel-vidia80
craype-dl-plugin-ftp: craype-dl-plugin-ftp/22.06.1.2
craype-dl-plugin-py3:
craype-hugepages128M: craype-hugepages128M
craype-hugepages16M: craype-hugepages16M
craype-hugepages1G: craype-hugepages1G
craype-hugepages256M: craype-hugepages256M
craype-hugepages2G: craype-hugepages2G
craype-hugepages2M: craype-hugepages2M
craype-hugepages32M: craype-hugepages32M
craype-hugepages4M: craype-hugepages4M
craype-hugepages512M: craype-hugepages512M
craype-hugepages64M: craype-hugepages64M
craype-hugepages8M: craype-hugepages8M
craype-network-none: craype-network-none
craype-network-ofi: craype-network-ofi
craype-network-ucx: craype-network-ucx
craype-x86-milan: craype-x86-milan
craype-x86-milan-x: craype-x86-milan-x
craype-x86-rome: craype-x86-rome
craypkg-gen:
cudatoolkit:
cudnn:
darshan:
dmtcp: dmtcp/3.0.0
dvs: dvs/2.15_4.4.232-2.4_1.1__geddbfef3

e4s:
eigen: eigen/3.4.0
espresso:
evp-patch: evp-patch
fast-mkl-amd: fast-mkl-amd/fast-mkl-amd
forge:
fpm: fpm/0.9.0
gcc:
gcc-mixed:
gdb4hpc:
globus-tools: globus-tools/1.0
gpu: gpu/1.0
gpu-test: gpu-test/1.1
gromacs:
gsl: gsl/2.7
hip:
idl: idl/8.5
impi: impi/2021.6.0
intel: intel/2023.1.0
intel-classic: intel-classic/2023.1.0
intel-classic-mixed: intel-classic-mixed/2023.1.0
intel-llvm: intel-llvm/2023-WW13
intel-mixed: intel-mixed/2023.1.0
intel-oneapi: intel-oneapi/2023.1.0
intel-oneapi-mixed: intel-oneapi-mixed/2023.1.0
iobuf: iobuf/2.0.10
jamo:
jgi: jgi/python-jamo
julia:
lammps: lammps/2022.11.03
libfabric: libfabric/1.15.2.0
llvm:
lmod: lmod
math-libreal32: math-libreal32/2023q2
mathematica: mathematica/13.0.1
matlab: matlab/R2021b
matlab-mcr: matlab-mcr/R2021b
mongodb:
mpich: mpich/4.1.1
mumps: mumps/5.5.1-gcc-11.2.0
mvasp: mvasp/5.4.4-cpu
namd:
nccl:

nersc_cr: nersc_cr/23.06
nvhpc:
nvhpc-mixed:
nvidia:
nvidia-mixed:
opencoarrays: opencoarrays/2.10.1
openmpi: openmpi/5.0.0rc12
papi:
parallel: parallel/20210922
paraview: paraview/5.11.1
perftools: perftools
perftools-base:
perftools-lite: perftools-lite
perftools-lite-events: perftools-lite-events
perftools-lite-gpu: perftools-lite-gpu
perftools-lite-hbm: perftools-lite-hbm
perftools-lite-loops: perftools-lite-loops
perftools-preload: perftools-preload
petsc: petsc/3.19.3-cpu-complex
pwanalyzer:
python:
pytorch:
qchem:
sanitizers4hpc:
settag: settag
spack:
spin: spin/2.0
taskfarmer: taskfarmer/1.5
tensorflow:
texlive: texlive/2022
totalview:
training: training/perlmutter-jan2022
upcxx:
upcxx-cuda:
upcxx-extras: upcxx-extras
valgrind:
valgrind4hpc:
vasp:
vasp-tpc:
wannier90:
wrf:
xalt: xalt/2.10.2
xpmem: xpmem/2.5.2-2.4_3.50__gd0f7936.shasta



Modules Loaded at Login

Modules Loaded by Default:

- | | | |
|--|----------------------------|-----------------------------|
| 1) craype-x86-milan | 7) cray-libsci/23.02.1.1 | 13) xalt/2.10.2 |
| 2) libfabric/1.15.2.0 | 8) cray-mpich/8.1.25 | 14) Nsight-Compute/2022.1.1 |
| 3) craype-network-ofi | 9) craype/2.7.20 | 15) Nsight-Systems/2022.2.1 |
| 4) xpmem/2.5.2-2.4_3.50__gd0f7936.shasta | 10) gcc/11.2.0 | 16) cudatoolkit/11.7 |
| 5) PrgEnv-gnu/8.3.3 | 11) perftools-base/23.03.0 | 17) craype-accel-nvidia80 |
| 6) cray-dsmml/0.2.2 | 12) cpe/23.03 | 18) gpu/1.0 |

- CPU Architecture
- Default Programming Environment and Compiler
- GPU Architecture and CUDA-Aware MPI

Modules with Lmod

Most Common

- `module list`
- `module load/unload`
- `module swap`
- `module show`
- `module spider`



Cool Tricks

- `module --redirect -r spider . | grep <string>`
- `ml -t`

More information: `man module` or <https://docs.nersc.gov/environment/lmod/>

Modules with Lmod

No Longer
Recommended

- `module avail`

*Only shows packages that can be loaded into the current module environment (due to hierarchy) – use `module spider` instead

```
epalmer@perlmutter:login34:~> $
```

More information: `man module` or <https://docs.nersc.gov/environment/lmod/>

Modules with Lmod

No Longer
Recommended

- `module avail`

*Only shows packages that can be loaded into the current module environment (due to hierarchy) – use `module spider` instead

```
epalmer@perlmutter:login34:~> $
```

More information: `man module` or <https://docs.nersc.gov/environment/lmod/>

Loading Modules Modifies Your Environment

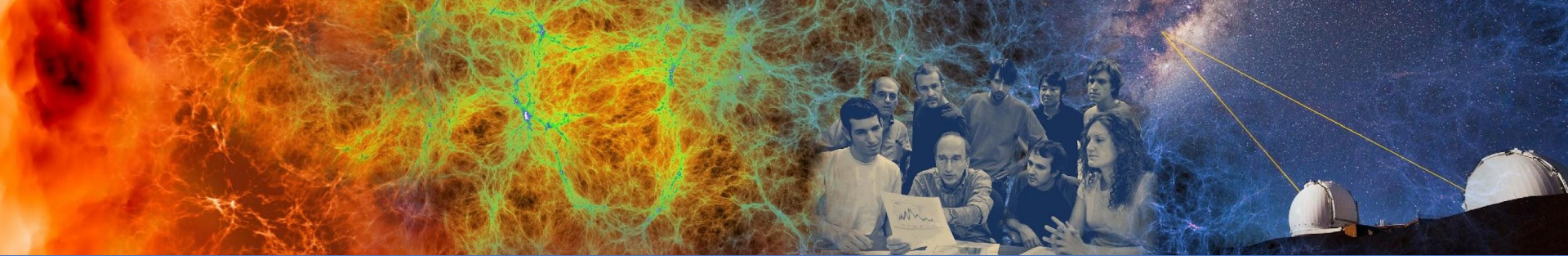
```
epalmer@perlmutter:login25:~/Training> $ module show cray-hdf5
```

```
-----  
/opt/cray/pe/lmod/modulefiles/compiler/gnu/8.0/cray-hdf5/1.12.1.5.lua:  
-----
```

```
family("hdf5")  
conflict("PrgEnv-pathscale")  
help([[Release info: /opt/cray/pe/hdf5/1.12.1.5/release_info]])  
whatis("The HDF5 Technology suite includes tools and applications for managing, manipulating, viewing, and  
prepending paths to the PATH environment variable, and prepending paths to the PKG_CONFIG_PATH, PE_PKGCONFIG_LIBS, and PE_HDF5_PKGCONFIG_LIBS environment variables, and prepending paths to the PE_FORTRAN_PKGCONFIG_LIBS, PE_CXX_PKGCONFIG_LIBS, and CRAY_HDF5_DIR environment variables, and setting the CRAY_HDF5_VERSION, CRAY_HDF5_PREFIX, HDF5_DIR, and HDF5_ROOT environment variables, and prepending paths to the CRAY_LD_LIBRARY_PATH and MODULEPATH environment variables.")  
prepend_path("PATH", "/opt/cray/pe/hdf5/1.12.1.5/bin")  
prepend_path("PKG_CONFIG_PATH", "/opt/cray/pe/hdf5/1.12.1.5/gnu/9.1/lib/pkgconfig")  
prepend_path("PE_PKGCONFIG_LIBS", "hdf5_hl:hdf5")  
setenv("PE_HDF5_PKGCONFIG_LIBS", "hdf5_hl:hdf5")  
prepend_path("PE_FORTRAN_PKGCONFIG_LIBS", "hdf5hl_fortran:hdf5_fortran")  
setenv("PE_HDF5_FORTRAN_PKGCONFIG_LIBS", "hdf5hl_fortran:hdf5_fortran")  
prepend_path("PE_CXX_PKGCONFIG_LIBS", "hdf5_hl_cpp:hdf5_cpp")  
setenv("PE_HDF5_CXX_PKGCONFIG_LIBS", "hdf5_hl_cpp:hdf5_cpp")  
setenv("CRAY_HDF5_DIR", "/opt/cray/pe/hdf5/1.12.1.5")  
setenv("PE_HDF5_DIR", "/opt/cray/pe/hdf5/1.12.1.5")  
setenv("CRAY_HDF5_VERSION", "1.12.1.5")  
setenv("CRAY_HDF5_PREFIX", "/opt/cray/pe/hdf5/1.12.1.5/gnu/9.1")  
setenv("HDF5_DIR", "/opt/cray/pe/hdf5/1.12.1.5/gnu/9.1")  
setenv("HDF5_ROOT", "/opt/cray/pe/hdf5/1.12.1.5/gnu/9.1")  
prepend_path("CRAY_LD_LIBRARY_PATH", "/opt/cray/pe/hdf5/1.12.1.5/gnu/9.1/lib")  
prepend_path("MODULEPATH", "/opt/cray/pe/lmod/modulefiles/hdf5/gnu/8.0/cray-hdf5/1.12.1")
```



Path Changes
Environment Variables
Other Info



Programming Environments: Configuring Compilers and Libraries



BERKELEY LAB



U.S. DEPARTMENT OF
ENERGY

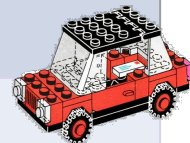
Office of
Science

Compiling: Your Machine vs. Perlmutter

What you may have done on another machine:

```
epalmer@home_machine:~> gcc helloworld.c -o helloworld.ex
```

- Not MPI-enabled code.



```
epalmer@home_cluster:~> mpicc helloworld.c -o helloworld.ex
```

- MPI compiler wrapper includes MPI libraries



What we recommend for Perlmutter:

```
epalmer@perlmutter:~> module load PrgEnv-gnu  
epalmer@perlmutter:~> cc helloworld.c -o helloworld.ex
```

- HPE compiler wrapper includes MPI libraries, optimizations and more.



Compiler Wrappers and a Useful Option

-v and -craype-verbose will show all the inputs added by the craype (Cray Programming Environment) to the compiler when the wrappers (CC, cc, ftn) are used.

```
epalmer@nid005015:~/Training> cc -craype-verbose helloworld.c -o helloworld.ex
```

```
gcc -march=znver3 -D__CRAY_X86_MILAN -D__CRAY_NVIDIA80  
-D__CRAYXT_COMPUTE_LINUX_TARGET -D__TARGET_LINUX__ helloworld.c -o  
helloworld.ex -Wl,-rpath=/opt/cray/pe/gcc-libs -Wl,-Bdynamic  
-I/opt/cray/pe/mpich/8.1.17/ofi/gnu/9.1/include  
-I/opt/cray/pe/libsci/21.08.1.2/GNU/9.1/x86_64/include  
-I/opt/cray/pe/dsmml/0.2.2/dsmml//include  
-I/opt/nvidia/hpc_sdk/Linux_x86_64/22.5/cuda/11.7/nvvm/include  
-I/opt/nvidia/hpc_sdk/Linux_x86_64/22.5/cuda/11.7/extras/CUPTI/include ...(and more)
```

Compiler Wrapper Includes A Lot

```
epalmer@nid005015:~/Training> cc -craype-verbose helloworld.c -o hello
```

```
gcc -march=znver3 -D__CRAY_X86_MILAN -D__CRAY_NVIDIA80 -D__CRAYXT_COMPUTE_LINUX_TARGET  
-D__TARGET_LINUX__ helloworld.c -fopenmp -o hello -Wl,-rpath=/opt/cray/pe/gcc-libs -Wl,-Bdynamic  
-l/opt/cray/pe/mpich/8.1.17/ofi/gnu/9.1/include -l/opt/cray/pe/libsci/21.08.1.2/GNU/9.1/x86_64/include  
-l/opt/cray/pe/dsmml/0.2.2/dsmml//include -l/opt/nvidia/hpc_sdk/Linux_x86_64/22.5/cuda/11.7/nvvm/include  
-l/opt/nvidia/hpc_sdk/Linux_x86_64/22.5/cuda/11.7/extras/CUPTI/include  
-l/opt/nvidia/hpc_sdk/Linux_x86_64/22.5/cuda/11.7/extras/Debugger/include  
-l/opt/cray/xpmem/2.4.4-2.3_12.2_gff0e1d9.shasta/include -L/opt/cray/pe/mpich/8.1.17/ofi/gnu/9.1/lib  
-L/opt/cray/pe/mpich/8.1.17/gtl/lib -L/opt/cray/pe/libsci/21.08.1.2/GNU/9.1/x86_64/lib -L/opt/cray/pe/dsmml/0.2.2/dsmml//lib  
-L/opt/nvidia/hpc_sdk/Linux_x86_64/22.5/cuda/11.7/lib64/stubs -L/opt/nvidia/hpc_sdk/Linux_x86_64/22.5/cuda/11.7/lib64  
-L/opt/nvidia/hpc_sdk/Linux_x86_64/22.5/cuda/11.7/nvvm/lib64  
-L/opt/nvidia/hpc_sdk/Linux_x86_64/22.5/cuda/11.7/extras/CUPTI/lib64  
-L/opt/nvidia/hpc_sdk/Linux_x86_64/22.5/cuda/11.7/extras/Debugger/lib64  
-L/opt/nvidia/hpc_sdk/Linux_x86_64/22.5/math_libs/11.7/lib64 -L/global/common/software/nersc/pm-2021q4/sw/darshan/3.3.1/lib  
-L/opt/cray/xpmem/2.4.4-2.3_12.2_gff0e1d9.shasta/lib64 -Wl,--as-needed,-lcupti,-lcudart,--no-as-needed -lcuda  
-Wl,-rpath=/global/common/software/nersc/pm-2021q4/sw/darshan/3.3.1/lib -Wl,-no-as-needed -ldarshan -lz  
-Wl,--as-needed,-lsci_gnu_82_mpi_mp,--no-as-needed -Wl,--as-needed,-lsci_gnu_82_mp,--no-as-needed -ldl  
-Wl,--as-needed,-lmpi_gnu_91,--no-as-needed -lmpi_gtl_cuda -Wl,--as-needed,-ldsmml,--no-as-needed -lxpmem  
-Wl,--as-needed,-lgfortran,-lquadmath,--no-as-needed -Wl,--as-needed,-lmvec,--no-as-needed -Wl,--as-needed,-lm,--no-as-needed  
-Wl,--as-needed,-lpthread,--no-as-needed -Wl,--disable-new-dtags
```

Same Wrapper, Different Compiler

```
epalmer@perlmutter:~> module load PrgEnv-nvidia  
epalmer@perlmutter:~> cc -craype-verbose helloworld.c -o helloworld.ex
```

```
nvc -tp=zen3 -acc -gpu=cc80 -D__CRAY_X86_MILAN -D__CRAY_NVIDIA80  
-D__CRAYXT_COMPUTE_LINUX_TARGET -pgf90libs helloworld.c -o helloworld.ex  
-l/opt/cray/pe/mpich/8.1.25/of/nvidia/20.7/include ...
```

```
epalmer@perlmutter:~> module load PrgEnv-intel  
epalmer@perlmutter:~> cc -craype-verbose helloworld.c -o helloworld.ex
```

```
icx -march=core-avx2 -mtune=core-avx2 -D__CRAY_X86_MILAN -D__CRAY_NVIDIA80  
-D__CRAYXT_COMPUTE_LINUX_TARGET helloworld.c -o helloworld.ex  
-Wl,-rpath=/opt/intel/oneapi/compiler/2023.1.0/linux/compiler/lib/intel64  
-l/opt/cray/pe/mpich/8.1.25/of/intel/19.0/include ...
```

PrgEnvs, Compilers and Libraries

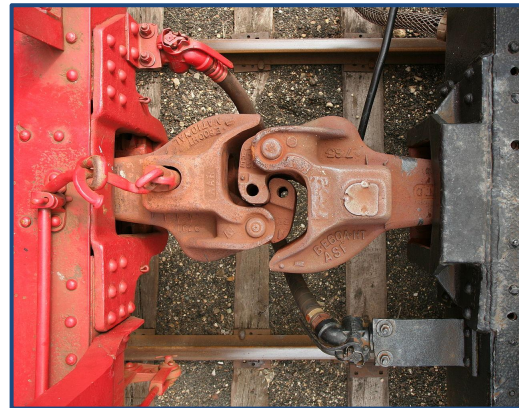
| Module | Compiler | C++ | | C | | Fortran | | MPI Library |
|---------------|----------|-----|-------------------|----|-------------------|---------|-----------|-------------|
| PrgEnv-gnu | GNU | CC | g++ | cc | gcc | ftn | gfortran | cray-mpich |
| PrgEnv-nvidia | NVHPC | CC | nvc++ | cc | nvc | ftn | nvfortran | cray-mpich |
| PrgEnv-intel | Intel | CC | icpx | cc | icx | ftn | ifort | cray-mpich |
| PrgEnv-cray | Cray | CC | crayCC (Clang) | cc | craycc (Clang) | ftn | crayftn | cray-mpich |

For additional compilers and more info: <https://docs.nersc.gov/development/compilers/base/>

Automatic Links Provided By The Wrappers

- Depending on modules loaded, compiler wrappers link:

MPI, LAPACK, Blas, ScaLAPACK,
and more, **automatically**.



- Cray modules, such as `cray-hdf5`, `cray-fftw`, etc. are also linked automatically by the compiler wrappers when loaded into the user environment.

Note: Several scientific libraries such as, LAPACK, ScaLAPACK, Blas and QDWH, are included in `cray-libsci`. For more information use: `man intro_libsci`.

Manually Specify Cray Compile Wrappers

Build systems such as CMake or Autotools (Makefiles) may be coded to search for the environment variables:

CC, for the C compiler ;
CXX, for the C++ compiler;
FC, for the Fortran compiler .

In this case, the Cray compile wrappers can be specified with:

```
CC=$(which cc) CXX=$(which CC) FC=$(which ftn)
```

Or at the configure step,

```
./configure CC=cc CXX=CC FC=ftn F77=ftn
```

More info: <https://docs.nersc.gov/development/build-tools/autoconf-make/>
<https://docs.nersc.gov/development/build-tools/cmake/>

Example: Compile Directions for SLATE

From INSTALL.md:

```
Configure and compile the SLATE library and its tester,  
then install the headers and library. This will also compile  
BLAS++, LAPACK++, and TestSweeper.
```

```
**Option 1: Makefile**
```

```
# create make.inc file, for example:  
CXX = mpicxx      # MPI compiler wrappers recommended  
FC   = mpif90  
blas = openblas  
CXXFLAGS = -DSLATE_HAVE_MT_BCAST
```

```
Compile and install:
```

```
make && make install
```

This build will require specifying:

CC=cc

CXX=CC

FC=ftn

Two Comments about Linking:

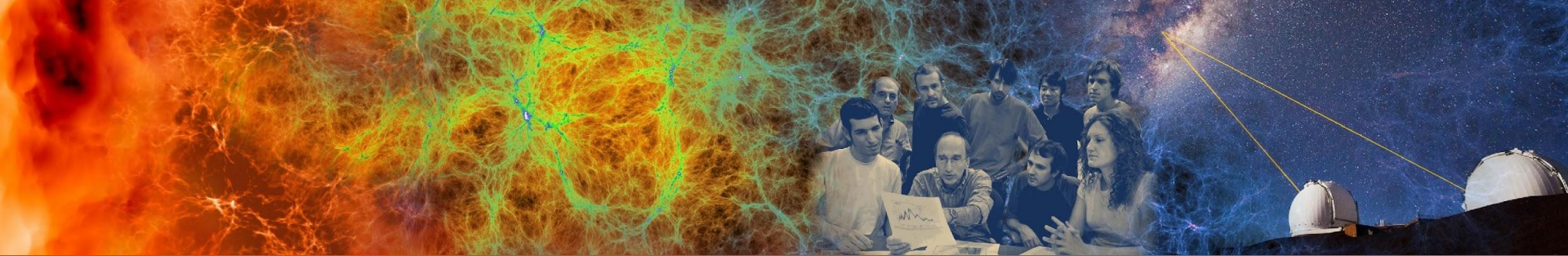
- Many modules prepend the `LIBRARY_PATH` so it is not necessary to specify library locations, e.g.

```
elvis@login01:~> $ module load gsl
elvis@login01:~> $ CC gsl_test.cpp -lgsl -lgslcblas -o gsl_test
```

Dynamic vs. Static Linking

- Cray wrappers build dynamically linked executables by default. (The kind that give, “error while loading shared libraries: ...” when they get misplaced at runtime.)
- On Perlmutter static compilation with `-static` or `CRAYPE_LINK_TYPE=static` can fail and is not supported.

More info: <https://docs.nersc.gov/development/compilers/wrappers/>



Examples of Compiling Code

Example CPU Code Compile with MPI and OpenMP

```
int main(int argc, char *argv[]) {
    int numprocs, rank, namelen;
    char processor_name[MPI_MAX_PROCESSOR_NAME];
    int iam = 0, np = 1;
    MPI_Init(&argc, &argv);
    MPI_Comm_size(MPI_COMM_WORLD, &numprocs);
    MPI_Comm_rank(MPI_COMM_WORLD, &rank);
    MPI_Get_processor_name(processor_name, &namelen);

    #pragma omp parallel default(shared) private(iam, np)
    {
        np = omp_get_num_threads();
        iam = omp_get_thread_num();
        printf("Hello from thread %d out of %d from process
              %d out of %d on %s\n",
              iam, np, rank, numprocs, processor_name);
    }
    MPI_Finalize();
}
```

Hellohybrid.c – contains both MPI and OpenMP commands.

Example from:

<https://rcc.uchicago.edu/docs/runnning-jobs/hybrid/index.html>

Some Good-to-Know Compiler Settings

| GNU | Cray | Nvidia | Description/ Comment |
|----------|---------------|-------------------------------------|--|
| -O0 | -O0 | -O1 | Default Optimization Level |
| -Ofast | -Ofast, -flto | -O4, -fast | Aggressive Optimization (some may cause non-bit-identical output) |
| -fopenmp | -fopenmp | -mp (CPU), -mp=gpu (GPU offload) | Enable OpenMP (not default) |
| | | -acc | Enable OpenACC |
| -g, -O0 | -g, -O0 | -g (-O0 by default) | Debug |
| -v | -v | -v | Verbose |

For more information:

man gcc/gfortran

man craycc/crayftn

man nvc/nvfortran

<https://docs.nersc.gov/development/compilers/>

Compile with MPI and OpenMP (CPU only)

```
epalmer@nid006368:~/NERSC_User_Training/EX1> #In this example, we will compile a  
code with MPI and OpenMP
```

Modules Loaded:
PrgEnv-gnu

Compile line: `cc hellohybrid.c -fopenmp -o hellohybrid`

Compile with MPI and OpenMP (CPU only)

```
epalmer@nid006368:~/NERSC_User_Training/EX1> █
```

Modules Loaded:
PrgEnv-gnu

Compile line: `cc hellohybrid.c -fopenmp -o hellohybrid`

Compile with MPI and OpenMP on GPU

```
epalmer@nid003676:~/NERSC_User_Training/EX1> ls
hellohybrid.c
epalmer@nid003676:~/NERSC_User_Training/EX1> module list

Currently Loaded Modules:
  1) craype-x86-milan          10) cudatoolkit/11.7
  2) libfabric/1.15.0.0       11) craype-accel-nvidia80
  3) craype-network-ofi      12) gpu/1.0
  4) perftools-base/22.06.0  13) nvidia/22.5
  5) xpmem/2.4.4-2.3_12.2__gff0e1d9.shasta 14) craype/2.7.16
  6) xalt/2.10.2              15) cray-dsmml/0.2.2
  7) darshan/3.3.1           16) cray-mpich/8.1.17
  8) Nsight-Compute/2022.1.1 17) cray-libsci/21.08.1.2
  9) Nsight-Systems/2022.2.1 18) PrgEnv-nvidia/8.3.3

epalmer@nid003676:~/NERSC_User_Training/EX1> cc
```

Modules Loaded:
PrgEnv-nvidia,
gpu

“-Minfo” is an optional command shows which parts were converted to NVIDIA GPU code.

Compile line: `cc hellohybrid.c -mp=gpu -Minfo -o hellohybrid`

Compile with MPI and OpenMP on GPU

```
epalmer@nid003676:~/NERSC_User_Training/EX1> █
```

Modules Loaded:
PrgEnv-nvidia,
gpu

“-Minfo” is an optional command shows which parts were converted to NVIDIA GPU code.

```
Compile line: cc hellohybrid.c -mp=gpu -Minfo -o hellohybrid
```

CUDA-Aware MPI

Modules Loaded by Default:

| | | |
|--|----------------------------|-----------------------------|
| 1) craype-x86-milan | 7) cray-libsci/23.02.1.1 | 13) xalt/2.10.2 |
| 2) libfabric/1.15.2.0 | 8) cray-mpich/8.1.25 | 14) Nsight-Compute/2022.1.1 |
| 3) craype-network-ofi | 9) craype/2.7.20 | 15) Nsight-Systems/2022.2.1 |
| 4) xpmem/2.5.2-2.4_3.50__gd0f7936.shasta | 10) gcc/11.2.0 | 16) cudatoolkit/11.7 |
| 5) PrgEnv-gnu/8.3.3 | 11) perftools-base/23.03.0 | 17) craype-accel-nvidia80 |
| 6) cray-dsmml/0.2.2 | 12) cpe/23.03 | 18) gpu/1.0 |

Output of module show gpu:

```
-----  
/opt/nersc/pe/modulefiles/gpu/1.0.lua:  
-----  
family("hardware")  
load("cudatoolkit")  
load("craype-accel-nvidia80")  
setenv("MPICH_GPU_SUPPORT_ENABLED", "1")
```

Note: The gpu module enables support for CUDA-Aware MPI – Allowing MPI to copy data to and from GPUs. module load cpu will turn it off.

Compile a CUDA Code with CUDA-Aware MPI

```
epalmer> # In this example, we will compile a CUDA code with CUDA-aware MPI
epalmer> ls
kernels.cu  kernels.h  vecAdd.cpp
epalmer> module list
```

Modules Loaded:
PrgEnv-nvidia,
gpu

Note: The flag, `MPICH_GPU_SUPPORT_ENABLED` turns CUDA-Aware MPI on or off. Loading the `gpu` module, sets this flag to 1, thereby enabling the feature.

Compile a CUDA Code with CUDA-Aware MPI

```
epalmer> █
```

Modules Loaded:
PrgEnv-nvidia,
gpu

Note: The flag, `MPICH_GPU_SUPPORT_ENABLED` turns CUDA-Aware MPI on or off. Loading the `gpu` module, sets this flag to 1, thereby enabling the feature.

Compile Commands for Code with OpenACC

Modules Loaded:

PrgEnv-nvidia,
gpu

Compile with OpenACC enabled:

```
cc helloacc.c -acc -Minfo=acc -o helloacc
```

*Optional command shows which parts were converted to NVIDIA GPU code.

Manually Specify Include, Library Location and Links

```
epalmer> # In this example, we will show how to manually include and link libraries during  
the compile step. The example code I will use, requires [
```


Manually Specify Include, Library Location and Links

```
epalmer> []
```

More Resources on Compiling Code

NERSC Docs:

- Compiling and Building Software:

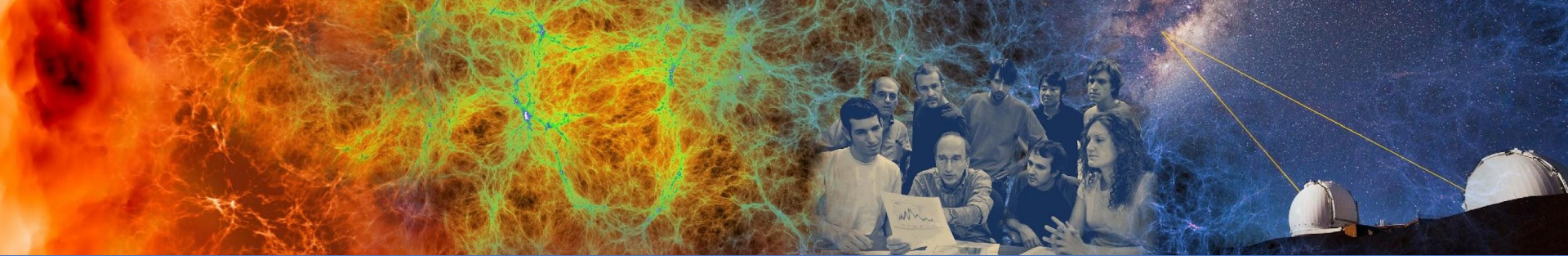
<https://docs.nersc.gov/systems/perlmutter/#compilingbuilding-software>

- Base Compilers: 

<https://docs.nersc.gov/development/compilers/base/#base-compilers-on-nersc-systems>

- Compiler Wrappers:

<https://docs.nersc.gov/development/compilers/wrappers/#compiler-wrappers>



Spack and E4S: Even More Software!



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Spack and The Extreme-Scale Scientific Software Stack (E4S)



| module | Version | Command | Installed Packages | Can I Install More? |
|-----------|---------|-----------------------|--------------------|---------------------|
| spack | 0.19.2 | module load spack | 532 of 6752 | Yes |
| e4s | 22.11 | module load e4s | 531 | Yes |
| e4s/23.05 | 23.05 | module load e4s/23.05 | 680 | Yes |

- E4S at NERSC is a curated software stack that get additional testing and support – delivered via the SPACK package manager

More info: Spack – <https://docs.nersc.gov/development/build-tools/spack/>

E4S – <https://docs.nersc.gov/applications/e4s/>

Install or Load a Spack Package

| Steps | Already Installed in Spack/E4S | Available via Spack/E4S |
|-------|---|--|
| 1 | <code>spack env activate <env></code> | |
| | Select an environment. *Needed for E4S modules, not required for Spack module | |
| 2 | <code>spack find -v</code> | <code>spack list</code> |
| | Match package config. and compiler | Search list of ~6700 avail. packages |
| 3 | <code>spack load <package></code> | <code>spack info <package></code> |
| | Load desired package/variant | Get information about package options |
| 4 | | <code>spack install <package></code> |
| | | Installs the package into Spack |
| 5 | | <code>spack load <package></code> |

Bonus: `spack load --sh <package>`

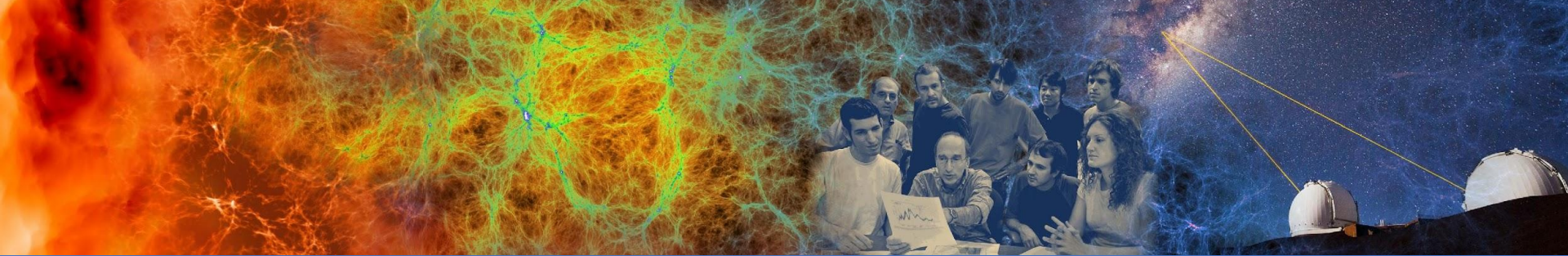
Show modifications made to your environment when a package is loaded.

Example: Install a Dependency with Spack

```
+phalana+piro+rol+ryttaa+saacae+shanda+shyly+stak+strakho+stratimikos+superia-di
st+teko+tepus+tperra+trifinoscouplings+zoltan+zoltan2_gotype=long_long
umap@2.1.0%gcc@11.2.0
umpire@2022.03.1%gcc@11.2.0
upcxx@2022.0.0%gcc@11.2.0
vtk-m@1.9.0%gcc@11.2.0
zfp@0.5.5%gcc@11.2.0

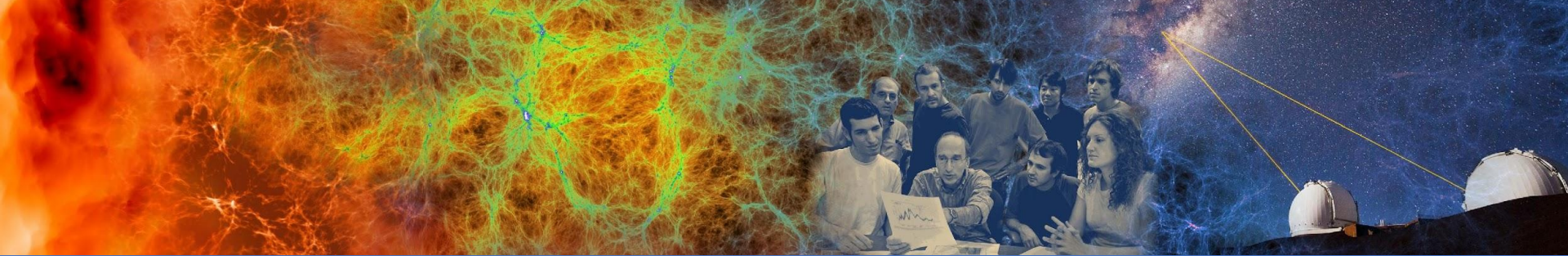
==> Installed packages
-- linux-sles15-zen3 / gcc@11.2.0 -----
gsl@2.7.1-external-cblas build.system=autotools
==> 1 installed package
epalmer># There it is
epalmer># To get the location of the libraries I need I will use a spack command
and store the output in a bash variable
epalmer>export GSL_ROOT=$(spack location -i gsl@2.7.1)
epalmer>echo $GSL_ROOT
/global/common/software/spackecp/perlmutter/e4s-22.11/03104/spack/opt/spack/linu
x-sles15-zen3/gcc-11.2.0/gsl-2.7.1-usionwbcmq56cvupqse43j4yv275154d
epalmer>ls $GSL_ROOT
bin include lib share
epalmer># So I can use the variable to access the location
epalmer># Now I will compile
epalmer>
```

Suppose my example code, `gsl_test.cpp`, requires `gsl` ver. 2.7.1.



Key Suggestions:

- Use module spider
- Use compiler wrappers –CC, cc, and ftn– with PrgEnv modules
- Additional software is available via Spack and E4S



Thanks for your Attention!

Documentation for Programming Environments and Compilation:

- Modules – <https://docs.nersc.gov/environment/lmod/>
- Programming Environment & Compilers – <https://docs.nersc.gov/systems/perlmutter/#compilingbuilding-software>
- Shell Scripts – https://docs.nersc.gov/environment/shell_startup/
- Spack – <https://docs.nersc.gov/development/build-tools/spack/>
- E4S – <https://docs.nersc.gov/applications/e4s/perlmutter/22.05/>

More questions? Need help? ... <http://help.nersc.gov/>



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