

Modules & Application Software

The number of software packages that are installed together with the operating system on cluster nodes is kept light on purpose. Additional packages and applications are provided by a module system, which enables you to easily customise your working environment on the cluster. This module system is called Lmod¹⁾. Furthermore, we can provide different versions of the software which you can use on demand. Loading a module, software specific settings are applied, e.g. changing environment variables like PATH, LD_LIBRARY_PATH and MANPATH.

Alternatively, you can manage software packages on the cluster yourself by building software from source, by means of EasyBuild or by using Singularity containers. Python packages can also be installed using the Conda manager. The first three possibilities, in addition to the module system, are described in the current section, whereas Conda usage in the cluster is explained in [this section](#).

We have adopted a systematic software naming and versioning convention in conjunction with the software installation system EasyBuild²⁾.

Software installation on the cluster utilizes a hierarchical software module naming scheme. This means that the command `module avail` does not display all installed software modules right away. Instead, only the modules that are immediately available for loading are displayed. More modules become available after their prerequisite modules are loaded. Specifically, loading a compiler module or MPI implementation module will make available all the software built with those applications. This way, we hope the prerequisites for certain software become apparent.

At the top level of the module hierarchy, there are modules for compilers, toolchains and software applications that come as a binary and thus do not depend on compilers. Toolchain modules organize compilers, MPI implementations and numerical libraries. Currently the following toolchain modules are available:

- Compiler only toolchains
 - GCC: GCC updated binutils
 - iccifort: Intel compilers, GCC
- Compiler & MPI toolchains
 - gomp: GCC, OpenMPI
 - iimpi: iccifort, Intel MPI
 - iompi: iccifort, OpenMPI
- Compiler & MPI & numerical libraries toolchains
 - foss: gomp, OpenBLAS, FFTW, ScaLAPACK
 - intel: iimpi, Intel MKL
 - iomkl: iompi, Intel MKL

Note that Intel compilers newer than 2020.x are provided by the toolchain module `intel-compilers`. It is strongly recommended to use this module as after 2023 the intel compiler modules `iccifort` will be removed.

Working with modules

This section explains how to use software modules.

List the entire list of possible modules

```
module spider
```

The same in a more compact list

```
module -t spider
```

Search for specific modules that have “string” in their name

```
module spider string
```

Detailed information about a particular version of a module (including instructions on how to load the module)

```
module spider name/version
```

Searches for all module names and descriptions that contain the specified string

```
module key string
```

List modules immediately available to load

```
module avail
```

Some software modules are hidden from the `avail` and `spider` commands. These are mostly modules for system library packages that other user applications depend on. To list hidden modules, you may provide the `--show-hidden` option to the `avail` and `spider` commands:

```
module --show-hidden avail
module --show-hidden spider
```

A hidden module has a dot (.) in front of its version numbers (eg. `zlib/.1.2.8`).

List currently loaded modules

```
module list
```

Load a specific version of a module

```
module load name/version
```

If only a name is given, the command will load the default version which is marked with a (D) in the

`module avail` listing (usually the latest version). Loading a module may automatically load other modules it depends on.

It is not possible to load two versions of the same module at the same time.

To switch between different modules

```
module swap old new
```

To unload the specified module from the current environment

```
module unload name
```

To clean your environment of all loaded modules

```
module purge
```

Show what environment variables the module will set

```
module show name/version
```

Save the current list of modules to “name” collection for later use

```
module save name
```

Restore modules from collection “name”

```
module restore name
```

List of saved collections

```
module savelist
```

To get the complete list of options provided by Lmod through the command `module type` the following

```
module help
```

Exercise: Working with modules

As an example, we show how to load the `gnuplot` module.

List loaded modules

```
module list
```

```
No modules loaded
```

Find available `gnuplot` versions

```
module -t spider gnuplot
```

```
gnuplot/4.6.0  
gnuplot/5.0.3
```

Determine how to load the selected gnuplot/5.0.3 module

```
module spider gnuplot/5.0.3
```

```
-----  
----  
gnuplot: gnuplot/5.0.3  
-----  
----
```

Description:

Portable interactive, function plotting utility - Homepage:
<http://gnuplot.sourceforge.net/>

This module can only be loaded through the following modules:

GCC/4.9.3-2.25 OpenMPI/1.10.2

Help:

Portable interactive, function plotting utility - Homepage:
<http://gnuplot.sourceforge.net/>

Load required modules

```
module load GCC/4.9.3-2.25 OpenMPI/1.10.2
```

```
Module for GCCcore, version .4.9.3 loaded  
Module for binutils, version .2.25 loaded  
Module for GCC, version 4.9.3-2.25 loaded  
Module for numactl, version .2.0.11 loaded  
Module for hwloc, version .1.11.2 loaded  
Module for OpenMPI, version 1.10.2 loaded
```

And finally load the selected gnuplot module

```
module load gnuplot/5.0.3
```

```
Module for OpenBLAS, version 0.2.15-LAPACK-3.6.0 loaded  
Module for FFTW, version 3.3.4 loaded  
Module for ScaLAPACK, version 2.0.2-OpenBLAS-0.2.15-LAPACK-3.6.0 loaded  
Module for bzip2, version .1.0.6 loaded  
Module for zlib, version .1.2.8 loaded  
.....  
.....
```

In order to simplify the procedure of loading the gnuplot module, the current list of loaded modules can be saved in a “mygnuplot” collection (the name string “mygnuplot” is, of course, arbitrary) and then loaded again when needed as follows:

Save loaded modules to “mygnuplot”

```
module save mygnuplot
```

```
Saved current collection of modules to: mygnuplot
```

If “mygnuplot” not is specified, the name “default” will be used.

Remove all loaded modules (or open a new shell)

```
module purge
```

```
Module for gnuplot, version 5.0.3 unloaded
```

```
Module for Qt, version 4.8.7 unloaded
```

```
Module for libXt, version .1.1.5 unloaded
```

```
.....
```

```
.....
```

List currently loaded modules. This selection is empty now.

```
module list
```

```
No modules loaded
```

List saved collections

```
module savelist
```

```
Named collection list:
```

```
1) mygnuplot
```

Load gnuplot module again

```
module restore mygnuplot
```

```
Restoring modules to user's mygnuplot
```

```
Module for GCCcore, version .4.9.3 loaded
```

```
Module for binutils, version .2.25 loaded
```

```
Module for GCC, version 4.9.3-2.25 loaded
```

```
Module for numactl, version .2.0.11 loaded
```

```
.....
```

```
.....
```

List of available software

In this section, you will find user guides for some of the software packages installed in the cluster. The guides provided can, of course, not replace documentation that comes with the application. Please read that as well.

A wide variety of application software is available in the cluster system. These applications are located on a central storage system that is accessible by the module system Lmod via an NFS export. Issue the command `module spider` on the cluster system or visit the [page](#) for a comprehensive list of available software. If you really need a different version of an already installed application, or one that is currently not installed, please get in touch. The main prerequisite for use of a software within the cluster system is its availability for Linux. Furthermore, if the application requires a license, we need to clarify additional questions.

Some select Windows applications can also be executed on the cluster system with the help of Wine or Singularity containers. For information on Singularity, see [Singularity Containers](#).

A current list of available software

Usage instructions

- [Abaqus](#)
- [ANSYS / CFX](#)
- [Code-Server \(VS Code\) on LUIS Cluster](#)
- [COMSOL](#)
- [Conda](#)
- [CPMD](#)
- [FEKO](#)
- [Jupyter on LUIS Cluster](#)
- [MATLAB](#)
- [mpiFileUtils](#)
- [NFFT](#)
- [uv - Python package manager](#)

Build software from source code

Note: We recommend using [EasyBuild](#) (see next section) if you want to make your software's build process reproducible and accessible through a module environment that EasyBuild automatically creates. EasyBuild comes with pre-configured recipes for installing thousands of scientific applications.

Sub-clusters of the cluster system have different CPU architectures that provide different instruction set capabilities/extensions. The command `lcpuarchs` (available on the login nodes) lists all available

CPU types.

```
login03:~$ lcpuarchs -v

CPU arch names      Cluster partitions
-----
avx2                LUIS[haku, lena, vis, jhub]
                   FCH[ai, ai, ainlp, ainlp, gih, pci, fi, imuk, p4d, p4d]

avx512
LUIS[gpu, gpu, gpu, gpu, test, mpp, single, mpp, share, amo, taurus, smp, smp]
FCH[tnt, tnt, ai, h100, gih, isd, isd, isd, isd, stahl, iwes, iwes, enos, pcikoe, pcikoe, p
cikoe, pcikoe, pcikoe, isu, phd, phd, gpu, itp, itp, imes, gpu, imes]

LUIS - LUIS clusters. Available to all users around the clock
FCH  - Research Cluster Housing nodes. Available to all users
      on weekdays from 20:00 to 08:00 and on weekends

CPU of this machine: avx512

For more verbose output type: lcpuarchs -vv
```

The technical sequence of these architectures is (oldest) -sse-avx-avx2-avx512- (newest). Cpus capable of executing instructions from newer instruction sets usually are able to execute commands from older extensions, so e.g. avx512 includes sse.

Software compiled to use a newer cpu instruction set will usually not typically abort with an “**Illegal instruction**” error when run on an older cpu. The important message here is that compilers may automatically set flags for the platform you are currently working on. If you compile your program on a node providing avx512 instructions (e. g. the amo sub-cluster) using the gcc compiler option -march=native, the program will usually not run on older nodes that are only equipped with cpus providing, say, avx instructions. To check which instruction set extensions a cpu architecture provides, you can run the command “lscpu”, which lists them in the “flags” section.

This section explains how to build a software on the cluster system to avoid the aforementioned issue if you want to be able to submit jobs to all compute nodes without specifying the CPU type. Beware, though, that compatibility usually comes at a price and allowing a compiler to use the newer instruction sets will usually boost performance. Depending on your workload, the effects/speedup on newer cpu architectures may even be called “drastic”. But there's usually no better way to tell than testing.

In the example below we want to compile a sample software my-soft in version 3.1.

In your [HOME](#) (or, perhaps better, in your [\\$SOFTWARE](#) directory, if all members of your project want to use the software) directory, create build/install directories for each available CPU architecture listed by the command `lcpuarchs -s`, as well as a directory source to storing the installation sources

Note: you can usually refer to a variable using `$variable_name` and all will be well. In the following examples, however, we demonstrate the use of curly brackets around the variable's designator, which will ensure proper separation of variables even in case of ambiguities (which in theory could occur in

long paths composed out of several variables). For all normal purposes, `$HOME` and `${HOME}` or `$LUIS_CPU_ARCH` and `${LUIS_CPU_ARCH}` will be equivalent. If, however, you use spaces in your directories (like `dir="my directory"`), this will not be sufficient, you'll then also need to put double quotation marks " around the variable (like in `cd "${dir}"`).

```
login03:~$ mkdir -p ${HOME}/sw/source
login03:~$ eval "mkdir -p ${HOME}/sw/${$(lcpuarchs -ss)}/my-soft/3.1/{build,install}"
```

Copy software installation archive to the source directory

```
login03:~$ mv my-soft-3.1.tar.gz ${HOME}/sw/source
```

Build `my-soft` for each available CPU architecture by submitting an interactive job to each compute node type requesting the proper CPU type. For example, to compile `my-soft` for `avx512` nodes, first submit an interactive job requesting the `avx512` feature:

```
login03:~$ salloc --nodes=1 --constraint=CPU_ARCH:avx512 --cpus-per-task=4 --time=6:00:00 --mem=16G
```

Then unpack and build the software. Note the environment variable `${LUIS_CPU_ARCH}` that contains the cpu instruction set of the compute node reserved.

```
taurus-n034:~$ tar -zxvf ${HOME}/sw/source/my-soft-3.1.tgz -C
${HOME}/sw/${LUIS_CPU_ARCH}/my-soft/3.1/build
taurus-n034:~$ cd ${HOME}/sw/${LUIS_CPU_ARCH}/my-soft/3.1/build
taurus-n034:~$ ./configure --prefix=${HOME}/sw/${LUIS_CPU_ARCH}/my-soft/3.1/install && make && make install
```

Finally, use the environment variable `${LUIS_CPU_ARCH}` in your job scripts to access the correct installation path of `my-soft` executable for the current compute node. Note that you may need to set/update the `${LD_LIBRARY_PATH}` environment variable to point to the location of your software's shared libraries.

[my-soft-job.sh](#)

```
#!/bin/bash -l
#SBATCH --job-name=my-soft
#SBATCH --nodes=1
#SBATCH --ntasks-per-node=16
#SBATCH --mem=60G
#SBATCH --time=12:00:00
#SBATCH --constraint="[CPU_ARCH:avx512|CPU_ARCH:avx2]"
#SBATCH --output my-soft-job_%j.out
#SBATCH --error my-soft-job_%j.err
#SBATCH --mail-user=myemail@...uni-hannover.de
#SBATCH --mail-type=BEGIN,END,FAIL
```

```
# Change to work dir
cd ${SLURM_SUBMIT_DIR}

# Load modules
module load my_necessary_modules

# run my_soft
export LD_LIBRARY_PATH=${HOME}/sw/${LUIS_CPU_ARCH}/my-soft/3.1/install/lib:${LD_LIBRARY_PATH}
srun $HOME/sw/${LUIS_CPU_ARCH}/my-soft/3.1/install/bin/my-soft.exe --input file.input
```

You can certainly consider combining the software build and execution steps into a single batch job script. However, it is recommended that you first perform the build steps **interactively** before adding them to a job script to ensure the software compiles without errors. For example, such a job script might look like this:

my-soft-job.sh

```
#!/bin/bash -l
#SBATCH --job-name=my-soft
#SBATCH --nodes=1
#SBATCH --ntasks-per-node=32
#SBATCH --mem=120G
#SBATCH --time=12:00:00
#SBATCH --constraint=CPU_ARCH:avx512
#SBATCH --output my-soft-job_%j.out
#SBATCH --error my-soft-job_%j.err
#SBATCH --mail-user=myemail@...uni-hannover.de
#SBATCH --mail-type=BEGIN,END,FAIL

# Change to work dir
cd ${SLURM_SUBMIT_DIR}

# Load modules
module load my_necessary_modules

# install software if the executable does not exist
[ -e "${HOME}/sw/${LUIS_CPU_ARCH}/my-soft/3.1/install/bin/my-soft.exe" ] || {
  mkdir -p ${HOME}/sw/${LUIS_CPU_ARCH}/mysof/3.1/{build,install}
  tar -zxvf ${HOME}/sw/source/my-soft-3.1.tgz -C
  ${HOME}/sw/${LUIS_CPU_ARCH}/my-soft/3.1/build
  cd $HOME/sw/${LUIS_CPU_ARCH}/my-soft/3.1/build
  ./configure --prefix=${HOME}/sw/${LUIS_CPU_ARCH}/my-soft/3.1/install
  make
  make install
}

# run my_soft
```

```
export LD_LIBRARY_PATH=${HOME}/sw/${LUIS_CPU_ARCH}/my-soft/3.1/install/lib:${LD_LIBRARY_PATH}
srun ${HOME}/sw/${LUIS_CPU_ARCH}/my-soft/3.1/install/bin/my-soft.exe --input file.input
```

EasyBuild

Note: If you want to manually build the software from source code, please refer to the [section](#) above.

EasyBuild is a software build and installation framework that allows you to manage (scientific) software on High Performance Computing (HPC) systems in an efficient way.

EasyBuild framework

The EasyBuild framework is available in the cluster through the module EasyBuild-custom. This module defines the location of the EasyBuild configuration files, recipes and installation directories. You can load the module using the command:

```
module load EasyBuild-custom
```

EasyBuild software and modules will be installed by default under the following directory:

```
$HOME/my.soft/software/${LUIS_CPU_ARCH}
$HOME/my.soft/modules/${LUIS_CPU_ARCH}
```

Here, the variable ARCH, which stores the CPU type of the machine on which the above module load command was executed, will currently be either haswell, sandybridge or skylake. The command `lcpuarchs` executed on the cluster login nodes lists all currently available values of ARCH. You can override the default software and module installation directory, and the location of your EasyBuild configuration files (MY_EASYBUILD_REPOSITORY) by exporting the following environment variables before loading the EasyBuild module:

```
export EASYBUILD_INSTALLPATH=/your/preferred/installation/dir
export MY_EASYBUILD_REPOSITORY=/your/easybuild/repository/dir
module load EasyBuild-custom
```

If other project members should also have access to the software, the recommended location is a subdirectory in [\\$SOFTWARE](#).

How to build your software

After you load the EasyBuild environment as explained in the section above, you will have the command `eb` available to build your code using EasyBuild. If you want to build the code using a given

configuration `<filename>.eb` and resolving dependencies, you will use the flag `-r` as in the example below:

```
eb <filename>.eb -r
```

The build command just needs the configuration file name with the extension `.eb` and not the full path, provided that the configuration file is in your search path: the command `eb --show-config` will print the variable `robot-paths` that holds the search path. More options are available - please have a look at the short help message typing `eb -h`. For instance, using the search flag `-S`, you can check if any EasyBuild configuration file already exists for a given program name:

```
eb -S <program_name>
```

You will be able to load the modules created by EasyBuild in the directory defined by the `EASYBUILD_INSTALLPATH` variable using the following commands:

```
module use $EASYBUILD_INSTALLPATH/modules/${LUIS_CPU_ARCH}/all
module load <modulename>/version
```

The command `module use` will prepend the selected directory to your `MODULEPATH` environment variable, therefore the command `module avail` will show modules of your software as well.

If you want the software module to be automatically available when opening a new shell in the cluster, modify your `~/ .bashrc` file as follows:

```
echo 'export EASYBUILD_INSTALLPATH=/your/preferred/installation/dir' >>
~/ .bashrc
echo 'module use $EASYBUILD_INSTALLPATH/modules/${LUIS_CPU_ARCH}/all' >>
~/ .bashrc
```

Note that to preserve the dollar sign in the second line above, the string must be enclosed in single quotes.

Further Reading

- [EasyBuild documentation](#)
- [Easyconfigs repository](#)

Apptainer Containers (replaces Singularity)



Apptainer will replace Singularity on the the LUH-Clusters. Currently you can use both commands `apptainer` and `singularity` because the last one is a symlink to `apptainer`. This may change in the future.

Please note: This instruction has been written for Apptainer 1.3.3-*

Please note: If you would like to fully manage your apptainer container images directly on the cluster, including build and/or modify actions, please contact us and ask for the permission “apptainer fakeroot” to be added to your account (because you will need it).

Apptainer containers on the cluster

Apptainer enables users to execute containers on High-Performance Computing (HPC) cluster like they are native programs or scripts on a host computer. For example, if the cluster system is running CentOS Linux, but your application runs in Ubuntu, you can create an Ubuntu container image, install your application into that image, copy the image to an approved location on the cluster and run your application using Apptainer in its native Ubuntu environment.

The main advantage of Apptainer is that containers are executed as an unprivileged user on the cluster system and, besides the local storage TMPDIR, they can access the network storage systems like HOME, BIGWORK and PROJECT, as well as GPUs that the host machine is equipped with.

Additionally, Apptainer properly integrates with the Message Passing Interface (MPI), and utilizes communication fabrics such as InfiniBand and Intel Omni-Path.

If you want to create a container and set up an environment for your jobs, we recommend that you start by reading [the Apptainer documentation](#). The basic steps to get started are described below.

Building Apptainer container using a recipe file

If you already have a pre-build container ready for use, you can simply upload the container image to the cluster and execute it. See the [section](#) below about running container images.

Below we will describe how to build a new or modify an existing container directly on the cluster. A container image can be created from scratch using a recipe file, or fetched from some remote container repository. In this sub-section, we will illustrate a recipe file method. In the next one, we will take a glance at remote container repositories.

Using a Apptainer recipe file is the recommended way to create containers if you want to build reproducible container images. This example recipe file builds a RockyLinux 9 container:

[rocky9.def](#)

```
BootStrap: yum
OSVersion: 9
MirrorURL: https://ftp.uni-
hannover.de/rocky/{OSVERSION}/BaseOS/$basearch/os
Include: yum

%setup
  echo "This section runs on the host outside the container during
  bootstrap"
```

```
%post
echo "This section runs inside the container during bootstrap"

# install packages in the container
yum -y groupinstall "Development Tools"
yum -y install wget vim python3 epel-release
yum -y install python3-pip

# install tensorflow
pip3 install --upgrade tensorflow

# enable access to BIGWORK and PROJECT storage on the cluster system
mkdir -p /bigwork /project

%runscript
echo "This is what happens when you run the container"

echo "Arguments received: $*"
exec /usr/bin/python3 "$@"

%test
echo "This test will be run at the very end of the bootstrapping
process"

/usr/bin/python3 --version
```

This recipe file uses the yum bootstrap module to bootstrap the core operation system, RockyLinux 9, within the container. For other bootstrap modules (e.g.. docker) and details on apptainer recipe files, refer to [the online documentation](#).

The next step is to build a container image on one of the cluster login servers.

Note: your account must be authorized to use the --fakeroot option. Please contact us at cluster-help@luis.uni-hannover.de.

Note: Currently, the --fakeroot option is enabled only on the cluster login nodes.

```
username@login01$ apptainer build --fakeroot rocky9.sif rocky9.def
```

This creates an image file named `rocky9.sif`. By default, apptainer containers are built as read-only SIF (Apptainer Image Format) image files. Having a container in the form of a file makes it easier to transfer it to other locations both within the cluster and outside of it. Additionally, a SIF file can be signed and verified.

Note that a container as the SIF file can be built on any storage of the cluster you have a write access to. However, it is recommended to build containers either in your `$BIGWORK` or in some directory under `/tmp` (or use the variable `$MY_APPTAINER`) on the login nodes.

Note: Containers located only under the paths `$BIGWORK`, `$SOFTWARE` and `/tmp` are allowed to be executed using `shell`, `run` or `exec` commands, see the [section](#) below,

The latest version of the `apptainer` command can be used directly on any cluster node without prior activation.

Downloading containers from external repositories

Another easy way to obtain and use a Apptainer container is to retrieve pre-build images directly from external repositories. Popular repositories are [Docker Hub](#) or [Apptainer Library](#). You can go there and search if they have a container that meets your needs. For docker images, use the [search form at Docker Hub](#) instead.

In the following example we will pull the latest python container from Docker Hub and save it in a file named `python_latest.sif`:

```
username@login01$ apptainer pull docker://python:latest
```

The `build` sub-command can also be used to download images, where you can additionally specify your preferred container file name:

```
username@login01$ apptainer build my-ubuntu22.04.sif
library://library/default/ubuntu:22.04
```

How to modify existing Apptainer images

First you should check if you really need to modify the container image. For example, if you are using Python in an image and simply need to add new packages via `pip` you can do that without modifying the image by running `pip` in the container with the `--user` option.

To modify an existing SIF container file, you need to first convert it to a writable sandbox format.

Please note: Since the `--fakeroot` option of the `shell` and `build` sub-commands does not work with container sandbox when the container is located on a shared storage such as `BIGWORK`, `PROJECT` or `HOME`, the container sandbox must be stored locally on the login nodes. We recommend using the `/tmp` directory (or variable `$MY_APPTAINER`) which has sufficient capacity.

```
username@login01$ cd $MY_APPTAINER
username@login01$ apptainer build --sandbox rocky9-sandbox rocky9.sif
```

The `build` command above creates a sandbox directory called `rocky9-sandbox` which you can then `shell` into in writable mode and modify the container as desired:

```
username@login01$ apptainer shell --writable --fakeroot rocky9-sandbox
Apptainer> yum install -qy python3-matplotlib
```

After making all desired changes, you exit the container and convert the sandbox back to the SIF file using:

```
Apptainer> exit
username@login01$ apptainer build -F --fakeroot rocky9.sif rocky9-sandbox
```

Note: you can try to remove the sandbox directory *rocky9-sandbox* afterward but there might be a few files you can not delete due to the namespace mappings that happens. The daily */tmp* cleaner job will eventually clean it up.

Running container images

Please note: In order to run a Apptainer container, the container SIF file or sandbox directory must be located either in your *\$BIGWORK*, in your group's *\$SOFTWARE* or in the */tmp* directory.

There are four ways to run a container under Apptainer.

If you simple call the container image as an executable or use the Apptainer run sub-command it will carry out instructions in the *%runscript* section of the container recipe file:

How to call the container SIF file:

```
username@login01:~$ ./rocky9.sif --version
This is what happens when you run the container
Arguments received: --version
Python 3.8.6
```

Use the run sub-command:

```
username@login01:~$ apptainer run rocky9.sif --version
This is what happens when you run the container
Arguments received: --version
Python 3.8.6
```

The Apptainer *exec* sub-command lets you execute an arbitrary command within your container instead of just the *%runscript*. For example, to get the content of file */etc/os-release* inside the container:

```
username@login01:~$ apptainer exec rocky9.sif cat /etc/os-release
NAME="Rocky Linux"
VERSION="8.4 (Green Obsidian)"
....
```

The Apptainer *shell* sub-command invokes an interactive shell within a container. Note the *Apptainer>* prompt within the shell in the example below:

```
username@login01:$ apptainer shell rocky9.sif
Apptainer>
```

Note that all three sub-commands *shell*, *exec* and *run* let you execute a container directly from remote repository without first downloading it on the cluster. For example, to run an one-liner "Hello World" ruby program:

```
username@login01:$ aptainer exec library://sylabs/examples/ruby ruby -e 'puts "Hello World!";'
Hello World!
```

Please note: You can access (read & write mode) your HOME, BIGWORK and PROJECT (only login nodes) storage from inside your container. In addition, the /tmp (or TMPDIR on compute nodes) directory of a host machine is automatically mounted in a container. Additional mounts can be specified using the --bind option of the exec, run and shell sub-commands, see aptainer run --help.

Apptainer & parallel MPI applications

In order to containerize your parallel MPI application and run it properly on the cluster system you have to provide MPI library stack inside your container. In addition, the userspace driver for Mellanox InfiniBand HCAs should be installed in the container to utilize cluster InfiniBand fabric as a MPI transport layer.

This example Apptainer recipe file `ubuntu-openmpi.def` retrieves an Ubuntu container from Docker Hub, and installs required MPI and InfiniBand packages:

Ubuntu 20.04

[ubuntu-openmpi.def](#)

```
BootStrap: docker
From: ubuntu:focal

%post
# install openmpi & infiniband
apt-get update
apt-get -y install openmpi-bin openmpi-common libibverbs1 libmlx4-1

# enable access to BIGWORK storage on the cluster
mkdir -p /bigwork /project

# enable access to /scratch dir. required by mpi jobs
mkdir -p /scratch
```

Ubuntu 22.x - 24.x

[ubuntu-openmpi.def](#)

```
BootStrap: docker
```

```
From: ubuntu:latest

%post
# install openmpi & infiniband
apt-get update
apt-get -y install openmpi-bin openmpi-common ibverbs-providers

# enable access to BIGWORK storage on the cluster
mkdir -p /bigwork /project

# enable access to /scratch dir. required by mpi jobs
mkdir -p /scratch
```

Once you have built the image file `ubuntu-openmpi.sif` as explained in the previous sections, your MPI application can be run as follows (assuming you have already reserved a number of cluster compute nodes):

```
module load GCC/10.2.0 OpenMPI/4.0.5
mpirun aptainer exec ubuntu-openmpi.sif /path/to/your/parallel-mpi-app
```

The above lines can be entered at the command line of an interactive session, or can also be inserted into a batch job script.

Further Reading

- [Apptainer home page](#)
- [Apptainer Library](#)
- [Docker Hub](#)

1)

https://lmod.readthedocs.io/en/latest/010_user.html

2)

<https://easybuild.readthedocs.io/en/latest/>

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