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Preface

Welcome to the User Manual for NVIDIA Base Command Manager 10.

0.1 About This Manual

Name Changes From Version 9.2 To 10

The cluster manager software was originally developed by Bright Computing and the name “Bright” featured previously in the product, repositories, websites, and manuals.

Bright Computing was acquired by NVIDIA in 2022. The corresponding name changes, to be consistent with NVIDIA branding and products, are a work in progress. There is some catching up to do in places. For now, some parts of the manual still refer to Bright Computing and Bright Cluster Manager. These remnants will eventually disappear during updates.

BCM in particular is a convenient abbreviation that happens to have the same letters as the former Bright Cluster Manager. With the branding change in version 10, Base Command Manager is the official full name for the product formerly known as Bright Cluster Manager, and BCM is the official abbreviation for Base Command Manager.

This manual is intended for the end users of a cluster running NVIDIA Base Command Manager (BCM, also known as “the cluster manager”). This manual tends to see things from a user perspective, and covers the basics of using BCM’s user environment to run compute jobs on the cluster. Although it does cover some aspects of general Linux usage, it is by no means comprehensive in this area. Readers are expected to have some familiarity with the basics of a Linux environment from the regular user point of view.

Regularly updated production versions of the NVIDIA Base Command Manager 10 manuals are available on updated clusters by default at /cm/shared/docs/cm. The latest updates are always online at https://docs.nvidia.com/base-command-manager.

The manuals constantly evolve to keep up with the development of the BCM environment and the addition of new hardware and/or applications. The manuals also regularly incorporate feedback from administrators and users, and any comments, suggestions or corrections will be very gratefully accepted at manuals@brightcomputing.com.

There is also a feedback form available via Base View for administrators, via the Account icon, following the clickpath:

Account→Help→Feedback

0.2 Getting User-Level Support

A user is first expected to refer to this manual or other supplementary site documentation when dealing with an issue. If that is not enough to resolve the issue, then support for an end-user is typically provided by the cluster administrator, who is often a unix or Linux system administrator with some cluster experience. Commonly, the administrator has configured and tested the cluster beforehand, and therefore has a good idea of its behavior and quirks. The initial step when calling in outside help is thus often to call in the cluster administrator.
1

Introduction

This manual is intended for cluster users who need a quick introduction to the NVIDIA Base Command Manager, which manages a Beowulf cluster configuration. It explains how to use the MPI and batch environments, how to submit jobs to the queuing system, and how to check job progress. The specific combination of hardware and software installed may differ depending on the specification of the cluster, which means that parts of this manual may not be relevant to the user’s particular cluster.

1.1 What Is A Beowulf Cluster?

1.1.1 Background And History

In the history of the English language, Beowulf is the earliest surviving epic poem written in English. It is a story about a hero with the strength of many men who defeated a fearsome monster called Grendel.

In computing, a Beowulf class cluster computer is a multiprocessor architecture used for parallel computations, i.e., it uses many processors together so that it has the brute force to defeat certain “fearsome” number-crunching problems.

The architecture was first popularized in the Linux community when the source code used for the original Beowulf cluster built at NASA was made widely available. The Beowulf class cluster computer design usually consists of one head node and one or more regular nodes connected together via Ethernet or some other type of network. While the original Beowulf software and hardware has long been superseded, the name given to this basic design remains “Beowulf class cluster computer”, or less formally “Beowulf cluster”.

1.1.2 Brief Hardware And Software Description

On the hardware side, commodity hardware is generally used in Beowulf clusters to keep costs down. These components are usually x86-compatible processors produced at the Intel and AMD chip foundries, standard Ethernet adapters, InfiniBand interconnects, and switches. From around 2019 the ARMv8 processor architecture is also gaining attention.

On the software side, free and open-source software is generally used in Beowulf clusters to keep costs down. For example: the Linux operating system, the GNU C compiler collection and open-source implementations of the Message Passing Interface (MPI) standard.

The head node controls the whole cluster and serves files and information to the nodes. It is also the cluster’s console and gateway to the outside world. Large Beowulf clusters might have more than one head node, and possibly other nodes dedicated to particular tasks, for example consoles or monitoring stations. In most cases compute nodes in a Beowulf system are dumb—in general, the dumber the better—with the focus on the processing capability of the node within the cluster, rather than other abilities a computer might generally have. A node may therefore have

- one or more processing elements. The processors may be standard CPUs, as well as GPUs, FPGAs, MICs, and so on.
• enough local memory—memory contained in a single node—to deal with the processes passed on to the node

• a connection to the rest of the cluster

Nodes are configured and controlled by the head node, and do only what they are told to do. One of the main differences between Beowulf and a Cluster of Workstations (COW) is the fact that Beowulf behaves more like a single machine rather than many workstations. In most cases, the nodes do not have keyboards or monitors, and are accessed only via remote login or possibly serial terminal. Beowulf nodes can be thought of as a CPU + memory package which can be plugged into the cluster, just like a CPU or memory module can be plugged into a motherboard to form a larger and more powerful machine. A significant difference is that the nodes of a cluster have a relatively slower interconnect.

1.2 Brief Network Description

A Beowulf Cluster consists of a login, compile and job submission node, called the head, and one or more compute nodes, often referred to as worker nodes. A second (fail-over) head node may be present in order to take control of the cluster in case the main head node fails. Furthermore, a second fast network may also have been installed for high-performance low-latency communication between the (head and the) nodes (see figure 1.1).

The login node is used to compile software, to submit a parallel or batch program to a job queuing system and to gather/analyze results. Therefore, it should rarely be necessary for a user to log on to one of the nodes and in some cases node logins are disabled altogether. The head, login and compute nodes usually communicate with each other through a gigabit Ethernet network, capable of transmitting information at a maximum rate of 1000 Mbps. In some clusters 10 gigabit Ethernet (10GE, 10GBE, or 10GigE) is used, capable of up to 10 Gbps rates, while higher rates than that are also available.

Sometimes an additional network is used by the cluster for even faster communication between the compute nodes. This particular network is mainly used for programs dedicated to solving large scale computational problems, which may require multiple machines and could involve the exchange of vast amounts of information. One such network topology is InfiniBand, commonly capable of transmitting information at a maximum effective data rate of about 124Gbps and about 1.2µs end-to-end latency on small packets, for clusters in 2013. The commonly available maximum transmission rates will increase over the years as the technology advances.
Applications relying on message passing benefit greatly from lower latency. The fast network is usually complementary to a slower Ethernet-based network.
2.1 Login To The Cluster Environment

The login node is the node where the user logs in and works from. Simple clusters have a single login node, but large clusters sometimes have multiple login nodes to improve the availability of the cluster. In most clusters, the login node is also the head node from where the cluster is monitored and installed. On the login node:

- applications can be developed
- code can be compiled and debugged
- applications can be submitted to the cluster for execution
- running applications can be monitored

To carry out an ssh login to the cluster, a terminal session can be started from Unix-like operating systems:

Example

$ ssh myname@cluster.hostname

On a Windows operating system, an SSH client such as for PuTTY (http://www.putty.org) can be downloaded. Another standard possibility is to run a Unix-like environment such as Cygwin (http://www.cygwin.com) within the Windows operating system, and then run the SSH client from within it.

A Mac OS X user can use the Terminal application from the Finder, or under Application/Utilities/Terminal.app. An X11 windowing environment must be installed for it to work. XQuartz is the recommended X11 windowing environment, and is indeed the only Apple-backed X11 version available from OS X 10.8 onward.

When using the SSH connection, the cluster’s address must be added. When the connection is made, a username and password must be entered at the prompt.

If the administrator has changed the default SSH port from 22 to something else, the port can be specified with the -p <port> option:

$ ssh -X -p <port> <user>@<cluster>

The -X option can be dropped if no X11-forwarding is required. X11-forwarding allows a GUI application from the cluster to be displayed locally.

Optionally, after logging in, the password used can be changed using the passwd command:

$ passwd
2.2 Setting Up The User Environment

By default, each user uses the bash shell interpreter. In that case, each time a user login takes place, a file named .bashrc is executed to set up the shell environment for the user. The shell and its environment can be customized to suit user preferences. For example,

- the prompt can be changed to indicate the current username, host, and directory, for example: by setting the prompt string variable:
  \[
  \text{PS1}=\"[\text{\u@h \h} \W]\$\"\]
- the size of the command history file can be increased, for example: \text{export HISTSIZE=100}
- aliases can be added for frequently used command sequences, for example: \text{alias lart='ls -alrt'}
- environment variables can be created or modified, for example: \text{export MYVAR=MYSTRING}
- the location of software packages and versions that are to be used by a user (the path to a package) can be set.

Because there is a huge choice of software packages and versions, it can be hard to set up the right environment variables and paths for software that is to be used. Collisions between different versions of the same package and non-matching dependencies on other packages must also be avoided. To make setting up the environment easier, BCM provides preconfigured environment modules (section 2.3).

2.3 Environment Modules

For a user to compile and run computational jobs on a cluster, a special shell environment is typically set up for the software that is used.

However, setting up the right environment for a particular software package and version can be tricky, and it can be hard to keep track of how it was set up.

For example, users want to have a clean way to bring up the right environment for compiling code according to the various MPI implementations, but can easily get confused about which libraries have been used, and can end up with multiple libraries with similar names installed in a disorganized manner.

A user might also like to conveniently test new versions of a software package before permanently installing the package.

Within a Linux distribution running without special utilities, setting up environments can be complex. However, BCM makes use of the environment modules package, which provides the module command. The module command is a special utility to make taking care of the shell environment much easier.

2.3.1 Available Commands

Practical use of the modules commands is given in sections 2.3.2 and 2.3.3.

For reference, the help text for the module command can be viewed as follows:

Example

```
[me@cluster ~]$ module --help
Module Version: Release 4.4.0 (2019-11-17)
Usage: module [options] [command] [args ...]
Loading / Unloading commands:
  add | load modulefile [...] Load modulefile(s)
  rm | unload modulefile [...] Remove modulefile(s)
```
purge
reload | refresh
switch | swap [mod1] mod2

List / Searching commands:
list [-t|-l] List loaded modules
avail [-d|-l] [-t|-l] [-S|-C] [--in-depth|--no-in-depth] [mod ...]
list all or matching available modules
aliases List all module aliases
whatis [modulefile ...] Print whatis information of modulefile(s)
apropos | keyword | search str Search all name and whatis containing str
is-loaded [modulefile ...] Test if any of the modulefile(s) are loaded
is-avail modulefile [...] Is any of the modulefile(s) available
info-loaded modulefile Get full name of matching loaded module(s)

Collection of modules handling commands:
save [collection|file] Save current module list to collection
restore [collection|file] Restore module list from collection or file
saverm [collection] Remove saved collection
saveshow [collection|file] Display information about collection
savelist [-t|-l] List all saved collections
is-saved [collection ...] Test if any of the collection(s) exists

Shell's initialization files handling commands:
initlist List all modules loaded from init file
initadd modulefile [...] Add modulefile to shell init file
initrm modulefile [...] Remove modulefile from shell init file
initswitch mod1 mod2 Switch mod1 with mod2 from init file
initclear Clear all modulefiles from init file

Environment direct handling commands:
prepend-path [-d c] var val [...] Prepend value to environment variable
append-path [-d c] var val [...] Append value to environment variable
remove-path [-d c] var val [...] Remove value from environment variable

Other commands:
help [modulefile ...] Print this or modulefile(s) help info
display | show modulefile [...] Display information about modulefile(s)
test [modulefile ...] Test modulefile(s)
use [-a|-p] dir [...] Add dir(s) to MODULEPATH variable
unuse dir [...] Remove dir(s) from MODULEPATH variable
is-used [dir ...] Is any of the dir(s) enabled in MODULEPATH
path modulefile Print modulefile path
paths modulefile Print path of matching available modules
clear [-f] Reset Modules-specific runtime information
source scriptfile [...] Execute scriptfile(s)
config [--dump-state|name [val]] Display or set Modules configuration

Switches:
-t | --terse Display output in terse format
-l | --long Display output in long format
-d | --default Only show default versions available
-L | --latest Only show latest versions available
-S | --starts-with
Search modules whose name begins with query string
-C  --contains Search modules whose name contains query string
-i  --icase Case insensitive match
-a  --append Append directory to MODULEPATH
-p  --prepend Prepend directory to MODULEPATH
--auto Enable automated module handling mode
--no-auto Disable automated module handling mode
-f  --force By-pass dependency consistency or confirmation dialog

Options:
-h  --help This usage info
-V  --version Module version
-D  --debug Enable debug messages
-v  --verbose Enable verbose messages
-s  --silent Turn off error, warning and informational messages
--paginate Pipe mesg output into a pager if stream attached to terminal
--no-pager Do not pipe message output into a pager
--color[=WHEN] Colorize the output; WHEN can be 'always' (default if omitted), 'auto' or 'never'

2.3.2 Managing Environment Modules As A User
There is a good chance the cluster administrator has set up the user’s account, fred for example, so that some modules are loaded already by default. In that case, the modules loaded into the user’s environment can be seen with the module list command:

Example

[fred@basecm10 ~]$ module list
Currently Loaded Modulefiles:
1) shared 2) cmsh 3) cmd 4) cluster-tools/10.0 5) cm-setup/10.0 6) gcc/9.2.0

If there are no modules loaded by default, then the module list command just returns nothing.

How does a user know what modules are available? The “module avail” command lists all modules that are available for loading (some output elided):

Example

[fred@basecm10 ~]$ module avail
------------------------------------ /cm/local/modulefiles ------------------------------------
cluster-tools/10.0   cmd   freeipmi/1.6.4   luajit   openldap
               cm/image/10.0  cmjob   gcc/9.2.0  module-git   python3
               cm-scale/cm-scale.module  cmsh   ipmitool/1.8.18   module-info   python37
               cm-set-up/10.0   dot   lua/5.3.5   null   shared
----------------------------------- /cm/shared/modulefiles ------------------------------------
blacs/openmpi/gcc/64/1.1patch00  intel/compiler/32/2019/19.0.5
blas/gcc/64/3.8.0  intel/compiler/64/(default)
bonnie++/1.98  intel/compiler/64/2019/19.0.5
cm-pmix3/3.1.4  intel/mpi/64/(default)
default-environment  intel/mpi/64/2019/5.281
fft2/openmpi/gcc/64/double/2.1.5  iozone/3.487
fft2/openmpi/gcc/64/float/2.1.5  lapack/gcc/64/3.8.0
fft3/openmpi/gcc/64/3.3.8  mpi/ge/gcc/64/3.3.2
globals/openmpi/gcc/64/5.7  mlapich2/gcc/64/2.3.2
hdf5/1.10.1  metcdf/gcc/64/gcc/64/4.7.3

2.3 Environment Modules

In the list there are two kinds of modules:

- **local modules**, which are specific to the node, or head node only
- **shared modules**, which are made available from a shared storage, and which only become available for loading after the shared module is loaded.

Modules can be loaded using the `add` or `load` options. A list of modules can be added by spacing them:

Example

```
[fred@basecm10 ~]$ module add shared gcc openmpi/gcc
```

Tab completion works for suggesting modules for the `add`/`load` commands. If the tab completion suggestion is unique, even though it is not the full path, then it is still enough to specify the module. For example, looking at the possible available modules listed by the `avail` command previously, it turns out that specifying `gcc` is enough to specify `gcc/9.2.0` because there is no other directory path under `gcc/` besides `9.2.0` anyway.

To remove one or more modules, the `module unload` or `module rm` command is used.

To remove all modules from the user’s environment, the `module purge` command is used.

The user should be aware that some loaded modules can conflict with others loaded at the same time. This can happen with MPI modules. For example, loading `openmpi/gcc` without removing an already loaded `intel/mpi/64` can result in conflicts about which compiler should be used.

**The shared Module**

The shared module provides access to shared libraries. By default these are under `/cm/shared`.

The shared module is special because often other modules, as seen under `/cm/shared/modulefiles`, depend on it. So, if it is to be loaded, then it is usually loaded first, so that the dependent modules can use it.

The shared module is obviously a useful local module, and is therefore often configured to be loaded for the user by default. Setting the default environment modules is discussed in section 2.3.3.

### 2.3.3 Changing The Default Environment Modules

If a user has to manually load up the same modules every time upon login it would be inefficient. That is why an initial default state for modules can be set up by the user, by using the `module init*` subcommands:

- **module initadd**: add a module to the initial state
- **module initrm**: remove a module from the initial state
- **module initlist**: list all modules loaded initially
- **module initclear**: clear all modules from the list of modules loaded initially

**Example**
[fred@basecm10 ~]$ module initclear
[fred@basecm10 ~]$ module initlist
bash initialization file $HOME/.bashrc loads modules:

[fred@basecm10 ~]$ module initadd shared gcc openmpi/gcc
[fred@basecm10 ~]$ module initlist
bash initialization file $HOME/.bashrc loads modules:
  shared gcc openmpi/gcc

In the preceding example, the modules defined for the new initial environment for the user are
loaded from the next login onward.

Example

[fred@basecm10 ~]$ module list
No Modulefiles Currently Loaded.
[fred@basecm10 ~]$ exit
logout
Connection to basecm10 closed
[root@basejumper ~]# ssh fred@basecm10
fred@basecm10's password:
...
[fred@basecm10 ~]$ module list
Currently Loaded Modulefiles:
  1) shared  2) gcc/9.2.0  3) openmpi/gcc/64/1.10.7
[fred@basecm10 ~]$

If the user is unsure about what the module does, it can be checked using “module whatis”:

$ module whatis openmpi/gcc
----------------------------------- /cm/shared/modulefiles ------------------------------------
openmpi/gcc/64/1.10.7: adds OpenMPI to your environment variables

The man pages for module and modulefile give further details on usage.

2.4 Compiling Applications

Compiling an application is usually done on the head node or login node. Typically, there are several
compilers available on the head node in BCM. These compilers provide different levels of optimization,
standards conformance, and support for accelerators.

For example: The GNU compiler collection, Intel compilers, and the NVIDIA HPC SDK compilers.
The following table summarizes the available compiler commands on a cluster with these compilers:

<table>
<thead>
<tr>
<th>Language</th>
<th>GNU</th>
<th>NVIDIA HPC</th>
<th>Intel</th>
</tr>
</thead>
<tbody>
<tr>
<td>C</td>
<td>gcc</td>
<td>nvc and nvcc</td>
<td>icc</td>
</tr>
<tr>
<td>C++</td>
<td>g++</td>
<td>nvc++ and nvcc</td>
<td>icc</td>
</tr>
<tr>
<td>Fortran77</td>
<td>gfortran</td>
<td>nvfortran</td>
<td>ifort</td>
</tr>
<tr>
<td>Fortran90</td>
<td>gfortran</td>
<td>nvfortran</td>
<td>ifort</td>
</tr>
<tr>
<td>Fortran95</td>
<td>gfortran</td>
<td>nvfortran</td>
<td>ifort</td>
</tr>
</tbody>
</table>

GNU compilers are the de facto standard on Linux and are installed by default. They are provided
under the terms of the GNU General Public License. Commercial compilers by Portland and Intel are
available as packages via the BCM YUM repository, and require the purchase of a license to use them. To
make a compiler available to be used in a user’s shell commands, the appropriate environment module
(section 2.3) must be loaded first. On most clusters two versions of GCC are available:
2.4 Compiling Applications

1. The version of GCC that comes along with the Linux distribution. For example, for CentOS 7.x:

   Example

   ```
   [fred@basecm10 ~]$ which gcc; gcc --version | head -1
   /usr/bin/gcc
   gcc (GCC) 4.8.5 20150623 (Red Hat 4.8.5-39)
   ```

2. The latest version suitable for general use that is packaged as a module by BCM:

   Example

   ```
   [fred@basecm10 ~]$ module load gcc
   [fred@basecm10 ~]$ which gcc; gcc --version | head -1
   /cm/local/apps/gcc/9.2.0/bin/gcc
   gcc (GCC) 9.2.0
   ```

   To use the latest version of GCC, the `gcc` module in a default BCM cluster must be loaded. To revert to the version of GCC that comes natively with the Linux distribution, the `gcc` module must be unloaded.

   The compilers—GCC, Intel, Portland—in the preceding table are ordinarily used for applications that run on a single node. However, the applications used may fork, thread, and run across as many nodes and processors as they can access if the application is designed that way.

   The standard, structured way of running applications in parallel is to use the MPI-based libraries (Chapter 3). These are the MPICH, MVAPIC, Open MPI, and Intel MPI libraries. The libraries link to the underlying compilers of the preceding table.

   If the parallel library environment has been loaded, then the following MPI compiler commands automatically use the underlying compilers:

<table>
<thead>
<tr>
<th>Language</th>
<th>C</th>
<th>C++</th>
<th>Fortran77</th>
<th>Fortran90</th>
</tr>
</thead>
<tbody>
<tr>
<td>Command</td>
<td>mpicc</td>
<td>mpicxx</td>
<td>mpif77</td>
<td>mpif90</td>
</tr>
</tbody>
</table>

2.4.1 Open MPI And Mixing Compilers

BCM comes with multiple Open MPI packages corresponding to the different available compilers. However, sometimes mixing compilers is desirable. For example, C-compilation may be preferred using `icc` from Intel, while Fortran90-compilation may be preferred using `gfortran` from the GNU Project. In such cases it is possible to override the default compiler path environment variable, for example:

   ```
   [fred@basecm10 ~]$ module list
   Currently Loaded Modulefiles:
   1) shared  2) gcc/9.2.0  3) openmpi/gcc/64/1.10.7
   [fred@basecm10 ~]$ mpicc --version --showme; mpif90 --version --showme
   gcc --version
   gfortran --version
   [fred@basecm10 ~]$ export OMPI_CC=icc; export OMPI_FC=openf90
   [fred@basecm10 ~]$ mpicc --version --showme; mpif90 --version --showme
   icc --version
   openf90 --version
   ```

   Variables that may be set are OMPI_CC, OMPI_CXX, OMPI_FC, and OMPI_F77. More on overriding the Open MPI wrapper settings is documented in the man pages of `mpicc` in the environment section.
3

Using MPI

3.1 Introduction

The Message Passing Interface (MPI) is a standardized and portable message passing system designed by a group of researchers from academia and industry to function on a wide variety of parallel computers. The standard defines the syntax and semantics of a core of library routines useful to a wide range of users writing portable message-passing programs in Fortran or the C programming language. MPI libraries allow the compilation of code so that it can be used over a variety of multi-processor systems from SMP nodes to NUMA (non-Uniform Memory Access) systems and interconnected cluster nodes.

Depending on the cluster hardware, the interconnect available may be Ethernet, Infini-Band/OmniPath.

Typically, the cluster administrator has already custom-configured the cluster in a way to suit the workflow of users. However, users that are interested in understanding and using the MPI options available, should find this chapter helpful.

3.2 MPI Libraries

MPI libraries that are commonly integrated by the cluster administrator with NVIDIA Base Command Manager are

- MPICH (https://www.mpich.org/)
- MVAPICH (http://mvapich.cse.ohio-state.edu/)
- OpenMPI (https://www.open-mpi.org/)

The preceding MPI libraries can be used with compilers from GCC, or Intel.

The following MPI library implementations may also be integrated with BCM:

- Open MPI version 4 library with CUDA-awareness. NVIDIA GPUs using CUDA can make use of this library.

3.3 MPI Packages And Module Paths

By default, the only MPI implementations installed are the ones that work using GCC.

The cluster administrator can make available for the user a variety of different MPI implementations for different compilers, by installing the appropriate packages. The actual combination of compiler and MPI implementation that a user needs for job runs can then be loaded by the user with the help of modules (section 2.3.2).
### 3.3.1 MPI Packages That Can Be Installed, And Their Corresponding Module Paths

The following packages are made available from the BCM repositories at the time of writing (October 2022):

**Argonne National Laboratory Ethernet implementation for MPI-1, MPI-2, MPI-3**

<table>
<thead>
<tr>
<th>package name</th>
<th>module path</th>
</tr>
</thead>
<tbody>
<tr>
<td>mpich-ge-gcc-64</td>
<td>mpich/ge/gcc</td>
</tr>
<tr>
<td>mpich-ge-intel-64</td>
<td>mpich/ge/intel</td>
</tr>
</tbody>
</table>

**MVAPICH2 InfiniBand implementation for MPI-3**

<table>
<thead>
<tr>
<th>package name</th>
<th>module path</th>
</tr>
</thead>
<tbody>
<tr>
<td>mvapich2-gcc-64</td>
<td>mvapich2/gcc</td>
</tr>
<tr>
<td>mvapich2-intel-64</td>
<td>mvapich2/intel</td>
</tr>
</tbody>
</table>

**MVAPICH2 InfiniBand implementation with performance scaled messaging for MPI-3**

<table>
<thead>
<tr>
<th>package name</th>
<th>module path</th>
</tr>
</thead>
<tbody>
<tr>
<td>mvapich2-psmgcc-64</td>
<td>mvapich2/psmgcc</td>
</tr>
<tr>
<td>mvapich2-psmintel-64</td>
<td>mvapich2/psmintel</td>
</tr>
</tbody>
</table>

**Open MPI version 1 implementation for Ethernet with MPI-3**

<table>
<thead>
<tr>
<th>package name</th>
<th>module path</th>
</tr>
</thead>
<tbody>
<tr>
<td>openmpi-ge-gcc-64</td>
<td>openmpi/gcc</td>
</tr>
<tr>
<td>openmpi-ge-intel-64</td>
<td>openmpi/intel</td>
</tr>
</tbody>
</table>

**Open MPI version 1 implementation for Ethernet and InfiniBand with MPI-3**

<table>
<thead>
<tr>
<th>package name</th>
<th>module path</th>
</tr>
</thead>
<tbody>
<tr>
<td>openmpi-geib-gcc-64</td>
<td>openmpi/gcc</td>
</tr>
<tr>
<td>openmpi-geib-intel-64</td>
<td>openmpi/intel</td>
</tr>
</tbody>
</table>

**Open MPI version 3 implementation for Ethernet and InfiniBand with MPI-3**

<table>
<thead>
<tr>
<th>package name</th>
<th>module path</th>
</tr>
</thead>
<tbody>
<tr>
<td>openmpi3-geib-gcc-64</td>
<td>openmpi3/gcc</td>
</tr>
<tr>
<td>openmpi3-ge-gcc-64 (only Ethernet, only GCC)</td>
<td>openmpi3/gcc</td>
</tr>
<tr>
<td>openmpi3-ge-intel-64 (only Ethernet, only Intel compiler)</td>
<td>openmpi3/intel</td>
</tr>
</tbody>
</table>

**Open MPI version 4 implementation for InfiniBand with MPI-3**

<table>
<thead>
<tr>
<th>package name</th>
<th>module path</th>
</tr>
</thead>
<tbody>
<tr>
<td>openmpi4-gcc (GCC)</td>
<td>openmpi4/gcc</td>
</tr>
<tr>
<td>openmpi4-intel (Intel compiler)</td>
<td>openmpi4/intel</td>
</tr>
</tbody>
</table>
3.3 MPI Packages And Module Paths

3.3.2 Finding The Installed MPI Packages And Their Available Module Paths

Finding The Installed MPI Packages

The packages installed on the cluster can be found with the `rpm -qa` query on a RHEL system.

Example

```
[fred@basecm10 ~]$ # search for packages starting with (open)mpi, mvapich, intel-mpi, cm-openmpi
[fred@basecm10 ~]$ rpm -qa | egrep '(^mpi|^openmpi|^mvapich|^intel-mpi|cm-openmpi)'
openmpi-geib-gcc-64-1.10.7-656_cm9.2.x86_64
mpich-ge-gcc-64-3.4.2-197_cm9.2.x86_64
mvapich2-gcc-64-2.3.7-213_cm9.2.x86_64
openmpi4-gcc-4.1.2-100042_cm9.2_d17293b429.x86_64
intel-mpi-2020-2019.9-100008_cm9.2_aca4120975.x86_64
cm-openmpi4-cuda11.7-ofed47-gcc9-4.1.4-100012_cm9.2_a9f8de3740.x86_64
openmpi4-intel-4.1.2-100042_cm9.2_d17293b429.x86_64
```

The interconnect and compiler implementation of a package can be worked out from looking at the name of the package.

Here, for example,

```
openmpi-geib-gcc-64-1.10.7-656_cm9.2.x86_64
```
implies: Open MPI version 1.10.7, compiled for both Gigabit Ethernet (ge) and InfiniBand (ib), with GCC (gcc, the GNU project cross-compiler) for a 64-bit architecture, packaged as a (BCM) cluster manager (cm) package, for version 10.0 for the x86_64 architecture.

Finding The Available MPI Modules

The corresponding module paths can be found by a search through the available modules:

Example

```
[fred@basecm10 ~]$ # search for modules starting with (open)mpi or mvapich
[fred@basecm10 ~]$ module -l avail | egrep ('^openmpi|mpi|mvapich|intel/mpi|cm-openmpi)'
intel/mpi/64/ default
intel/mpi64/2020/4.304 2021/10/23 22:18:12
mpich/ge/gcc64/3.4.2 2021/11/26 22:33:47
mvapich2/gcc/64/2.3.7 2022/07/14 13:02:23
```
Tab-completion when searching for modules is another approach.

As in the case for the MPI library, for a module the interconnect and compiler implementation can likewise be worked out from looking at the name of the module. So, if a user would like to use an Open MPI implementation that works with an AMD64 node, using GCC, then loading the module `openmpi/gcc/64` should be suitable.

### 3.4 The Appropriate Interconnect, Compiler, And MPI Implementation For A Module

#### 3.4.1 Interconnects

Jobs can use particular networks for inter-node communication. The hardware for these networks can be Ethernet or InfiniBand, while the software can be a particular MPI implementation.

**Gigabit Ethernet**

Gigabit Ethernet is an interconnect that is commonly available in consumer computing. For HPC the next generation 10 Gigabit Ethernet and beyond have also been in use for some time. For Ethernet, no additional modules or libraries are needed. The Open MPI, MPICH and MVAPICH implementations all work with any Ethernet technology.

**InfiniBand**

InfiniBand is a high-performance switched fabric that has come to dominate the connectivity in HPC applications. An InfiniBand interconnect has a lower latency and somewhat higher throughput than a comparably-priced Ethernet interconnect. This is because of special hardware, as well as special software that shortcuts the networking stack in the operating system layer. This typically provides significantly greater performance for most HPC applications. Open MPI and MVAPICH are suitable MPI implementations for InfiniBand.

#### 3.4.2 Selecting A Compiler And MPI implementation

Once the appropriate compiler module has been loaded, the associated MPI implementation is selected by loading the corresponding library module. In the following simplified list, `<compiler>` indicates a choice of `gcc`, or `intel`:

- `mpich/ge/<compiler>`
- `mvapich2/<compiler>`
- `openmpi/<compiler>`
- `openmpi3/<compiler>`
- `openmpi4/<compiler>`

Section 3.3.1 should be referred to for a more complete list.

### 3.5 Compiling And Carrying Out An MPI Run

After the appropriate MPI module has been added to the user environment, the user can start compiling applications.
For a cluster using Ethernet interconnectivity, the `mpich` and `openmpi` implementations may be used. For a cluster using InfiniBand, the `mvapich`, `mvapich2` and `openmpi` implementations may be used. Open MPI’s `openmpi` implementations first attempt to use InfiniBand, but revert to Ethernet if InfiniBand is not available.

In this section the loading of modules, compilation, and runs are illustrated.

### 3.5.1 Example MPI Run

This example covers an MPI run, which can be run inside and outside of a queuing system.

Depending on the libraries and compilers installed on the system by the cluster administrator, the availability of the packages providing these modules may differ. To see a full list of modules on the system the command `module avail` can be typed.

#### Examples Of Loading MPI Modules

To use `mpirun`, the relevant environment modules must be loaded. For example, to use the `mpich` over Gigabit Ethernet (`ge`) GCC implementation:

```bash
$ module add mpich/ge/gcc
```

or to use the `openmpi4` Open MPI version 4 MPI-3 Intel implementation:

```bash
$ module add openmpi4/intel
```

Similarly, to use the `mvapich2` InfiniBand GCC implementation:

```bash
$ module add mvapich2/gcc
```

#### Keeping Loading Of Modules Clean

The preceding modules can actually be loaded concurrently, and it works as expected. Paths supplied by the most recently-loaded module override the paths of any previous modules.

For example, if `mpich/ge/gcc` is loaded first, then `openmpi4/intel`, and then `mvapich2/gcc`, as suggested in the preceding excerpts, then the modules might be listed as:

```bash
$ module list
Currently Loaded Modulefiles:
Currently Loaded Modulefiles:
  1) shared  3) cmd  5) cm-setup/9.2  7) openmpi4/intel/4.1.2
  2) cmsh  4) cluster-tools/9.2  6) mpich/ge/gcc/64/3.4.2  8) mvapich2/gcc/64/2.3.7
```

The path of the MPI compiler `mpicc` is defined by the last module in the modulefiles stack, which is the MVAPICH GCC implementation, as can be seen with:

```bash
$ which mpicc
/cm/shared/apps/mvapich2/gcc/64/2.3.7/bin/mpicc
```

After removing it, then the path changes to the path supplied by the previous module in the stack, the Open MPI Intel implementation:

```bash
$ module remove mvapich2/gcc/64/2.3.7
$ which mpicc
/cm/shared/apps/openmpi4/intel/4.1.2/bin/mpicc
```

After removing that module too, the path again changes to the path supplied by the previous module in the stack, the MPICH GCC implementation:

```bash
$ module remove openmpi4/intel/4.1.2
$ which mpicc
/cm/shared/apps/mpich/ge/gcc/64/3.4.2/bin/mpicc
```

However, because loading modules on top of each other can cause confusion, a user should generally try adding modules in a simple, clean manner.
Examples Compiling And Preparing The MPI Application

The code must be compiled with MPI libraries and an underlying compiler. The correct library command can be found in the following table:

<table>
<thead>
<tr>
<th>Language</th>
<th>C</th>
<th>C++</th>
<th>Fortran77</th>
<th>Fortran90</th>
</tr>
</thead>
<tbody>
<tr>
<td>Command</td>
<td>mpicc</td>
<td>mpixx</td>
<td>mpif77</td>
<td>mpif90</td>
</tr>
</tbody>
</table>

An MPI application `myapp.c`, built in C, could then be compiled as:

```
$ mpicc myapp.c
```

The `mpicc` compilation requires the underlying compiler (GCC, Intel) already be available. By default, Linux systems have a version of the GCC compiler already in their paths, even if no modules have been loaded. So if the module for an MPI implementation based on GCC is loaded without explicitly loading the GCC compiler, then the `mpicc` compilation still works in this case.

The `a.out` binary that is created can then be executed using the `mpirun` command (section 3.5.1).

Creating A Machine File

A machine file contains a list of nodes which can be used by MPI programs.

The workload management system creates a machine file based on the nodes allocated for a job when the job is submitted with the workload manager job submission tool. So if the user chooses to have the workload management system allocate nodes for the job, then creating a machine file is not needed.

However, if an MPI application is being run “by hand” outside the workload manager, then the user is responsible for creating a machine file manually. Depending on the MPI implementation, the layout of this file may differ.

Machine files can generally be created in two ways:

- Listing the same node several times to indicate that more than one process should be started on each node:

  node001
  node001
  node002
  node002

- Listing nodes once, but with a suffix for the number of CPU cores to use on each node:

  node001:2
  node002:2

Running The Application

Creating A Simple Parallel Processing Executable

A simple “hello world” program designed for parallel processing can be built with MPI. After compiling it, it can be used to send a message about how and where it is running:

```
[fred@basecm10 ~]$ cat hello.c
#include <stdio.h>
#include <mpi.h>

int main (int argc, char *argv[])
{
  int id, np, i;
  char processor_name[MPI_MAX_PROCESSOR_NAME];
  int processor_name_len;
```
MPI_Init(&argc, &argv);

MPI_Comm_size(MPI_COMM_WORLD, &np);
MPI_Comm_rank(MPI_COMM_WORLD, &id);
MPI_Get_processor_name(processor_name, &processor_name_len);

for(i=1;i<2;i++)
{printf("Hello world from process %03d out of %03d, processor name %s\\n",
    id, np, processor_name);
}

MPI_Finalize();
return 0;

[fred@basecm10 ~]$ module add openmpi/gcc  #or as appropriate depending on installed package
[fred@basecm10 ~]$ mpicc hello.c -o hello

The preceding compilation works for MPI based on gcc because a version of gcc is already available by default with the distribution. Its version can be found with, for example:

[fred@basecm10 ~]$ gcc --version | head -1
gcc (GCC) 8.5.0 20210514 (Red Hat 8.5.0-10)

If the user has a need for another version of the gcc compiler, then that can be loaded before the compilation.

[fred@basecm10 ~]$ module load gcc/11.2.0
[fred@basecm10 ~]$ mpicc hello.c -o hello

If the user would like to use the Intel compiler with MPI instead, then the steps would instead be something like:

[fred@basecm10 ~]$ module load intel/compiler/64/2020/19.1.3 intel/mpi/64/2020/4.304
[fred@basecm10 ~]$ mpicc hello.c -o helloforintel

If in doubt about how things are compiled, then the libraries that the binary was compiled with can be checked using ldd. For example, the GCC-related libraries used could be checked with:

[fred@basecm10 ~]$ ldd hello |grep gcc |awk '{print $1,$2,$3}'
libmpi.so.12 => /cm/shared/apps/openmpi/gcc/64/1.10.7/lib64/libmpi.so.12
libopen-rte.so.12 => /cm/shared/apps/openmpi/gcc/64/1.10.7/lib64/libopen-rte.so.12
libopen-pal.so.13 => /cm/shared/apps/openmpi/gcc/64/1.10.7/lib64/libopen-pal.so.13

The compiled binary can be run:

[fred@basecm10 ~]$ ./hello
Hello world from process 000 out of 001, processor name basecm10

However, it still runs on a single processor, and on the node it is started on, unless it is submitted to the system in a special way.

Running An MPI Executable In Parallel Without A Workload Manager

Compute node environment provided by user's .bashrc: After the relevant module files are chosen (section 3.5.1) for MPI, an executable compiled with MPI libraries runs on nodes in parallel when submitted with mpirun. The executable running on other nodes loads environment modules on those other nodes by sourcing the .bashrc file of the user (section 2.3.3) and flags passed by the mpirun command. It is a good idea to ensure that the environment module stack used on the compute node is not confusing.
The environment of the user from the interactive shell prompt is not normally carried over automatically to the compute nodes during an `mpirun` submission. That is, compiling and running the executable normally only works on the local node without a special treatment. To have the executable run on the compute nodes, the right environment modules for the job must be made available on the compute nodes too, as part of the user login process to the compute nodes for that job. Usually the system administrator takes care of such matters in the default user configuration by setting up a default user environment (section 2.3.3), with reasonable `init.rm` and `init.add` options. Users are then typically allowed to set up their personal default overrides to the default administrator settings, by placing their own `init.rm` and `init.add` options to the `module` command according to their needs, or by specifying an overriding environment in the job submissions.

**Running `mpirun` outside a workload manager:** When using `mpirun` manually, outside a workload manager environment, the number of processes (-np) as well as the number of hosts (-machinefile) should be specified. For example, on a cluster with 2 compute-nodes and a machine file as specified on page 18:

**Example**

```bash
$ module add mvapich2/gcc gcc/11.2.0 # or as appropriate
$ mpicc hello.c -o hello
$ mpirun -np 4 -machinefile mpirun.hosts ~/hello
```

```
Hello world from process 0 of 4 on node001.cm.cluster
Hello world from process 1 of 4 on node002.cm.cluster
Hello world from process 2 of 4 on node001.cm.cluster
Hello world from process 3 of 4 on node002.cm.cluster
```

If the cluster has no InfiniBand connectors, then the preceding `mpirun` fails, because MVAPICH requires InfiniBand. That kind of failure displays an output such as:

```
rdma_open_hca(575).............: No IB device found
```

Open MPI implementations are more forgiving. They check for InfiniBand and if that is unavailable they go ahead with Ethernet. However they have their own quirks. For example, an attempt to carry out an MPI run with Open MPI might be as follows:

**Example**

```bash
$ module initclear; module initadd openmpi/gcc
$ module add openmpi/gcc # or as appropriate
$ mpicc hello.c -o hello
$ mpirun -np 4 -machinefile mpirun.hosts hello
```

```
bash: orted: command not found
```

```
ORTE was unable to reliably start one or more daemons.
This usually is caused by:
```

```
... output snipped
```

It is generally a good idea to read through the man page for `mpirun` for the MPI implementation that the user is using. In this case, the man page for `mpirun` for Open MPI reveals that here the problem is that the environment in the interactive shell is not carried over to the compute nodes during an `mpirun` submission. So the path for `mpirun` should be specified with with either the `-prefix` option, or as an absolute path:

**Example**

```bash
```
3.5 Compiling And Carrying Out An MPI Run

The output of the preceding hello.c program is actually printed in random order. This can be modified as follows, so that only process 0 prints to the standard output, and other processes communicate their output to process 0:

```c
#include "mpi.h"
#include "string.h"
#include <stdio.h>

int main( int argc, char *argv[] )
{
    int numprocs, myrank, namelen, i;
    char processor_name[MPI_MAX_PROCESSOR_NAME];
    char greeting[MPI_MAX_PROCESSOR_NAME + 80];
    MPI_Status status;

    MPI_Init( &argc, &argv );
    MPI_Comm_size( MPI_COMM_WORLD, &numprocs );
    MPI_Comm_rank( MPI_COMM_WORLD, &myrank );
    MPI_Get_processor_name( processor_name, &namelen );

    sprintf( greeting, "Hello world, from process %d of %d on %s",
            myrank, numprocs, processor_name );

    if ( myrank == 0 ) {
        printf( "%s\n", greeting );
        for ( i = 1; i < numprocs; i++ ) {
            MPI_Recv( greeting, sizeof( greeting ), MPI_CHAR,
                      i, 1, MPI_COMM_WORLD, &status );
            printf( "%s\n", greeting );
        }
    } else {
        MPI_Send( greeting, strlen( greeting ) + 1, MPI_CHAR,
                  0, 1, MPI_COMM_WORLD );
    }

    MPI_Finalize( );
    return 0;
}
```

Example

```
[fred@basecm10 ~]$ module clear
[fred@basecm10 ~]$ module add intel/compiler/64 intel/mpi
[fred@basecm10 ~]$ module list
Currently Loaded Modulefiles:
  1) intel/compiler/64/2020/19.1.3  2) intel/mpi/64/2020/4.304
[fred@basecm10 ~]$ mpicc hello.c -o hello
[fred@basecm10 ~]$ mpirun -np 4 -machinefile mpirun.hosts ./hello
Hello world, from process 0 of 4 on node002
```
Hello world, from process 1 of 4 on node002
Hello world, from process 2 of 4 on node002
Hello world, from process 3 of 4 on node003

Running the executable with mpirun outside the workload manager as shown does not take the resources of the cluster into account. To handle running jobs with cluster resources is of course what workload managers such as Slurm are designed to do. Workload managers also typically take care of what environment modules should be loaded on the compute nodes for a job, via additions that the user makes to a job script.

Running an application through a workload manager via a job script is introduced in Chapter 4. Appendix A contains a number of simple MPI programs.

3.5.2 Hybridization

OpenMP is an implementation of multi-threading. This is a method of parallelizing whereby a parent thread—a series of instructions executed consecutively—forks a specified number of child threads, and a task is divided among them. The threads then run concurrently, with the runtime environment allocating threads to different processors and accessing the shared memory of an SMP system.

MPI can be mixed with OpenMP to achieve high performance on a cluster/supercomputer of multicore nodes or servers. MPI creates processes that reside on the level of node, while OpenMP forks threads on the level of a core within an SMP node. Each process executes a portion of the overall computation, while inside each process, a team of threads is created through OpenMP directives to further divide the problem. This kind of execution makes sense due to:

- the ease of programming that OpenMP provides
- OpenMP not necessarily requiring copies of data structure, which allows for designs that overlap computation and communication
- overcoming the limits of parallelism within the SMP node is of course still possible by using the power of other nodes via MPI.

Example

```c
#include<mpi.h>
#include <omp.h>
#include <stdio.h>
#include<stdlib.h>

int main(int argc , char** argv) {
    int size, myrank,namelength;
    char processor_name[MPI_MAX_PROCESSOR_NAME];
    MPI_Init(&argc,&argv);
    MPI_Comm_rank(MPI_COMM_WORLD,&myrank);
    MPI_Comm_size(MPI_COMM_WORLD,&size);
    MPI_Get_processor_name(processor_name,&namelength);
    printf("Hello I am Processor %d on %s of %d\n",myrank,processor_name,
    size);
    int tid = 0; int n_of_threads = 1;
    #pragma omp parallel default(shared) private(tid, n_of_threads)
    {
        #if defined (_OPENMP)
            n_of_threads= omp_get_num_threads();
            tid = omp_get_thread_num();
        #endif
        printf("Hybrid Hello World: I am thread # %d out of %d\n", tid, n_o
```
3.5 Compiling And Carrying Out An MPI Run

```c
f_threads);
}
MPI_Finalize();
return 0;
}
```

The program can be compiled as follows:

```
fred@basecm10 ~]$ mpicc -o hybridhello omphello.c -fopenmp
```

To specify the number of OpenMP threads per MPI task the environment variable `OMP_NUM_THREADS` must be set.

**Example**

```
fred@basecm10 ~]$ export OMP_NUM_THREADS=3
```

The number of threads specified by the variable can then be run over the hosts specified by the `mpirun.hosts` file:

```
fred@basecm10 ~]$ mpirun -np 2 -hostfile mpirun.hosts ./hybridhello
```

**Benefits And Drawbacks Of Using OpenMP**

The main benefit to using OpenMP is that it can decrease memory requirements, with usually no reduction in performance. Other benefits include:

- Potential additional parallelization opportunities besides those exploited by MPI.
- Less domain decomposition, which can help with load balancing as well as allowing for larger messages and fewer tasks participating in MPI collective operations.
- OpenMP is a standard, so any modifications introduced into an application are portable and appear as comments on systems not using OpenMP.
- By adding annotations to existing code and using a compiler option, it is possible to add OpenMP to a code somewhat incrementally, almost on a loop-by-loop basis. The vector loops in a code that vectorize well are good candidates for OpenMP.

There are also some potential drawbacks:

- OpenMP can be hard to program and/or debug in some cases.
- Effective usage can be complicated on NUMA systems due to locality considerations.
- If an application is network- or memory-bandwidth-bound, then threading it is not going to help. In this case it will be OK to leave some cores idle.
- In some cases a serial portion may be essential, which can inhibit performance.
- In most MPI codes, synchronization is implicit and happens when messages are sent and received. However, with OpenMP, much synchronization must be added to the code explicitly. The programmer must also explicitly determine which variables can be shared among threads and which ones cannot (parallel scoping). OpenMP codes that have errors introduced by incomplete or misplaced synchronization or improper scoping can be difficult to debug because the error can introduce race conditions which cause the error to happen only intermittently.
3.5.3 Support Thread Levels

MPI defines four “levels” of thread safety. The maximum thread support level is returned by the `MPI_Init_thread` call in the “provided” argument.

An environment variable `MPICH_MAX_THREAD_SAFETY` can be set to different values to increase the thread safety:

<table>
<thead>
<tr>
<th>MPICH_MAX_THREAD_SAFETY</th>
<th>Supported Thread Level</th>
</tr>
</thead>
<tbody>
<tr>
<td>not set</td>
<td>MPI_THREAD_SINGLE</td>
</tr>
<tr>
<td>single</td>
<td>MPI_THREAD_SINGLE</td>
</tr>
<tr>
<td>funneled</td>
<td>MPI_THREAD_FUNNELED</td>
</tr>
<tr>
<td>serialized</td>
<td>MPI_THREAD_SERIALIZED</td>
</tr>
<tr>
<td>multiple</td>
<td>MPI_THREAD_MULTIPLE</td>
</tr>
</tbody>
</table>

3.5.4 Further Recommendations

Users face various challenges with running and scaling large scale jobs on peta-scale production systems. For example: certain applications may not have enough memory per core, the default environment variables may need to be adjusted, or I/O may dominate run time.

Possible ways to deal with these are:

- Trying out various compilers and compiler flags, and finding out which options are best for particular applications.

- Changing the default MPI rank ordering. This is a simple, yet sometimes effective, runtime tuning option that requires no source code modification, recompilation or re-linking. The default MPI rank placement on the compute nodes is SMP style. However, other choices are round-robin, folded rank, and custom ranking.

- Using fewer cores per node is helpful when more memory per process than the default is needed. Having fewer processes to share the memory and interconnect bandwidth is also helpful in this case. For NUMA nodes, extra care must be taken.

- Hybrid MPI/OpenMP reduces the memory footprint. Overlapping communication with computation in hybrid MPI/OpenMP can be considered.

- Some applications may perform better when large memory pages are used.
Workload Management

4.1 What Is A Workload Manager?

A workload management system (also known as a queuing system, job scheduler or batch submission system) manages the available resources such as CPUs, GPUs, and memory for jobs submitted to the system by users.

Jobs are submitted by the users using job scripts. Job scripts are constructed by users and include requests for resources. How resources are allocated depends upon policies that the system administrator sets up for the workload manager.

4.2 Why Use A Workload Manager?

Workload managers are used so that users do not manually have to keep track of node usage in a cluster in order to plan efficient and fair use of cluster resources.

Users may still perhaps run jobs on the compute nodes outside of the workload manager, if that is administratively permitted. However, running jobs outside a workload manager tends to eventually lead to an abuse of the cluster resources as more people use the cluster. This leads to inefficient use of available resources. It is therefore usually forbidden as a policy by the system administrator on production clusters.

4.3 How Does A Workload Manager Function?

A workload manager uses policies to ensure that the resources of a cluster are used efficiently, and must therefore track cluster resources and jobs. A workload manager is therefore generally able to:

- Monitor:
  - the node status (up, down)
  - all available resources (available cores, memory on the nodes)
  - the state of jobs (queued, on hold, deleted, failed, completed)

- Modify:
  - the status of jobs (freeze/hold the job, resume the job, delete the job)
  - the priority and execution order for jobs
  - the run status of a job. For example, by adding checkpoints to freeze a job.
  - (optional) how related tasks in a job are handled according to their resource requirements. For example, a job with two tasks may have a greater need for disk I/O resources for the first task, and a greater need for CPU resources during the second task.

Some workload managers can adapt to external triggers such as hardware failure, and send alerts or attempt automatic recovery.
4.4 Job Submission Process

Whenever a job is submitted, the workload management system checks on the resources requested by the job script. It assigns cores, accelerators, local disk space, and memory to the job, and sends the job to the nodes for computation. If the required number of cores or memory are not yet available, it queues the job until these resources become available. If the job requests resources that are always going to exceed those that can become available, then the job accordingly remains queued indefinitely.

The workload management system keeps track of the status of the job and returns the resources to the available pool when a job has finished (that is, been deleted, has crashed or successfully completed).

4.5 What Do Job Scripts Look Like?

A job script looks very much like an ordinary shell script, and certain commands and variables can be put in there that are needed for the job. The exact composition of a job script depends on the workload manager used, but normally includes:

- commands to load relevant modules or set environment variables for the job to run
- directives for the workload manager for items associated with the job. These items can be a request for resources, output control, or setting the email addresses for messages to go to
- an execution (job submission) line

When running a job script, the workload manager is normally responsible for generating a machine file based on the requested number of processor cores (np), as well as being responsible for the allocation any other requested resources.

The executable submission line in a job script is the line where the job is submitted to the workload manager. This can take various forms.

Example

For the Slurm workload manager, the line might look like:

\texttt{\textit{srun a.out}}

Example

For PBS Professional it may simply be:

\texttt{\textit{mpirun ./a.out}}

Example

For GE it may look like:

\texttt{\textit{mpirun -np 4 -machinefile \$TMP/machines ./a.out}}

4.6 Running Jobs On A Workload Manager

The details of running jobs through the following workload managers is discussed later on, for:

- Slurm (Chapter 5)
- GE (Chapter 6)
- PBS Professional (Chapter 7)
4.7 Running Jobs In Cluster Extension Cloud Nodes Using cmjob

Extra computational power from cloud service providers such as the Amazon Elastic Compute Cloud (EC2) can be used by an appropriately configured cluster managed by NVIDIA Base Command Manager.

If the head node is running outside a cloud services provider, and at least some of the compute nodes are in the cloud, then this “hybrid” cluster configuration is called a Cluster Extension cluster, with the compute nodes in the cloud being the cloud extension of the cluster.

For a Cluster Extension cluster, job scripts to a workload manager should be submitted using NVIDIA Base Command Manager’s cmjob utility. The cmjob utility should be configured beforehand by the administrator (section 4.3 of the Cloudbursting Manual). The cmjob utility allows the job to be considered for running on the extension (the cloud nodes). Jobs that are to run on the local regular nodes (not in a cloud) are not dealt with by cmjob.

The cluster administrator must have created a certificate for cmjob use for the user that is using the cmjob job submission utility:

Example

```
[basecm10->user[fred*]]% set createcmjobcertificate yes
```

In addition, the environment module (section 2.3) cmjob is typically configured by the system administrator to load by default on the head node. It must be loaded for the cmjob utility to work.

The default settings for cmjob are configured by the cluster administrator. By default, the storage provided by Amazon’s S3 has an expiry date of 30 days (section 4.3 of the Cloudbursting Manual). Users planning to keep data in the cloud may wish to check with the administrator what the S3 expiry setting is for them, if they have not already been informed about it.

4.7.1 Introduction To Options For cmjob

The syntax for cmjob has the following form:

```
  cmjob [options] [arguments]
```

The submit option is the most important option for users. A basic example for job submission takes the following form with the submit option:

```
  cmjob submit <script>
```

Details for cmjob are given by the help option (-h|--help). The help option gives details for deeper level options, as well as at the top level. At the top level, the help text that is displayed is:

```
[fred@head ~]$ cmjob -h
usage: cmjob [-h]
   {submit, status, cancel, list-labels, list-files-for-label, create-label, delete-label, \
    upload, download, list-fsx-instances, create-fsx-instance, share-fsx-instance, delete-fsx-instance, \
    list-anf-volumes, create-anf-volume, share-anf-volume, delete-anf-volume}
   ...

Submit a job.
```

positional arguments:

```
   {submit, status, cancel, list-labels, list-files-for-label, create-label, delete-label, upload, \
    download, list-fsx-instances, create-fsx-instance, share-fsx-instance, delete-fsx-instance, \
    list-anf-volumes, create-anf-volume, share-anf-volume, delete-anf-volume}
    submit  Submit a job to execute on the compute nodes.
    status   Obtain the status of a certain job.
```
cancel  Cancel job with a specified job id.
list-labels  List all known labels
list-files-for-label  List all files stored under a certain label or list all files for all labels.
cancel  Create a label with the given name
delete-label  Delete the label with the given name
upload  Upload input data under some label.
download  Download labeled output data from previous jobs.
list-fsx-instances  List available FSx instances that can be used by this user.
create-fsx-instance  Create a new FSx instance.
share-fsx-instance  Share an owned FSx instance with other users. These users are allowed to use the instance, but will not be able to delete it.
delete-fsx-instance  Delete an existing FSx instance.
list-anf-volumes  List available ANF volumes that can be used by this user.
create-anf-volume  Create a new ANF volume.
share-anf-volume  Share an owned ANF volume with other users. These users are allowed to use the volume, but will not be able to delete it.
delete-anf-volume  Delete an existing ANF volume.

optional arguments:
-h, --help  show this help message and exit

$ cat myscript1
#!/bin/sh
hostname
$ cmjob submit myscript1
Submitting job: myscript1(slurm-2) [slurm:2] ... OK

Users can also run cmjob submit in more sophisticated ways. The submission can be done with some cloud-related options set. This can be done in two styles with cmjob:

1. command line style (page 28)
2. job directive style (page 30)

Command Line Style Options For cmjob submit
The main command line options for cmjob submit can be viewed via the help text:
usage: cmjob submit [-h] [--remote-output-list <file>]
        [-D <output file>[:label]] [-n <number>]
        [--job-name-prefix <prefix>] [-X EXTRA_FLAGS]
        [-w WLMANAGER]
        [--expected-output-size EXPECTED_OUTPUT_SIZE]
        [-i <input file>[@<new_name>][:<label>]]
        [--input-list <file>]
        [-o <output file>[@<new_name>][:<label>]]
        [--output-list <file>] [--fsx-instance <name | fsx_id>]
        [--on-demand-fsx] [--on-demand-anf] [-c <cert>] [-k <key>]
        [-u <head node url>] [-dry-run] [--debug] [-v]
        script

positional arguments:
  script                      The script that is to be executed on the compute
                           nodes. Must start with a shebang line.

optional arguments:
  -h, --help                 show this help message and exit
  --remote-output-list <file>
                           One remote file (located on the cloud nodes)
                           containing a list of output files. This remote file is
                           usually generated by the job itself.
  -D <output file>[:label], --no-download <output file>[:label]
                           Output files to not download from the object store.
                           This is useful when you want them to be consumed by a
                           subsequent job directly from the object store, or if
                           the output files are large. E.g. `cmjob -o file1:label
                           --no-download file1:product produce.sh; cmjob -i
                           file1:product consume.sh` 
  -n <number>, --num-threads <number>
                           Number of parallel file transfers to/from S3.
  --job-name-prefix <prefix>
                           String that will be used as prefix in cloud job name.
  -X EXTRA_FLAGS, --extra-flags EXTRA_FLAGS
                           Extra flags for the workload manager during
                           submission. (e.g: -X "-p partition"
  -w WLMANAGER, --wlmanager WLMANAGER
                           Workload manager used for job submission. If no name
                           is specified, then a workload manager is worked out
                           from the modules currently loaded.
  --expected-output-size EXPECTED_OUTPUT_SIZE
                           In gigabytes. Together with the size of the input
                           files, this controls the size of the allocated storage
                           volume.
  -i <input file>[@<new_name>][:<label>], --input <input file>[@<new_name>][:<label>]
                           One or more input files for the job. Can be specified
                           multiple times.
  --input-list <file>            One file containing a list of input files.
  -o <output file>[@<new_name>][:<label>], --output <output file>[@<new_name>][:<label>]
                           One or more output files produced by a cmjob job that
                           will be stored in the object store (and downloaded to
                           the head node unless --no-download is used). Can be
                           specified multiple times.
--output-list <file> 
One file containing a list of output files.

--fsx-instance <name | fsx_id>
Use a previously created FSx device instead of a storage node to read and write data from.

--on-demand-fsx
Do not use storage nodes, but create and use an on-demand FSx instance instead. The volume will be deleted upon job completion. Use --output flag to copy the output data out of the volume before it's deleted.

--anf-volume <name | id>
Use a previously created ANF volume instead of a storage node to read and write data from.

--on-demand-anf
Do not use storage nodes, but create and use an on-demand ANF volume instead. The volume will be deleted upon job completion. Use --output flag to copy the output data out of the volume before it's deleted.

-c <cert>, --cert <cert>
The cert file to use when connection to the cmdaemon api.

-k <key>, --key <key>
The key to use when connecting to the cmdaemon api.

-u <head node url>, --url <head node url>
The url of the cmdaemon api to connect to.

--dry-run
Don't submit job description to cmdaemon.

--debug
Enable debug output.

-v, --verbose
Enhance verbosity of cmjob commands.

Note: All of these options (except for -h | --help) can be used on the command line, or in the <script>

Job-directive Style #CMJOB Directives For cmjob submit
All cmjob submit command line options can also be specified in a job-directive style format in the job script itself, using the "#CMJOB" tag to indicate an option.

Example

$ cat myscript2
#!/bin/sh
#CMJOB --input-list=/home/user/myjob.in
#CMJOB --output-list=/home/user/myjob.out
#CMJOB --remote-output-list=/home/user/file-which-will-be-created
#CMJOB --input=/home/user/onemoreinput.dat
#CMJOB --input=/home/user/myexec
myexec

$ cmjob submit myscript2
Submitting job: myscript2(slurm-2) [slurm:2] ... OK

Workload Manager Job-directives And Workload Manager Flag Options For cmjob
Beside the two cmjob submit styles using options and directives, there is another way, independent of those, to pass on job handling instructions. This is using by workload manager job specifications.

Workload manager batch job tags are tags associated with the workload manager, such as #SBATCH for Slurm (Chapter 5), or #PBS for PBS Professional (Chapter 7). These tags can also be used by the user to specify flags to be sent to the workload manager when submitting the job via cmjob submit. This can be done by

- using the tag itself (e.g: #SBATCH for Slurm, #PBS for PBS Professional ...) These can be used along with the #CMJOB tags.
4.7 Running Jobs In Cluster Extension Cloud Nodes Using cmjob

Example

$ cat myscript3
#!/bin/sh
#SBATCH -p partition
hostname
$ cmjob submit myscript3
Submitting job: myscript3(slurm-3) [slurm:3] ... OK

• or by using the cmjob flag option --extra-flags to specify the flag associated with the tag.

Example

$ cat myscript4
#!/bin/sh
hostname
$ cmjob submit --extra-flags="-p partition" myscript4
Submitting job: myscript4(slurm-4) [slurm:4] ... OK

4.7.3 An Example Session Of Running cmjob With Cloud Storage

In this section a job is submitted with cmjob to a cloud compute node. For this to work, the cluster administrator should have already arranged for the cloud director and cloud compute nodes to be up.

If cloud storage is needed for a job, then cmjob causes it to be powered up. If it is not needed, then it is powered down. If AWS is used, then the cloud storage gets input from cmjob via an S3 bucket. By default the S3 bucket storage expires after 30 days.

The job submitted via cmjob could be something like the slurmhello.sh script from section 5.3.1, but with appropriate cmjob options.

Cluster administrators can use cmsh to list cloud jobs that are running via cmjob. The jobs are listed under the cloudJobs submode.

In the following example, where a job has been submitted, the cloud storage is not yet up:

Example

[root@basecm10 ~]# cmsh
[basecm10]% cmjob
[basecm10->cmjob[cmjob]]% cloudJobs
[basecm10->cmjob[cmjob]->cloudJobs]% list
Name (key) jobStatus storageNode region user
---------- ----------------------------------------------- --------------------- --------- ----
slurm-39  Waiting for available storage node... us-east-1 fred

The cmjob utility takes care of the storage requirements and directives specified by the user. In the preceding example it brings up cloud storage automatically.

Tracking Job Status With cmsh

The job status can be viewed using cmsh by the cluster administrator, and also by users with the appropriate profile privileges. Once the storage node is up, a job typically displays a sequence similar to the following in the jobStatus column:

Example

Name (key) jobStatus storageNode region user
---------- ----------------------------------------------- --------------------- --------- ----
slurm-39  Attaching EBS volume... us-east-1-cstorag001 us-east-1 fred
followed by:

```
slurm-39 Transferring input data from S3 to the storage+ us-east-1-cstorage001 us-east-1 fred
```

followed by:

```
slurm-39 Running... us-east-1-cstorage001 us-east-1 fred
```

followed by:

```
slurm-39 Transferring result from the storage node to S3 us-east-1-cstorage001 us-east-1 fred
```

followed by (for jobs with a data label):

```
slurm-39 Deregistering job as producer for labels us-east-1-cstorage001 us-east-1 fred
```

followed by:

```
slurm-39 Detaching EBS volume... us-east-1-cstorage001 us-east-1 fred
```

followed by:

```
slurm-39 Finished us-east-1-cstorage001 us-east-1 fred
```

### Tracking Job Status With `cmjob`

Regular users without `cmsh` profile privileges can track the same job status sequence via the `cmjob status <job name>` command:

**Example**

```
[fred@basecm10 ~]$ cmjob status slurm-39
```

```
+---------------+--------------------------------------------------------------------+
<table>
<thead>
<tr>
<th>Property</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Job name</td>
<td>slurm-39</td>
</tr>
<tr>
<td>Job ID</td>
<td>39</td>
</tr>
<tr>
<td>Job status</td>
<td>Transferring input data from S3 to the storage node... (0m 2s ago)</td>
</tr>
<tr>
<td>Total runtime</td>
<td>3m 31s</td>
</tr>
<tr>
<td>Storage type</td>
<td>NFSViaStorageNode</td>
</tr>
<tr>
<td>Job script</td>
<td>slurmhello.sh</td>
</tr>
<tr>
<td>Region</td>
<td>us-east-1</td>
</tr>
<tr>
<td>Job Queue</td>
<td>us-east-1</td>
</tr>
</tbody>
</table>
+---------------+--------------------------------------------------------------------+
[fred@basecm10 ~]$
```

**Running cmjob With Labels For Data**

Labels can be assigned to data placed in the cloud by a job—job A for example. Data that that is left in the cloud for the next job—job B, for example—can then be accessed by job B using that label, and then be kept in the cloud under a new label. This can be a more efficient way to run chained jobs. The data can also be taken out from the cloud and stored locally, even if this is not efficient.

In other words, a label allows users to avoid having to move data in and out of the cloud when jobs use the same data, which can save time and money.

**Scripts for the example session:** The following example session is based on the following job scripts, which use basic Unix utilities such as `seq`, `paste`, and `bc` to do some simple arithmetic:
4.7 Running Jobs In Cluster Extension Cloud Nodes Using cmjob

[fred@basecm10 ~]$ cat generate.sh
#!/bin/bash
#SBATCH -n 1
#SBATCH -p us-east-1

## make a column of numbers from 1..10
## save to file 1to10
seq 10 > 1to10

[fred@basecm10 ~]$ cat basicsum.sh
#!/bin/bash
#SBATCH -n 1
#SBATCH -p us-east-1

## sum up the column of numbers in 1to10 file
## save to file basicsum
paste -s -d+ 1to10 | bc > basicsum

[fred@basecm10 ~]$ cat square.sh
#!/bin/bash
#SBATCH -n 1
#SBATCH -p us-east-1

## Take 1to10 column and make a column of squares
## save to file squares

{ for i in $(cat 1to10)
do
echo "$i*$i" | bc
done
} > squares

[fred@basecm10 ~]$ cat squaressum.sh
#!/bin/bash
#SBATCH -n 1
#SBATCH -p us-east-1

## sum up the column of numbers in file squares
## send output to file squaredsum
paste -s -d+ squares | bc > squaredsum

Example session run for user fred: The Slurm workload manager and cmjob modules should already be loaded (module load slurm cmjob).

The cloud director and compute nodes should also be up and running.

- Submission, labels, and no downloading:
The session run starts with a user fred running:

[fred@basecm10 ~]$ cmjob submit --output 1to10:mylabel --no-download 1to10:mylabel generate.sh
Submitting workload job...
Submitted workload job slurm-74

which submits the generate.sh script. The script is run in the cloud, and it produces the numbers 1 to 10 on the compute node, which the user can pretend is a huge amount of data that is best kept in the cloud. The output is placed in a file 1to10. The --output option means that the output file
is given the label mylabel. The --no-download option means the output—accessed using the label mylabel—is prevented from being downloaded back to the directory of fred on premises, which may be sensible for a huge amount of data that may be used by another job.

The job status can be checked with cmjob status or cmjob status slurm-74:

```
[fred@basecm10 ~]$ cmjob status slurm-74
+---------------+-------------------+
| Property      | Value             |
+---------------+-------------------+
| Job name      | slurm-74          |
| Job ID        | 74                |
| Job status    | Finished          |
| Total runtime | 3m 11s            |
| Storage type  | NFSViaStorageNode |
| Job script    | generate.sh       |
| Region        | us-east-1         |
| Job Queue     | us-east-1         |
+---------------+-------------------+
```

The label status can be checked with cmjob list-labels:

```
[fred@basecm10 ~]$ cmjob list-labels
+---------+-----------------------------------+---------------------------------------+
| Label   | Created at                         | Expires at                           |
+---------+-----------------------------------+---------------------------------------+
| mylabel | 2020-04-20 00:26:27 - 15m 54s ago | 2020-05-20 02:00:00 - 30d 1h from now |
+---------+-----------------------------------+---------------------------------------+
```

The files stored in the object store under a label can be checked using cmjob list-files-for-label:

```
[fred@basecm10 ~]$ cmjob list-files-for-label .
+------------------+---------+-----------------------------------+------+
| Name             | Label   | Last modified                     | Size |
+------------------+---------+-----------------------------------+------+
| /home/fred/1to10 | mylabel | 2020-04-20 00:30:17 - 12m 11s ago | 21   |
+------------------+---------+-----------------------------------+------+
```

If needed, the 1to10 file can be checked to see if it exists, and how it looks. It can be brought down from the cloud with the download option of cmjob:

```
[fred@basecm10 ~]$ cmjob download --output 1to10:mylabel
Submitting download job...
Submitted download job download-54
[fred@basecm10 ~]$ cat 1to10
1
2
3
4
5
6
7
8
9
10
```
• Using a large object already in the cloud and bringing out a result locally:
The next job that is to be run, can pick up the labeled object in the cloud and do something with it. In this case, it picks up the numbers as input from the labeled object file in the cloud, and sums the numbers up with the script in the cloud compute node. The output of the script, which the user can pretend is considerably smaller than the input, can then be placed as output in the local directory of fred using the -output option:

    [fred@basecm10 ~]$ cmjob submit --input 1to10:mylabel --output basicsum basicsum.sh
    Submitting workload job...
    Submitted workload job slurm-75

After the job is finished (according to the report from cmjob status slurm-75), the file basicsum shows up locally:

    [fred@basecm10 ~]$ cat basicsum
    55

• Processing large data in the cloud and leaving the data there:
In the next case considered, the square.sh file can be run so that it uses the “large” file labeled with mylabel,1to10, and uses it as input data. It keeps the output file, squares, which is also a “large” file, in the object store in the cloud using the label numbers2:

    [fred@basecm10 ~]$ cmjob submit --input 1to10:mylabel --output squares:numbers2
    --no-download *:numbers2 square.sh
    Submitting workload job...
    Submitted workload job slurm-76

After that job is run, its output file, squares, can be picked up and checked locally with:

    [fred@basecm10 ~]$ cmjob download --output squares:numbers2
    Submitting download job...
    Submitted download job download-57
    [fred@basecm10 ~]$ cat squares
    1
    4
    9
    16
    25
    36
    49
    64
    81
    100

If the download is repeated, or otherwise risks overwriting a local file, then the user is prompted with a warning.
However, in any case, the idea is to avoid unnecessary downloads of the “large” file, so typically a download is not done for large data.

• Processing the new large data in the cloud and downloading a reduced end result:
Continuing on, this new “large” data can be processed with the script squaressum.sh, to sum the squares file in the object store in the cloud. The squares file is accessed in the cloud with the label numbers2, and reduced to a single number. The shrunk result is brought down from the cloud and kept locally as the file squaredsum:
Workload Management

[fred@basecm10 ~]$ cmjob submit --input squares:numbers2 --output squaredsum squaresum.sh

The job status can be checked with:

```
cmjob status <job name>
```

When the job is in the finished state, the user can run:

[fred@basecm10 ~]$ cat squaredsum
385

Processing Using High-Performance AWS FSx For Lustre Storage:

The user Fred may wish to use a high-performance filesystem in the cloud. Lustre is a
high-performance parallel filesystem aimed at HPC needs. Amazon offers AWS FSx with
Lustre in some regions of AWS, as documented at https://aws.amazon.com/about-aws/
global-infrastructure/regional-product-services/.

Region restrictions: Changes may need to be carried out to the region used, to make use of FSx.

For example, the scripts generate.sh and so on from page 32 onward used a region, us-east-1,
that at the time of writing (May 2020) did not have FSx. The region us-west-1 on the other hand
did have FSx.

So, if the user would like to have jobs using FSx, and the cluster administrator has set up the
FSx-enabled region, such as us-west-1, with a cloud director, cloud nodes, and cloud storage,
then the user should also change the Slurm queue option (the partition option, -p) in the scripts
generate.sh and so on from page 32.

[An aside for the cluster administrator: The cluster administrator can manage the Slurm queues
for the region via the jobqueue submode (cmsh → wlm → jobqueue, section 7.7.2 of the Administrator
Manual). A Slurm queue for cloud storage can also be created as an option when the cluster
administrator runs cm-cloud-storage-setup when setting up a region.]

Permission restrictions: The high performance of Lustre is expensive, so a cluster administrator
typically restricts its use. Some permissions may therefore be needed before a user can use Lustre.
[Aside for the administrator: Granting users FSx-related permissions is described on page 73 of
the Cloudbursting Manual.]

Three ways to launch FSx: Once the appropriate tokens are set, the user can launch FSx volumes
for their jobs in three ways—on-demand, user-managed, or admin-managed. The user should check
with the cluster administrator which of these is available:

1. **On-demand:** With this way, the option --on-demand-fsx can be added to cmjob. This creates
an FSx volume on demand, instead of a storage node, and the FSx volume is deleted after the
job ends.

   For example, a session can be run, where the user Fred submits a job in a region that supports
   FSx, with

   ```
   [fred@basecm10 ~]$ cmjob submit --on-demand-fsx --output 1to10:mylabel2 generate.sh
   ```

   When the job is completed, the FSx volume that came up on demand, is deleted.

2. **User-managed:** A more hands-on way to manage an FSx instance is with a create-use-delete
sequence that is run by the user.
4.7 Running Jobs In Cluster Extension Cloud Nodes Using cmjob

(a) Creation can be carried out in this format:

```
cmjob create-fsx-instance <FSx instance name>
```

The command creates an FSx instance with a name `<FSx instance name>` that the user specifies. On execution, the FSx instance ID is displayed. FSx instance IDs can also be seen if the command `cmjob list-fsx-instances` is run.

(b) Jobs can then be submitted with the `--fsx-instance` option, in a format similar to:

```
cmjob submit --fsx-instance <FSx instance name, or FSx instance ID>
```

and the jobs then use the pre-created instance.

(c) Unlike the `--on-demand-fsx` option, the FSx volume is not deleted when the job ends. So, after use, the instance must be deleted explicitly with a command in a form such as:

```
cmjob delete-fsx-instance <FSx instance name, or FSx instance ID>
```

Thus a session can be run where a user fred creates an FSx instance, `fsxa`, with:

```
Example

[fred@basecm10 ~]$ cmjob create-fsx-instance fsxa
FSx instance 'fsxa' is being created. The default capacity will be used.
The instance is being created with id fs-07f668884b4eda47c in region 'us-west-1' and will be made available in VPC 'vpc-07f3150b0bdaa9e'. Capacity: 1200GiB.
This might take several minutes. Use 'list-fsx-instances' to see the progress.
[fred@basecm10 ~]$ cmjob list-fsx-instances
+------+----------------------+----------+----------+------+
| Name | ID | Status | Capacity | Jobs |
+------+----------------------+----------+----------+------+
| fsxa | fs-07f668884b4eda47c | CREATING | 1200 | |
+------+----------------------+----------+----------+------+
[fred@basecm10 ~]$ cmjob list-fsx-instances
+------+----------------------+-----------+----------+------+
| Name | ID | Status | Capacity | Jobs |
+------+----------------------+-----------+----------+------+
| fsxa | fs-07f668884b4eda47c | AVAILABLE | 1200 | |
+------+----------------------+-----------+----------+------+
```

Once the FSx storage is available, it can be used merely by adding the `--fsx-instance` option, along with the FSx storage identifier, to `cmjob`:

```
Example

[fred@basecm10 ~]$ cmjob submit --fsx-instance fsxa --output 1to10:mylabel3 generate.sh
Submitting workload job...
Submitted workload job slurm-4
```

Once the job is finished, and fred no longer requires the data, the instance should be deleted:

```
Example

[fred@basecm10 ~]$ cmjob delete-fsx-instance fsxa
About to delete this FSX instance
fsxa (fs-07f668884b4eda47c)
  owner: fred
  status: AVAILABLE
  capacity: 1200GB
  created at: 2020-05-01T21:55:52.162000+02:00
  managed: UserManaged
  shared with: no one
Enter 'yes' to continue> yes
Deleting FSx instance fs-07f668884b4eda47c
[fred@basecm10 ~]$
```
3. **Admin-managed:** With the admin-managed option, it is the cluster administrator who creates an FSx instance, and who shares it with one or more other regular users. Users are then able to use the instance just as in user-managed instances, but they cannot delete the instance or view the files of other users.

- **Processing Using High-Performance ANF Volumes:**

  ANF (Azure NetApp Files) is a high-performance filesystem available in Azure clouds. The procedures to use it are very similar to that for FSx volumes in AWS clouds.

  The cluster administrator can set up a cloud director, cloud nodes, and cloud storage, in an ANF-enabled region, such as westeurope. If a regular user would like to have jobs using ANF, and the user is using the scripts `generate.sh` and so on from page 32, then the user should set the Slurm queue option (the partition option, `-p`) so that that ANF-enabled region is used.

  [An aside for the cluster administrator: The cluster administrator can manage the Slurm queues for the region via the `jobqueue` submode (`cmsh → wlm → jobqueue`, section 7.7.2 of the Administrator Manual). A Slurm queue for cloud storage can also be created as an option when the cluster administrator runs `cm-cloud-storage-setup` when setting up a region.]

  **Permission restrictions:** ANF is expensive to use, so a cluster administrator typically restricts its use. Some permissions may therefore be needed before a user can use ANF.

  [Aside for the administrator: Granting users ANF-related permissions is described on page 95 of the Cloudbursting Manual.]

**Three ways to launch ANF:** Once the appropriate tokens are set, the user can launch ANF volumes for their jobs in three ways—on-demand, user-managed, or admin-managed. The user should check with the cluster administrator which of these is available:

1. **On-demand:** With this way, the option `--on-demand-anf` can be added to `cmjob`. This creates an ANF volume on demand, instead of a storage node, and the ANF volume is deleted after the job ends.
   
   For example, a session can be run, where the user `fred` submits a job in a region that supports ANF, with
   
   ```bash
   [fred@basecm10 ~]$ cmjob submit --on-demand-anf --output 1to10:mylabel2 generate.sh
   ``
   
   When the job is completed, the ANF volume that came up on demand, is deleted.

2. **User-managed:** A more hands-on way to manage an ANF instance is with a create-use-delete sequence that is run by the user.
   
   (a) Creation can be carried out in this format:
   
   ```bash
   cmjob create-anf-volume <ANF volume name>
   ```
   
   The command creates an ANF volume with a name `<ANF volume name>` that the user specifies. On execution, the ANF instance ID is displayed. ANF volume IDs can also be seen if the command `cmjob list-anf-volumes` is run.
   
   (b) Jobs can then be submitted with the `--anf-volume` option, in a format similar to:
   
   ```bash
   cmjob submit --anf-volume <ANF volume name, or ANF volume ID>
   ```
   
   and the jobs then use the pre-created instance.
   
   (c) Unlike the `--on-demand-anf` option, the ANF volume is not deleted when the job ends. So, after use, the instance must be deleted explicitly with a command in a form such as:
   
   ```bash
   cmjob delete-anf-volume <ANF volume name, or ANF volume ID>
   ```
   
   Thus a session can be run where a user `fred` creates an ANF instance, `anfa`, with:
4.7 Running Jobs In Cluster Extension Cloud Nodes Using cmjob

Example

[fred@basecm10 ~]$ cmjob create-anf-volume anfa
ANF volume 'anfa' is being created. The default size will be used.
The volume named anfa (anfa-jrcjlSJXgP) is created in location 'westeurope' and will be made available in resource group 'pj-mojo-westeurope-bcm'. Capacity: 4TiB.
This might take several minutes. Use 'list-anf-volumes' to see the progress.
[fred@basecm10 ~]$ cmjob list-anf-volumes

<table>
<thead>
<tr>
<th>Name</th>
<th>ID</th>
<th>Status</th>
<th>Capacity</th>
<th>Jobs</th>
</tr>
</thead>
<tbody>
<tr>
<td>anfa</td>
<td>anfa-jrcjlSJXgP</td>
<td>CREATING</td>
<td>4TiB</td>
<td></td>
</tr>
</tbody>
</table>

[fred@basecm10 ~]$ cmjob list-anf-volumes

<table>
<thead>
<tr>
<th>Name</th>
<th>ID</th>
<th>Status</th>
<th>Capacity</th>
<th>Jobs</th>
</tr>
</thead>
<tbody>
<tr>
<td>anfa</td>
<td>anfa-jrcjlSJXgP</td>
<td>AVAILABLE</td>
<td>4TiB</td>
<td></td>
</tr>
</tbody>
</table>

Once the ANF volume is available, it can be used merely by adding the --anf-volume option, along with the ANF volume identifier, to cmjob:

Example

[fred@basecm10 ~]$ cmjob submit --anf-volume anfa --output 1to10:mylabel3 generate.sh
Submitting workload job...
Submitted workload job slurm-4

Once the job is finished, and fred no longer requires the data, the instance should be deleted:

Example

[fred@basecm10 ~]$ cmjob delete-anf-volume anfa
About to delete this ANF volume
anfa (anfa-jrcjlSJXgP)
owner: fred
status: AVAILABLE
size: 4TB
created at: 2020-05-10T23:16:07.871298Z
managed: UserManaged
shared with: no one
Enter 'yes' to continue> yes
Deleting ANF volume anfa
[fred@basecm10 ~]$ 

3. Admin-managed: With the admin-managed option, it is the cluster administrator who creates an ANF volume, and who shares it with one or more other regular users. Users are then able to use the instance just as in user-managed instances, but they cannot delete the instance or view the files of other users.
Slurm

Slurm is a workload management system developed originally at the Lawrence Livermore National Laboratory. Slurm used to stand for Simple Linux Utility for Resource Management. However Slurm has evolved since then, and its advanced state nowadays means that the acronym is obsolete.

Slurm has both a graphical interface and command line tools for submitting, monitoring, modifying and deleting jobs. It is normally used with *job scripts* to submit and execute jobs. Various settings can be put in the job script, such as number of processors, resource usage, and application specific variables.

The steps for running a job through Slurm are to:

- Create the script or executable that will be handled as a job
- Create a job script that sets the resources for the script/executable
- Submit the job script to the workload management system

The details of Slurm usage depends upon the MPI implementation used. The description in this chapter will cover using Slurm’s Open MPI implementation, which is quite standard. Slurm documentation can be consulted ([https://slurm.schedmd.com/mpi_guide.html](https://slurm.schedmd.com/mpi_guide.html)) if the implementation the user is using is very different.

### 5.1 Loading Slurm Modules And Compiling The Executable

In section 3.5.1 an MPI “Hello, world!” executable that can run in parallel is created and run in parallel outside a workload manager.

The executable can be run in parallel using the Slurm workload manager. For this, the Slurm module should first be loaded by the user on top of the chosen MPI implementation, in this case Open MPI:

#### Example

```
[fred@basecm10 ~]$ module list
Currently Loaded Modulefiles:
  1) gcc/11.2.0    2) openmpi/gcc/64/4.1.1
[fred@basecm10 ~]$ module add slurm; module list
Currently Loaded Modulefiles:
  1) gcc/11.2.0    2) openmpi/gcc/64/4.1.1    3) slurm/slurm/21.08.8
```

The “hello world” executable from section 3.5.1 can then be compiled and run for one task outside the workload manager, on the local host, as:

```
[fred@basecm10 ~]$ mpicc hello.c -o hello
[fred@basecm10 ~]$ mpirun -np 1 hello
```

Adding a full path to `mpirun`, and adding a machine file, would allow it to run on the machine file hosts, instead of just locally.
5.2 Running The Executable With `salloc`

Running it as a job managed by Slurm can be done interactively with the Slurm allocation command, `salloc`, as follows:

```
[fred@basecm10 ~]$ salloc mpirun hello
```

Slurm is more typically run as a batch job (section 5.3). However execution via `salloc` uses the same options, and it is more convenient as an introduction because of its interactive behavior.

In a default BCM configuration, Slurm auto-detects the compute node cores available, and by default spreads the tasks across the cores as part of the allocation request. Specifying a machine file is therefore not required.

To change how Slurm spreads the executable across compute nodes is typically determined by the options in the following table:

<table>
<thead>
<tr>
<th>Short Option</th>
<th>Long Option</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>-N</td>
<td>--nodes=</td>
<td>Request this many nodes on the cluster.</td>
</tr>
<tr>
<td>-n</td>
<td>--ntasks=</td>
<td>Request this many tasks on the cluster.</td>
</tr>
<tr>
<td>-c</td>
<td>--cpus-per-task=</td>
<td>Request this many CPUs per task.</td>
</tr>
<tr>
<td>(none)</td>
<td>--ntasks-per-node=</td>
<td>Request this number of tasks per node.</td>
</tr>
</tbody>
</table>

Specifying `-N1` (indicating 1 node) or `-n4` (indicating 4 tasks) is specified.

The full options list and syntax for `salloc` can be viewed with “`man salloc`”.

The requirement of specified options to `salloc` must be met before the executable is allowed to run. So, for example, if `-nodes=4` and the cluster only has 3 nodes, then the executable does not run.

5.2.1 Node Allocation Examples

The following session illustrates and explains some node allocation options and issues for Slurm using a cluster with just 1 compute node and 4 CPU cores:

**Default settings:** The `hello` MPI executable with default settings of Slurm runs successfully over the first (and in this case, the only) node that it finds:

```
[fred@basecm10 ~]$ salloc mpirun hello
salloc: Granted job allocation 572
Hello world from process 0 out of 4, host name node001
Hello world from process 1 out of 4, host name node001
Hello world from process 2 out of 4, host name node001
Hello world from process 3 out of 4, host name node001
salloc: Relinquishing job allocation 572
```

The preceding output also displays if `-N1` (indicating 1 node) is specified, or if `-n4` (indicating 4 tasks) is specified.

The node and task allocation is almost certainly not going to be done by relying on defaults. Instead, node specifications are supplied to Slurm along with the executable.

To understand Slurm node specifications, the following cases consider and explain where the node specification is valid and invalid.
5.2 Running The Executable With `salloc`

**Number of nodes requested:** The value assigned to the `-N` or `--nodes` option is the number of nodes from the cluster that is requested for allocation for the executable. In the current cluster example it can only be 1. For a cluster with, for example, 1000 compute nodes, it could be a number up to 1000.

A resource allocation request for 2 nodes with the `--nodes` option halts for the current cluster which only has 1 compute node:

```
[fred@basecm10 ~]$ salloc -N2 mpirun hello
salloc: Requested partition configuration not available now
salloc: Pending job allocation 573
salloc: job 573 queued and waiting for resources
```

The default behavior is to patiently wait for resources to become available. This makes sense if a cluster can increase its available resources within a reasonable time period. The user can interrupt an `salloc` in this state with a `ctrl-c`.

**Number of tasks requested per cluster:** The value assigned to the `-n` or `--ntasks` option is the number of tasks that are requested for allocation from the cluster for the executable. In the current cluster example, which has a single compute node with 4 cores, it can be 1 to 4 tasks. The default CPU resources available on a cluster are the number of available processor cores on the compute nodes, which is 4 on this cluster with a single compute node of 4 cores.

A resource allocation request for 5 tasks for this cluster halts because it exceeds the default resources available on the 4-core cluster:

```
[fred@basecm10 ~]$ salloc -n5 mpirun hello
salloc: Requested partition configuration not available now
salloc: Pending job allocation 574
salloc: job 574 queued and waiting for resources
```

Adding and configuring just one more compute node to the current cluster would allow the resource allocation to succeed, since that would provide at least one more core to the cluster.

**Number of tasks requested per node:** The value assigned to the `--ntasks-per-node` option is the number of tasks that are requested for allocation from each compute node on the cluster. In the current cluster example, it can be 1 to 4 tasks. A resource allocation request for 5 tasks per compute node with `--ntasks-per-node` results in a similar output on this cluster running a single compute node with 4 cores. It gives an output like:

```
[fred@basecm10 ~]$ salloc --ntasks-per-node=5 mpirun hello
salloc: Requested partition configuration not available now
salloc: Pending job allocation 575
salloc: job 575 queued and waiting for resources
```

Adding and configuring another 4-core node to the current cluster would still not allow resource allocation to succeed, because the request is for at least 5 cores per compute node, rather than per cluster.

**Restricting the number of tasks that can run per compute node:** A resource allocation request for 2 tasks per compute node with the `--ntasks-per-node` option, and simultaneously an allocation request for 1 task to run on the cluster using the `--ntasks` option, runs successfully, although it uselessly leaves 2 cores unused on the compute node:

```
[fred@basecm10 ~]$ salloc --ntasks-per-node=2 --ntasks=1 mpirun hello
salloc: Granted job allocation 576
Hello world from process 000 out of 002, processor name node001
Hello world from process 001 out of 002, processor name node001
salloc: Relinquishing job allocation 576
```
The other way round, that is, a resource allocation request for 1 task per node with the
--ntasks-per-node option, and simultaneously an allocation request for 2 tasks to run on the cluster using the --ntasks option, fails on this cluster running 1 compute node with 4 cores. This is because although 1 task can be allocated resources from the single node, resources for 2 tasks are being asked for on the cluster, which requires 2 nodes:

[fred@basecm10 ~]$ salloc --ntasks-per-node=1 --ntasks=2 mpirun hello
salloc: error: Job submit/allocate failed: Requested node configuration is not available
salloc: Job allocation 577 has been revoked.

5.3 Running The Executable As A Slurm Job Script

Instead of using options appended to the salloc command line as in section 5.2, it is usually more convenient to send jobs to Slurm with the sbatch command acting on a job script.

A job script is also sometimes called a batch file. In a job script, the user can add and adjust the Slurm options, which are the same as the salloc options of section 5.2. The various settings and variables that go with the application can also be adjusted.

5.3.1 Slurm Job Script Structure

A job script submission for the Slurm batch job script format is illustrated by the following:

[fred@basecm10 ~]$ cat slurmhello.sh
#!/bin/sh
#SBATCH -o my.stdout
#SBATCH --time=30 #time limit to batch job
#SBATCH -N 4
#SBATCH --ntasks=16
#SBATCH --ntasks-per-node=4
module add gcc/9.2.0 openmpi/gcc/64/1.10.7 slurm
mpirun hello

The structure is:

shebang line: shell definition line.
SBATCH lines: optional job script directives (section 5.3.2).
shell commands: optional shell commands, such as loading necessary modules.
application execution line: execution of the MPI application using sbatch, the Slurm submission wrapper.

In SBATCH lines, "#SBATCH" is used to submit options. The various meanings of lines starting with "#" are:

<table>
<thead>
<tr>
<th>Line Starts With</th>
<th>Treated As</th>
</tr>
</thead>
<tbody>
<tr>
<td>#</td>
<td>Comment in shell and Slurm</td>
</tr>
<tr>
<td>#SBATCH</td>
<td>Comment in shell, option in Slurm</td>
</tr>
<tr>
<td>#SBATCH</td>
<td>Comment in shell and Slurm</td>
</tr>
</tbody>
</table>
After the Slurm job script is run with the `sbatch` command (Section 5.3.4), the output goes into file `my.stdout`, as specified by the “-o” command.

If the output file is not specified, then the file takes a name of the form “`slurm-<jobnumber>.out`”, where `<jobnumber>` is a number starting from 1.

The command “`sbatch --usage`” lists possible options that can be used on the command line or in the job script. Command line values override script-provided values.

### 5.3.2 Slurm Job Script Options

Options, sometimes called “directives”, can be set in the job script file using this line format for each option:

```
#SBATCH {option} {parameter}
```

Directives are used to specify the resource allocation for a job so that Slurm can manage the job optimally. Available options and their descriptions can be seen with the output of `sbatch --help`. The more overviewable usage output from `sbatch --usage` may also be helpful.

Some of the more useful ones are listed in the following table:

<table>
<thead>
<tr>
<th>Directive Description</th>
<th>Specified As</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name the job <code>&lt;jobname&gt;</code></td>
<td><code>#SBATCH -J &lt;jobname&gt;</code></td>
</tr>
<tr>
<td>Request at least <code>&lt;minnodes&gt;</code> nodes</td>
<td><code>#SBATCH -N &lt;minnodes&gt;</code></td>
</tr>
<tr>
<td>Request <code>&lt;minnodes&gt;</code> to <code>&lt;maxnodes&gt;</code> nodes</td>
<td><code>#SBATCH -N &lt;minnodes&gt;-&lt;maxnodes&gt;</code></td>
</tr>
<tr>
<td>Request at least <code>&lt;MB&gt;</code> amount of temporary disk space</td>
<td><code>#SBATCH --tmp &lt;MB&gt;</code></td>
</tr>
<tr>
<td>Run job for a time of <code>&lt;walltime&gt;</code> minutes</td>
<td><code>#SBATCH -t &lt;walltime&gt;</code></td>
</tr>
<tr>
<td>Run job at <code>&lt;time&gt;</code> (format: HH:MM MM/DD/YY)</td>
<td><code>#SBATCH --begin &lt;time&gt;</code></td>
</tr>
<tr>
<td>Set the working directory to <code>&lt;directorypath&gt;</code></td>
<td><code>#SBATCH --D &lt;directorypath&gt;</code></td>
</tr>
<tr>
<td>Set error log name to <code>&lt;jobname.err&gt;</code>*</td>
<td><code>#SBATCH -e &lt;jobname.err&gt;</code></td>
</tr>
<tr>
<td>Set output log name to <code>&lt;jobname.log&gt;</code>*</td>
<td><code>#SBATCH -o &lt;jobname.log&gt;</code></td>
</tr>
<tr>
<td>Mail <code>&lt;user@address&gt;</code> on job state change</td>
<td><code>#SBATCH --mail-user &lt;user@address&gt;</code></td>
</tr>
<tr>
<td>Mail on all state changes</td>
<td><code>#SBATCH --mail-type=ALL</code></td>
</tr>
<tr>
<td>Mail on job end</td>
<td><code>#SBATCH --mail-type=END</code></td>
</tr>
<tr>
<td>Run job in partition</td>
<td><code>#SBATCH -p &lt;destination&gt;</code></td>
</tr>
<tr>
<td>Run job using GPU with ID <code>&lt;number&gt;</code>, as described in section 8.5.2</td>
<td><code>#SBATCH --gres=gpu: &lt;number&gt;</code></td>
</tr>
</tbody>
</table>

*By default, both standard output and standard error go to a file: `slurm-%j%.out` where `<%j>` is the job number.

### 5.3.3 Slurm Environment Variables

Available environment variables include:

- `SLURM_CLUSTER_NAME` - name of the Slurm cluster
- `SLURM_CPUS_ON_NODE` - CPUs on allocated node
- `SLURM_JOB_ID` - job ID of executing job
- `SLURM_JOB_NODELIST` - list of nodes allocated to job
- `SLURM_JOB_NUM_NODES` - total number of nodes in job’s resource allocation
- `SLURM_JOB_PARTITION` - partition of job
- `SLURM_NODEID` - ID of the nodes allocated
- `SLURM_NTASKS` - total number of processes in current job (same as `-n|--ntasks=)
- `SLURM_PROCID` - MPI rank (or relative process ID) of the current process
SLURM_SUBMIT_DIR - directory from which job was launched
SLURM_TASK_PID - process ID of task started
SLURM_TASKS_PER_NODE - number of tasks to be run on each node (man page gives specification)

Typically, end users use SLURM_PROCID in a program so that an input of a parallel calculation depends on it. The calculation is thus spread across processors according to the assigned SLURM_PROCID, so that each processor handles the parallel part of the calculation with different values.

More information on environment variables is also to be found in the man page for sbatch.

5.3.4 Submitting The Slurm Job Script With sbatch

Submitting a Slurm job script created as in the previous section is done by executing the job script with sbatch:

```
[fred@basecm10 ~]$ sbatch slurmhello.sh
Submitted batch job 604
[fred@basecm10 ~]$ cat my.stdout
Hello world from process 001 out of 016, processor name node001
...
```

Queues in Slurm terminology are called “partitions”. Slurm has a default queue called defq. The administrator may have removed this or created others.

If a particular queue is to be used, this is typically set in the job script using the -p or --partition option:

```
#SBATCH --partition=bitcoinsq
```

It can also be specified as an option to the sbatch command during submission to Slurm.

5.3.5 Checking And Changing Queued Job Status With squeue, scancel, scontrol And sview

Job Queue Listing With squeue

After a job has gone into a queue, the queue status can be checked using the squeue command. The job number can be specified with the -j option to avoid seeing other jobs.

Example

```
[fred@basecm10 ~]$ squeue
       JOBID PARTITION   NAME     USER ST TIME NODES  NODELIST(REASON)
          673    defq   slurmhel   fred PD 0:00  5 (PartitionNodeLimit)
          683    defq   slurmhel   fred PD 0:00  4 (Resources)
          684    defq   slurmhel   fred PD 0:00  4 (Resources)
          685    defq   slurmhel   fred PD 0:00  4 (Resources)
          682    defq   slurmhel   fred R  0:02 4 node[001-004]
[fred@basecm10 ~]$ squeue -j 673
       JOBID PARTITION   NAME     USER ST TIME NODES  NODELIST(REASON)
          673    defq   slurmhel   fred PD 0:00  5 (PartitionNodeLimit)
```

The man page for squeue covers other options, and explains the meaning of the output.

A backstory for this squeue listing is that in JOBID 673 the user has tried to batch submit the slurmhello.sh job, while requesting 5 nodes, using the entry #SBATCH -N 5 in slurmhello.sh. However, since only 4 nodes were available for the cluster at the time, the submission is pending.

After modifying the entry to #SBATCH -N 4, and submitting the job several times, the jobs 674 to 681 complete successfully. The job with the ID 682 is seen in the R (running) state, and the rest are in a PD (pending) state, with the reasons being either

- Resources (that there are no resources yet available), or
- PartitionNodeLimit (the number of nodes required by this job cannot yet be met because it is outside of its partition’s current limits).
Job Canceling With `scancel`

Jobs can be canceled quietly by job ID with “`scancel <job number>`”. The `-v` option gives some feedback. Verbosity increases with `-vv`, `-vvv`, and `-vvvv`.

Example

```
[fred@basecm10 ~]$ scancel -v 673
scancel: Terminating job 673
```

Other cancel options, such as per state (with `-t|--state=`), per partition (with `-p|--partition=`), are also possible. The man page for `scancel` has details.

Queued Job Changes With `scontrol`

The `scontrol` command allows users to see and change the job directives while the job is still queued. For example, a user may have specified a job, using the `--begin` directive, to start at 10am the next day by mistake. To change the job to start at 10pm tonight, something like the following session may take place:

```
[fred@basecm10 ~]$ scontrol show jobid=254 | grep Time
  RunTime=00:00:04 TimeLimit=UNLIMITED TimeMin=N/A
  SubmitTime=2011-10-18T17:41:34 EligibleTime=2011-10-19T10:00:00
  StartTime=2011-10-18T17:44:15 EndTime=Unknown
  SuspendTime=None SecsPreSuspend=0
```

The parameter that should be changed is “`EligibleTime`”, which can be done as follows:

```
[fred@basecm10 ~]$ scontrol update jobid=254 EligibleTime=2011-10-18T22:00:00
```

GUI Slurm Control `sview`

An approximate GUI Slurm equivalent to `scontrol` is the `sview` tool. This allows the job to be viewed under its jobs tab, and the job to be edited with a right click menu item. It can also carry out many other functions, including canceling a job.

Webbrowser-accessible job viewing is possible from the workload tab of the `User Portal` (section 12.2).
Altair Grid Engine

Altair Grid Engine (GE) is a workload management and job scheduling system. It is a further development by Univa of Sun Grid Engine (SGE). SGE was developed by Sun Microsystems before 2011, and was at one time open source. Univa Grid Engine (UGE) was a proprietary fork of it. After Univa was acquired by Altair, UGE engine is now called Altair Grid Engine (GE).

GE has both a graphical interface and command line tools for submitting, monitoring, modifying and deleting jobs.

GE uses job scripts to submit and execute jobs. Various settings can be put in the job script, such as number of processors, resource usage and application-specific variables.

The steps for running a job through GE are to:

- Create a job script
- Select the directives to use
- Add the scripts and applications and runtime parameters
- Submit it to the workload management system

6.1 Writing A Job Script

A binary cannot be submitted directly to GE—a job script is needed for that. A job script can contain various settings and variables to go with the application. A job script format looks like:

```
#!/bin/bash
#$ Script options # Optional script directives
shell commands # Optional shell commands
application # Application itself
```

6.1.1 Directives

It is possible to specify options ('directives') to GE by using ‘#$’ in the script. The difference in the meaning of lines that start with the ‘#’ character in the job script file should be noted:

<table>
<thead>
<tr>
<th>Line Starts With</th>
<th>Treated As</th>
</tr>
</thead>
<tbody>
<tr>
<td>#</td>
<td>Comment in shell and GE</td>
</tr>
<tr>
<td>#$</td>
<td>Comment in shell, directive in GE</td>
</tr>
<tr>
<td># $</td>
<td>Comment in shell and GE</td>
</tr>
</tbody>
</table>
### 6.1.2 GE Environment Variables

Available environment variables for GE can be seen in the man page for qsub. Some of these are called pseudo environment variables:

- `$HOME` - Home directory on execution machine
- `$USER` - User ID of job owner
- `$JOB_ID` - Current job ID
- `$JOB_NAME` - Current job name; (like the `-N` option in qsub, qsh, qrsh, qlogin and qalter)
- `$HOSTNAME` - Name of the execution host
- `$TASK_ID` - Array job task index number

There are also about 30 environment variables beginning with the prefix `SGE_`.

### 6.1.3 Job Script Options

Options can be set in the job script file using this line format for each option:

```bash
#$ {option} {parameter}
```

Available options and their descriptions can be seen with the output of `qsub -help`:

<table>
<thead>
<tr>
<th>Option and parameter</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>-a date_time</code></td>
<td>request a start time</td>
</tr>
<tr>
<td><code>-adds list_attr_name attr_key attr_value</code></td>
<td>add additional sublist parameter</td>
</tr>
<tr>
<td><code>-ac context_list</code></td>
<td>add context variables</td>
</tr>
<tr>
<td><code>-ar ar_id</code></td>
<td>bind job to advance reservation</td>
</tr>
<tr>
<td><code>-A account_string</code></td>
<td>account string in accounting record</td>
</tr>
<tr>
<td>`-b y[es]</td>
<td>n[oe]`</td>
</tr>
<tr>
<td>`-binding [env</td>
<td>pe</td>
</tr>
<tr>
<td><code>-c ckpt_selector</code></td>
<td>define type of checkpointing for job</td>
</tr>
<tr>
<td><code>-ckpt ckpt-name</code></td>
<td>request checkpoint method</td>
</tr>
<tr>
<td><code>-clear</code></td>
<td>skip previous definitions for job</td>
</tr>
<tr>
<td><code>-clearp attr_name</code></td>
<td>clear/delete a parameter</td>
</tr>
<tr>
<td><code>-clears list_attr_name attr_key</code></td>
<td>clear/delete sublist parameter</td>
</tr>
<tr>
<td><code>-cwd</code></td>
<td>use current working directory</td>
</tr>
<tr>
<td><code>-C directive_prefix</code></td>
<td>define command prefix for job script</td>
</tr>
<tr>
<td><code>-dc simple_context_list</code></td>
<td>delete context variable(s)</td>
</tr>
<tr>
<td><code>-di date_time</code></td>
<td>request a deadline initiation time</td>
</tr>
<tr>
<td><code>-e path_list</code></td>
<td>specify standard error stream path(s)</td>
</tr>
</tbody>
</table>

...continued
<table>
<thead>
<tr>
<th>Option and parameter</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>-h</td>
<td>place user hold on job</td>
</tr>
<tr>
<td>-hard</td>
<td>consider following requests &quot;hard&quot;</td>
</tr>
<tr>
<td>-help</td>
<td>print this help</td>
</tr>
<tr>
<td>-hold_jid job_identifier_list</td>
<td>define jobnet interdependencies</td>
</tr>
<tr>
<td>-hold_jid_ad job_identifier_list</td>
<td>define jobnet array interdependencies</td>
</tr>
<tr>
<td>-i file_list</td>
<td>specify standard input stream file(s)</td>
</tr>
<tr>
<td>-j y[es]</td>
<td>n[oe]</td>
</tr>
<tr>
<td>-jc jc_name</td>
<td>derive job from the specified job class</td>
</tr>
<tr>
<td>-js job_share</td>
<td>share tree or functional job share</td>
</tr>
<tr>
<td>-jsv jsv_url</td>
<td>job submission verification script to be used</td>
</tr>
<tr>
<td>-l resource_list</td>
<td>request the given resources</td>
</tr>
<tr>
<td>-m mail_options</td>
<td>define mail notification events</td>
</tr>
<tr>
<td>-mods list_attr_name attr_key attr_value</td>
<td>modify sublist parameter</td>
</tr>
<tr>
<td>-masterl resource_list</td>
<td>request the given resources for the master task of a parallel job</td>
</tr>
<tr>
<td>-mbind wc_queue_list</td>
<td>bind master task to queue(s)</td>
</tr>
<tr>
<td>-notify</td>
<td>notify job before killing/suspending it</td>
</tr>
<tr>
<td>-now y[es]</td>
<td>n[oe]</td>
</tr>
<tr>
<td>-M mail_list</td>
<td>notify these e-mail addresses</td>
</tr>
<tr>
<td>-N name</td>
<td>specify job name</td>
</tr>
<tr>
<td>-o path_list</td>
<td>specify standard output stream path(s)</td>
</tr>
<tr>
<td>-p project_name</td>
<td>set job’s project</td>
</tr>
<tr>
<td>-p priority</td>
<td>define job’s relative priority</td>
</tr>
<tr>
<td>-par pe-allocation-rule</td>
<td>request overwriting pe allocation rule</td>
</tr>
<tr>
<td>-pe pe-name slot_range</td>
<td>request slot range for parallel jobs</td>
</tr>
<tr>
<td>-pty y[es]</td>
<td>n[oe]</td>
</tr>
<tr>
<td>-q wc_queue_list</td>
<td>bind job to queue(s)</td>
</tr>
<tr>
<td>-R y[es]</td>
<td>n[oe]</td>
</tr>
<tr>
<td>-r y[es]</td>
<td>n[oe]</td>
</tr>
<tr>
<td>-rou reporting_variable_list</td>
<td>write online usage of job to the reporting database</td>
</tr>
<tr>
<td>-rdi y[es]</td>
<td>n[oe]</td>
</tr>
</tbody>
</table>

...continued
### Table 6.1.3: GE Job Script Options...continued

<table>
<thead>
<tr>
<th>Option and parameter</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>-sc context_list</td>
<td>set job context (replaces old context)</td>
</tr>
<tr>
<td>-shell y[n]</td>
<td>start command with or without wrapping &lt;loginshell&gt; -c</td>
</tr>
<tr>
<td>-si session_id</td>
<td>execute GE requests as part of the specified session</td>
</tr>
<tr>
<td>-soft</td>
<td>consider following requests as soft</td>
</tr>
<tr>
<td>-sync sync_options</td>
<td>define sync notification events</td>
</tr>
<tr>
<td>-S path_list</td>
<td>command interpreter to be used</td>
</tr>
<tr>
<td>-t task_id_range</td>
<td>create a job-array with these tasks</td>
</tr>
<tr>
<td>-tc max_running_tasks</td>
<td>throttle the number of concurrent tasks</td>
</tr>
<tr>
<td>-tcon y[n]</td>
<td>run all tasks concurrently</td>
</tr>
<tr>
<td>-terse</td>
<td>tersed output, print only the job-id</td>
</tr>
<tr>
<td>-umask mask</td>
<td>set umask of the job</td>
</tr>
<tr>
<td>-v variable_list</td>
<td>export these environment variables</td>
</tr>
<tr>
<td>-verify</td>
<td>do not submit, just verify</td>
</tr>
<tr>
<td>-V</td>
<td>export all environment variables</td>
</tr>
<tr>
<td>-w e</td>
<td>w</td>
</tr>
<tr>
<td>-wd working_directory</td>
<td>use working_directory</td>
</tr>
<tr>
<td>-xdv docker_volumes</td>
<td>specify docker volume(s)</td>
</tr>
<tr>
<td>-xd docker_options</td>
<td>specify docker run options</td>
</tr>
<tr>
<td>-x --help</td>
<td>show a list of available docker run options</td>
</tr>
<tr>
<td>-xd_run_as_image_user y[n]</td>
<td>start autostart Docker job as the user defined in the Docker image</td>
</tr>
<tr>
<td>-@ file</td>
<td>read commandline input from file</td>
</tr>
</tbody>
</table>

More detail on these options and their use is found in the man page for qsub.

#### 6.1.4 The Executable Line

In a job script, the executable line is launched with the job launcher command after the directives lines have been dealt with, and after any other shell commands have been carried out to set up the execution environment.

**Using `mpirun` In The Executable Line**

The `mpirun` job-launcher command is used for executables compiled with MPI libraries. Executables that have not been compiled with MPI libraries, or which are launched without any specified number of nodes, run on a single free node chosen by the workload manager.

The executable line to run a program `myprog` that has been compiled with MPI libraries is run by placing the job-launcher command `mpirun` before it as follows:

```
[fred@basecm10 ~]$ mpirun myprog
```

#### 6.1.5 Job Script Examples

Some job script examples are given in this section. Each job script can use a number of variables and directives.
Single Node Example Script

An example script for GE:

```
[fred@basecm10 ~]$ cat application
#!/bin/sh
# Give the job a name
#$ -N sleep
# Use this interpreter
#$ -S /bin/sh
# Make sure that the .e and .o file arrive in the
# working directory
#$ -cwd
# Merge the standard out and standard error to one file
#$ -j y
sleep 60
echo Now it is: `date`
```

Parallel Example Script

For parallel jobs the -pe (parallel environment) option must be assigned to the script. Depending on the interconnect, there may be a choice between a number of parallel environments such as MPICH (Ethernet) or MVAPICH2 (InfiniBand).

```
#!/bin/sh
#
# Your job name
#$ -N My_Job
#
# Use current working directory
#$ -cwd
#
# Join stdout and stderr
#$ -j y
#
# pe (Parallel environment) request. Set your number of requested slots here.
# qconf -spl shows current PE list
#$ -pe mpich2 2
#
# Run job through bash shell
#$ -S /bin/bash
#
# If modules are needed, source modules environment:
# . /etc/profile.d/modules.sh
#
# Add any modules you might require:
module add gcc openmpi/gcc/64/1.10.7
#
# The following output will show in the output file. Used for debugging.
echo "Got $NSLOTS processors."
echo "Machines:"
cat machines
#
# Use MPIRUN to run the application. application must be compiled with right MPI library.
# That is openmpi here, compiled with:
# mpicc hello.c -o hello
```
mpirun ./hello

The number of available slots can be set by the administrator to an arbitrary value. However, it is typically set so that it matches the number of cores in the cluster, for efficiency reasons. More slots can be set than cores, but most administrators prefer not to do that.

In a job script, the user can request slots from the available slots. Requesting multiple slots therefore typically means requesting multiple cores. In the case of an environment that is not set up as a parallel environment, the request for slots is done with the -np option. For jobs that run in a parallel environment, the -pe option is used. Mixed jobs, running in both non-MPI and parallel environments are also possible if the administrator has allowed it in the complex attribute slots settings.

Whether the request is granted is decided by the workload manager policy set by the administrator. If the request exceeds the number of available slots, then the request is not granted.

If the administrator has configured the cluster to use cloud computing with Auto Scale (cm-scale) (Chapter 8 of the Administrator Manual), then the total number of slots available to a cluster changes over time automatically, as nodes are started up and stopped dynamically.

### 6.2 Submitting A Job

The GE module uge must be loaded first so that UGE commands can be accessed:

```
$ module add shared uge
```

With GE a job can be submitted with `qsub`. The `qsub` command has the following syntax:

```
qsub [ options ] [ jobscript | -- [ jobscript args ]]
```

After completion (either successful or not), output is put in the user's current directory, appended with the job number which is assigned by GE to the job name.

For example, for a jobscript with the job option `#$ -N myapp`, the job name is set to `myapp`. When the job completes, then by default there is an error and there is an output file. The names they take are of the form:

- `myapp.e<job ID>`
- `myapp.o<job ID>`

If the `-j y` job script option is set, then the error file is combined with the output file into a single `.o<job ID>` file.

#### 6.2.1 Submitting To A Specific Queue

Some clusters have specific queues for jobs which run are configured to house a certain type of job: long and short duration jobs, high resource jobs, or a queue for a specific type of node.

To see which queues are available on the cluster the `qstat` command can be used:

```
$ qstat -g c
```

```
+----------------+-------------+-------------+----------------+----------------+----------+----------------+----------------+
| CLUSTER QUEUE  | CLOAD | USED | RES | AVAIL | TOTAL  | aoACDS | cmuE |
|----------------+-------------+-------------+----------------+----------------+----------+----------------+----------------+|
| long.q         | 0.01     | 0   | 0   | 144   | 288    | 0       | 144            |
| default.q      | 0.01     | 0   | 0   | 144   | 288    | 0       | 144            |
+----------------+-------------+-------------+----------------+----------------+----------+----------------+----------------+|
```

The job is then submitted, for example to the `long.q` queue:

```
$ qsub -q long.q sleeper.sh
```

#### 6.2.2 Queue Assignment Required For Auto Scale (cm-scale)

If cm-scale is used with GE, then jobs must be assigned to a queue by default, or cm-scale ignores the job. If the cluster administrator has not configured GE to assign a queue to jobs by default, then the user can assign a queue to the job with the `-q` option in order to have cm-scale consider the job.
6.3 Monitoring A Job

The job status can be viewed with `qstat`. In this example the `sleeper.sh` script has been submitted. Using `qstat` without options will only display a list of jobs, with no queue status options:

```bash
fred@basecm10 ~]$ qstat
job-ID prior name user state submit/start at queue jclass slots ja-task-ID
---------------------------------------------------------------------------------------
  80  0.55500 myapp fred r 03/12/2020 23:02:20 all.q@node003 2
  81  0.55500 myapp fred qw 03/12/2020 23:02:06  2
  82  0.55500 myapp fred qw 03/12/2020 23:02:07  2
  83  0.55500 myapp fred qw 03/12/2020 23:02:07  2
```

More details are visible when using the `-f` (for full) option. Some of the details that can be seen are:

- The queue-type `qtype`, which can take a state of Batch (B), Interactive (I), or Parallel (P).
- The `resv/used/tot` column is the count of reserved/used/total free slots in the queue.
- The `states` column is the state of the queue.

```bash
fred@basecm10 ~]$ qstat -f
queuename qtype resv/used/tot. np_load arch states
---------------------------------------------------------------------------------
all.q@node001.cm.cluster BIP 0/1/1 0.01 lx-amd64
  88  0.55500 myapp fred r 03/12/2020 23:11:35  1
---------------------------------------------------------------------------------
all.q@node002.cm.cluster BIP 0/0/1 0.01 lx-amd64
---------------------------------------------------------------------------------
all.q@node003.cm.cluster BIP 0/1/1 0.00 lx-amd64
  88  0.55500 myapp fred r 03/12/2020 23:11:35  1
---------------------------------------------------------------------------------
```

The man page for `qstat` gives details on the job states and the queue states.

By default the `qstat` command shows only jobs belonging to the current user, i.e. the command is executed with the option `-u $user`. To see jobs from other users too, the following format is used:

```bash
fred@basecm10 ~]$ qstat -u "*"
```

6.4 Deleting A Job

A job can be deleted in GE with the following command

```bash
fred@basecm10 ~]$ qdel <jobid>
```

The job-id is the number assigned by GE when the job is submitted using `qsub`. Only jobs belonging to the logged-in user can be deleted. Using `qdel` will delete a user’s job regardless of whether the job is running or in the queue.
PBS Professional And OpenPBS

NVIDIA Base Command Manager works with PBS Professional and OpenPBS, which are the modern editions of the original Portable Batch System (PBS) software. The original PBS software was a workload management and job scheduling system to manage computing resources, and was originally developed at NASA in the 1990s.

In this manual, PBS is used as a shorter form to refer to both the modern editions.

PBS job scripts are used to submit and execute jobs. The user puts values into a job script for the resources being requested, such as the number of processors to be used, the memory to be used, or number of nodes required. Other values are also set for the runtime parameters and application-specific variables.

The steps for running a job through a PBS job script are:

- Creating an application to be run via the job script
- Creating the job script, adding directives, applications, runtime parameters, and application-specific variables to the script
- Submitting the script to the workload management system

This chapter covers the using the workload managers and job scripts with the PBS variants so that users can get a basic understanding of how they are used, and can get started with typical cluster usage.

In this chapter:

- section 7.1 covers the components of a job script and job script examples
- section 7.2.1 covers submitting, monitoring, and deleting a job with a job script

More on using PBS is to be found in the PBS Professional Guides, which can be accessed from: https://community.altair.com/community?id=altair_product_documentation.

7.1 Components Of A Job Script

To use PBS, a batch job script is created by the user. The job script is a shell script containing the set of commands that the user wants to run. It also contains the resource requirement directives and other specifications for the job. After preparation, the job script is submitted to the workload manager using the qsub command. The workload manager then tries to make the job run according to the job script specifications.

A job script can be resubmitted with different parameters (e.g. different sets of data or variables).

7.1.1 Sample Script Structure

A job script in PBS has a structure illustrated by the following basic example:

Example
#!/bin/bash
#
#PBS -l walltime=1:00:00
#PBS -l select=2:ncpus=1:mem=500mb
#PBS -j oe

cd ${HOME}/myprogs
mpirun myprog a b c

The first line is the standard “shebang” line used for scripts. The lines that start with #PBS are PBS
directive lines, described shortly in section 7.1.2. The last two lines are an example of setting any
remaining options or configuration settings up, so that the script can run. In this case, a change to the
directory myprogs is made, and then the executable myprog is run, with arguments a b c. The line that
runs the program is called the executable line (section 7.1.3).

To run the executable file in the executable line in parallel, the job launcher mpirun is placed imme-
diately before the executable file. The number of nodes the parallel job is to run on is assumed to have
been specified in the PBS directives.

7.1.2 Directives

Job Script Directives And qsub Options

A job script typically has several configurable values called job script directives, set with job script
directive lines. These are lines that start with a “#PBS”. Any directive lines beyond the first executable
line are ignored.

The lines are comments as far as the shell is concerned because they start with a “#”. However, at
the same time the lines are special commands when the job script is processed by the qsub command.
The difference is illustrated by the following:

- The following shell comment is only a comment for a job script processed by qsub:

  # PBS

- The following shell comment is also a job script directive when processed by qsub:

  #PBS

Job script directive lines with the #PBS part removed are the same as options applied to the qsub
command, so a look at the man pages of qsub describes the possible directives and how they are used. If
there is both a job script directive and a qsub command option set for the same item, then the qsub
option takes precedence.

Since the job script file is a shell script, the shell interpreter used can be changed to another shell
interpreter by modifying the first line (the “#!” line) to the preferred shell. Any shell specified by the
first line can also be overridden by using the “#PBS -S” directive to set the shell path.

Walltime Directive

The workload manager typically has default walltime limits per queue with a value limit set by the
administrator. The user can set a walltime limit for the job by setting the “#PBS -l walltime” directive
to a specific time. The time specified is the maximum time that the user expects the job should run for,
and it allows the workload manager to work out an optimum time to run the job. The job can then run
sooner than it would by default.

If the walltime limit is exceeded by a job, then the job is stopped, and an error message in the follow-
ing format is displayed:
Here, \textit{running time} is the time that the job actually took to run after it ran, while \textit{set time} is the time that the user set as the walltime resource limit.

If there is no default walltime set by the cluster administrator, and if the user submits a job with no walltime set too, then the job only runs when all jobs with a walltime have made room. So to avoid waiting a very long time, and to experience a reasonably consistent behavior on different clusters, the user really should set a walltime.

### Resource List Directives

Resource list directives specify arguments to the \texttt{-l} directive of the job script, and allow users to specify values to use instead of the system defaults.

For example, in the sample script structure earlier, a job walltime of one hour and a memory space of at least 500MB are requested (the script requires the size of the space be spelled in lower case, so “500mb” is used).

If a requested resource list value exceeds what is available, the job is queued until resources become available.

For example, if nodes only have 2000MB to spare and 4000MB is requested, then the job is queued indefinitely, and it is up to the user to fix the problem.

Resource list directives also allow, for example, the number of nodes (-l select=2:) and the number of processor cores for each node, \texttt{ncpus}, to be specified. If no value is specified, then the default is 1 node, and 1 core per node (\texttt{ncpus=1}).

As an aside: To avoid specifying \texttt{ncpus} with a directive, the default value for \texttt{ncpu} can be configured by the cluster administrator with the PBS \texttt{qmgr} configuration setting (section 7.11.1 of the \textit{Administrator Manual}):

**Example**

\texttt{qmgr -c "set server resources.default.ncpus = 4"}

A directive for a resource such as the wall time applies to the entire job. In PBS, such a resource is called a \textit{job-wide} resource.

A collection of resources allocated to a job, but with the resources being requested from the same physical compute node, is called a \textit{chunk}. A directive can be applied at chunk level with the \texttt{select=chunk name} option. Typically, an MPI job has one chunk per MPI process (rank).

Thus, requested resources can be job-wide only (for example, \texttt{walltime}), or chunk only (for example, \texttt{ncpus}). In a job, a resource cannot be used as chunk as well as job-wide—it can only be used as chunk (one or multiple chunks), or job-wide.

So, to run a job on 8 cores, the job-wide specification could be done with:

\texttt{#PBS -l select=8:ncpus=1}

The preceding specification requests 8 CPU cores, and the cores can be anywhere on the cluster.

To run a job on one chunk, the chunk specification could be done with:

\texttt{#PBS -l select=1:ncpus=8}

The preceding specification requests 8 CPU cores, and the cores must be on the same physical node.

Further examples of node resource specification are given in a table on page 61.

### Job Directives: Job Name, Logs, And IDs

If the name of the job script file is \texttt{jobname}, then by default the output and error streams are logged to \texttt{jobname.o<number>} and \texttt{jobname.e<number>} respectively, where \texttt{<number>} indicates the associated job number. The default paths for the logs can be changed by using the \texttt{-o} and \texttt{-e} directives respectively, while the base name (\texttt{jobname} here) can be changed using the \texttt{-N} directive.
Often, a user may simply merge both logs together into one of the two streams using the -j directive. Thus, in the preceding example, "-j oe" merges the logs to the output log path, while "-j eo" would merge it to error log path.

The job ID is an identifier based on the job number and the FQDN of the login node. For a login node called basecm10.cm.cluster, the job ID for a job number with the associated value <number> from earlier, would by default be <number>.basecm10.cm.cluster, but it can also simply be abbreviated to <number>.

Job Queues

Sending a job to a particular job queue is sometimes appropriate. An administrator may have set queues up so that some queues are for very long term jobs, or some queues are for users that require GPUs. Submitting a job to a particular queue <destination> is done by using the directive "#PBS -q <destination>".

Directives Summary

Some useful job directives are illustrated in the following table:

<table>
<thead>
<tr>
<th>Directive Description</th>
<th>Specified As</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name the job &lt;jobname&gt;</td>
<td>#PBS -N &lt;jobname&gt;</td>
</tr>
<tr>
<td>Run the job for a maximum runtime of &lt;walltime&gt;</td>
<td>#PBS -l &lt;walltime&gt;</td>
</tr>
<tr>
<td>Run the job for a maximum runtime of 3 hours 10 minutes and 30 seconds</td>
<td>#PBS -l walltime=03:10:30</td>
</tr>
<tr>
<td>Run the job at &lt;time&gt;</td>
<td>#PBS -a &lt;time&gt;</td>
</tr>
<tr>
<td>Set error log name to &lt;jobname.err&gt;</td>
<td>#PBS -e &lt;jobname.err&gt;</td>
</tr>
<tr>
<td>Set output log name to &lt;jobname.log&gt;</td>
<td>#PBS -o &lt;jobname.log&gt;</td>
</tr>
<tr>
<td>Join error messages to output log</td>
<td>#PBS -j eo</td>
</tr>
<tr>
<td>Join output messages to error log</td>
<td>#PBS -j oe</td>
</tr>
<tr>
<td>Mail to <a href="mailto:user@address">user@address</a></td>
<td>#PBS -M <a href="mailto:user@address">user@address</a></td>
</tr>
<tr>
<td>Mail on &lt;event&gt; where &lt;event&gt; takes the value of the letter in the parentheses (a) bort (b) egin (e) nd (n) do not send email</td>
<td>#PBS -m &lt;event&gt;</td>
</tr>
<tr>
<td>Queue is &lt;destination&gt;</td>
<td>#PBS -q &lt;destination&gt;</td>
</tr>
<tr>
<td>Login shell path is &lt;shellpath&gt;</td>
<td>#PBS -S &lt;shellpath&gt;</td>
</tr>
</tbody>
</table>

Almost every qsub sets a job attribute, and has a corresponding PBS directive with the same syntax as the option. The man page for qsub and the man page for pbs_job_attributes explain this in detail.

Resource Request Examples

As PBS evolved, the specification for requesting nodes has changed. The form:

#PBS -l nodes=3

is deprecated, and automatically converted to:

#PBS -l select=3

The deprecated changes are described in detail the man page for pbs_resources, in the section on BACKWARD COMPATIBILITY.
Examples of requests for \texttt{select=} options are shown in the following table:

<table>
<thead>
<tr>
<th>Resource Request Example Description</th>
<th>#PBS -l Specification</th>
</tr>
</thead>
<tbody>
<tr>
<td>8 nodes, anywhere on the cluster</td>
<td>\texttt{select=8}</td>
</tr>
<tr>
<td>2 nodes, 1 processor per node</td>
<td>\texttt{select=2:ncpus=1}</td>
</tr>
<tr>
<td>3 nodes, 8 processors per node</td>
<td>\texttt{select=3:ncpus=8}</td>
</tr>
<tr>
<td>5 nodes, 2 processors per node, and 1 GPU per node</td>
<td>\texttt{select=5:ncpus=2:ngpus=1}</td>
</tr>
<tr>
<td>5 nodes, 2 processors per node, 3 virtual processors for MPI code</td>
<td>\texttt{select=5:ncpus=2:mpiprocs=3}</td>
</tr>
<tr>
<td>5 nodes, 2 processors per node, using any GPU on the nodes</td>
<td>\texttt{select=5:ncpus=2:ngpus=1}</td>
</tr>
<tr>
<td>5 nodes, 2 processors per node, using a GPU with ID 0 from nodes</td>
<td>\texttt{select=5:ncpus=2:gpu_id=0}</td>
</tr>
</tbody>
</table>

Some of the examples illustrate requests for GPU resource usage. GPUs and the CUDA utilities for NVIDIA are introduced in Chapter 8. GPU usage is treated by the workload manager like the attributes of a resource which the cluster administrator will have pre-configured according to local requirements.

For further details on resources, the man page for \texttt{pbs\_resources} can be checked.

### 7.1.3 The Executable Line

In the job script structure (section 7.1.1), the executable line is launched with the job launcher command after the directives lines have been dealt with, and after any other shell commands have been carried out to set up the execution environment.

**Using \texttt{mpirun} In The Executable Line**

The \texttt{mpirun} command is used for executables compiled with MPI libraries. Executables that have not been compiled with MPI libraries, or which are launched without any specified number of nodes, run on a single free node chosen by the workload manager.

The executable line to run a program \texttt{myprog} that has been compiled with MPI libraries is run by placing the job-launcher command \texttt{mpirun} before it as follows:

\texttt{mpirun myprog}

### 7.1.4 Example Batch Submission Scripts

#### Node Availability

The following job script tests which out of 4 nodes requested with \texttt{"-l nodes=} are made available to the job in the workload manager:

**Example**

```bash
#!/bin/bash
#PBS -l walltime=1:00
#PBS -l nodes=4          #--- legacy
#PBS -l select=4
echo -n "I am on: "
hostname;

echo finding ssh-accessible nodes:
for node in $(cat ${PBS_NODEFILE}) ; do
  echo -n "running on: "
  /usr/bin/ssh $node hostname
done
```

The directive specifying `walltime` means the script runs at most for 1 minute. The `$PBS_NODEFILE` array used by the script is created and appended with hosts by the queuing system. The script illustrates how the workload manager generates a `$PBS_NODEFILE` array based on the requested number of nodes, and which can be used in a job script to spawn child processes. When the script is submitted, the output from the log will look like:

I am on: node001
finding ssh-accessible nodes:
running on: node001
running on: node002
running on: node003
running on: node004

This illustrates that the job starts up on a node, and that no more than the number of nodes that were asked for in the resource specification are provided.

The list of all nodes for a cluster can be found using the `pbsnodes` command (section 7.2.6).

**Using InfiniBand**

A sample PBS script for InfiniBand is:

```bash
#!/bin/bash
# ! Sample PBS file
# ! Name of job

#PBS -N MPI

# ! Number of nodes (in this case 8 nodes with 4 CPUs each)
# ! The total number of nodes passed to mpirun will be nodes*ppn
# ! Second entry: Total amount of wall-clock time (true time).
# ! 02:00:00 indicates 02 hours

#PBS -l walltime=02:00:00
#PBS -l select=8:ncpus=4

# ! Mail to user when job terminates or aborts
#PBS -m ae

# If modules are needed by the script, then source modules environment:
. /etc/profile.d/modules.sh

# Add any modules you might require:
module add shared mvapich/gcc pbspro

# ! Full path to application + application name
application="<application>"

# ! Run options for the application
options="<options>"

# ! Work directory
workdir="<work dir>"
```

~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~
### You should not have to change anything below this line ####
### Change the Working Directory

```bash
#!/ change the working directory (default is home directory)

cd $workdir

# echo Running on host $(hostname)
# echo Time is $(date)
# echo Directory is $(pwd)
# echo PBS job ID is $PBS_JOBID
# echo This job runs on the following machines:
# echo $(cat $PBS_NODEFILE | uniq)

mpirun_command="mpirun $application $options"

# Run the parallel MPI executable (nodes*ppn)
# echo Running $mpirun_command
# eval $mpirun_command
```

In the preceding script, no machine file is needed, since it is automatically built by the workload manager and passed on to the `mpirun` parallel job launcher utility. The job is given a unique ID and run in parallel on the nodes based on the resource specification.

#### 7.1.5 Links To PBS Resources

A number of useful links are:


- The man pages for `qsub`, `pbs_resources`, and `pbs_job_attributes`, for setting up job scripts.


#### 7.2 Submitting A Job

##### 7.2.1 Preliminaries: Loading The Modules Environment

To submit a job to the workload management system, the user must ensure that the following environment modules are loaded:

```bash
$ module add shared pbspro
```

Users can pre-load particular environment modules as their default using the “module init*” commands (section 2.3.3).

##### 7.2.2 Using `qsub`

The command `qsub` is used to submit jobs to the workload manager system. The command returns a unique job identifier, which is used to query and control the job and to identify output. The usage format of `qsub` and some useful options are listed here:

**Usage:** `qsub [options] <job script>`
### Option Hint Description

--- ----  -----------
-a at run the job at a certain time
-l list request certain resource(s)
-q queue job is run in this queue
-N name name of job
-S shell shell to run job under
-j join join output and error files

For example, a job script called `mpirun.job` with all the relevant directives set inside the script, may be submitted as follows:

**Example**

```
$ qsub mpirun.job
```

A job may be submitted to a specific queue `testq` as follows:

**Example**

```
$ qsub -q testq mpirun.job
```

The `man` page for `qsub` describes these and other options. The options correspond to PBS directives in job scripts (section 7.1.1). If a particular item is specified by a `qsub` option as well as by a PBS directive, then the `qsub` option takes precedence.

#### 7.2.3 Job Output

By default, the output from the job script `<scriptname>` goes into the current working directory for PBS.

By default, error output is written to `<scriptname>_.e<jobid>` and the application output is written to `<scriptname>_.o<jobid>`, where `<jobid>` is a unique number that the workload manager allocates. Specific output and error files can be set using the `-o` and `-e` options respectively. The error and output files can usefully be concatenated into one file with the `-j oe` or `-j eo` options. More details on this can be found in the `qsub` man page.

#### 7.2.4 Monitoring The Status Of A Job

To use the commands in this section, the appropriate workload manager module must be loaded:

```
$ module add pbspro
```

**qstat Basics**

The main component is `qstat`, which has several options. In this example, the most frequently used options are discussed.

In PBS, the command “`qstat -an`” shows what jobs are currently submitted or running on the queuing system. An example output is:

```
[fred@basecm10 ~]$ qstat -an

basecm10:           Req'd    Req'd    Elap
Job ID  Username  Queue  Jobname  SessID NDS TSK Memory Time S Time
------------------- --------- --------- ------------------- ------ --- --- ------ ---- ----
121.basecm10       noah  workq  pbjob  -- 3 6  --     01:00 Q  --
125.basecm10       noah  workq  pbjob  26615 3 3  --     01:00 R  --
```
The output shows the Job ID, the user who owns the job, the queue, the job name, the session ID for a running job, the number of nodes requested, the number of CPUs or tasks requested, the time requested (-l walltime), the job state (S) and the elapsed time. In this example, one job is seen to be running (R), and one is still queued (Q). The -n parameter causes nodes that are in use by a running job to display at the end of that line.

Possible job states include:

<table>
<thead>
<tr>
<th>Job States</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>E</td>
<td>Job is exiting after having run</td>
</tr>
<tr>
<td>F</td>
<td>Job is finished</td>
</tr>
<tr>
<td>H</td>
<td>Job is held</td>
</tr>
<tr>
<td>Q</td>
<td>job is queued, eligible to run or routed</td>
</tr>
<tr>
<td>R</td>
<td>job is running</td>
</tr>
<tr>
<td>S</td>
<td>job is suspended</td>
</tr>
<tr>
<td>T</td>
<td>job is being moved to new location</td>
</tr>
<tr>
<td>W</td>
<td>job is waiting for its execution time</td>
</tr>
</tbody>
</table>

The command “qstat -q” shows what queues are available. In the following example, there is one job running in the testq queue and 4 are queued.

$ qstat -q

server: master.cm.cluster

<table>
<thead>
<tr>
<th>Queue</th>
<th>Memory</th>
<th>CPU Time</th>
<th>Walltime</th>
<th>Node</th>
<th>Run</th>
<th>Queue</th>
<th>Lm</th>
<th>State</th>
</tr>
</thead>
<tbody>
<tr>
<td>testq</td>
<td>--</td>
<td>23:59:59</td>
<td>--</td>
<td>1</td>
<td>4</td>
<td>--</td>
<td>E</td>
<td>R</td>
</tr>
<tr>
<td>default</td>
<td>--</td>
<td>23:59:59</td>
<td>--</td>
<td>0</td>
<td>0</td>
<td>--</td>
<td>E</td>
<td>R</td>
</tr>
</tbody>
</table>

Viewing Job Details With qstat

With qstat -f the full output of the job is displayed. The output shows what the jobname is, where the error and output files are stored, and various other settings and variables.

$ qstat -f

Job Id: 137.pj-cruncher

Job_Name = pbjob

Job_Owner = noah@pj-cruncher.cm.cluster
resources_used.cput = 00:00:00
resources_used.mem = 5456kb
resources_used.ncpus = 3
resources_used.vmem = 384164kb
resources_used.walltime = 00:00:11
job_state = R

queue = workq

server = pj-cruncher

Checkpoint = u

ctime = Fri Mar 20 18:07:06 2020

Error_Path = pj-cruncher.cm.cluster:/home/noah/pbjob.e137

exec_host = node001/0+node002/0+node003/0

exec_vnode = (node001:ncpus=1)+(node002:ncpus=1)+(node003:ncpus=1)

Hold_Types = n
7.2.5 Deleting A Job
An already submitted job can be deleted using the qdel command:

$ qdel <jobid>

Multiple space-separated job IDs can be specified.

7.2.6 Nodes According To PBS
The nodes that the workload manager knows about can be viewed using the pbsnodes command.

The following output is from a cluster made up of nodes with 4 physical cores, as indicated by the value of 4 for pcpus. If the node is available to run scripts, then its state is free or time-shared. When a node is used exclusively (section 8.5.2) by one script, the state is job-exclusive.

For PBS Professional the display resembles (some output elided):

[noah@pj-cruncher ~]$ pbsnodes -a
node001
    Mom = node001.cm.cluster
    ntype = PBS
    state = free
pcpus = 4
resources_available.arch = linux
resources_available.host = node001
resources_available.mem = 404784kb
...
queue = workq
resv_enable = True
sharing = default_shared
...
node002
Mom = node002.cm.cluster
ntype = PBS
state = free
pcpus = 4
resources_available.arch = linux
resources_available.host = node002
resources_available.mem = 404784kb
...
queue = workq
resv_enable = True
sharing = default_shared
last_state_change_time = Fri Mar 20 18:09:47 2020
last_used_time = Fri Mar 20 18:09:47 2020
...
Using GPUs

GPUs (Graphics Processing Units) are chips that provide specialized parallel processing power. Originally, GPUs were designed to handle graphics processing as part of the video processor, but their ability to handle non-graphics tasks in a similar manner has become important for general computing. GPUs designed for general purpose computing task are commonly called General Purpose GPUs, or GPGPUs.

A GPU is suited for processing an algorithm that naturally breaks down into a process requiring many similar calculations running in parallel. GPUs cores are able to rapidly apply the instruction on multiple data points organized in a 2-D, and more recently, 3-D, image. The image is placed in a framebuffer. In the original chips, the data points held in the framebuffer were intended for output to a display, thereby accelerating image generation.

The similarity between multicore CPU chips and modern GPUs makes it at first sight attractive to use GPUs for general purpose computing. However, the instruction set on GPGPUs is used in a component called the shader pipeline. This is, as the name suggests, to do with a limited set of graphics operations, and so is by its nature rather limited. Using the instruction set for problems unrelated to shader pipeline manipulations requires that the problems being processed map over to a similar manipulation. This works best for algorithms that naturally break down into a process requiring an operation to be applied in the same way on many independent vertices and pixels. In practice, this means that 1-D vector operations are an order of magnitude less efficient on GPUs than operations on triangular matrices.

Modern GPGPU implementations have matured so that they can now sub-divide their resources between independent processes that work on independent data, and they provide programmer-friendlier ways of data transfer between the host and GPU memory.

Physically, one GPU is typically a built-in part of the motherboard of a node or a board in a node, and consists of several hundred processing cores. There are also dedicated standalone units, commonly called GPU Units, consisting of several GPUs in one chassis. Several of these can be assigned to particular nodes, typically via PCI-Express connections, to increase the density of parallelism even further.

NVIDIA Base Command Manager has several tools that can be used to set up and program GPUs for general purpose computations.

8.1 Packages

A number of different GPU-related packages are included in BCM. The CUDA versions supported are listed in section 9.1 of the Installation Manual.

The version implemented depends on how the system administrator has installed and configured CUDA.

8.2 Using CUDA

After installation of the packages, for general usage and compilation it is sufficient to load just the CUDA<version>/toolkit module, where <version> indicates the CUDA version number. At the time of
writing of this section (May 2023), for Rocky Linux version 9.1, the versions available are 11.7, 11.8, 12.0, and 12.1.

The toolkit comes with the necessary tools and the NVIDIA compiler wrapper to compile CUDA C code.

Extensive documentation on how to get started, the various tools, and how to use the CUDA suite is at https://docs.nvidia.com/cuda.

Also available are several other modules related to CUDA:

- cuda12.1/blas: Provides paths and settings for the CUBLAS library.
- cuda12.1/fft: Provides paths and settings for the CUFFT library.

8.3 Using OpenCL

OpenCL functionality is provided with the cuda<version>/toolkit environment module.

Examples of OpenCL code can be found in the $CUDA_SDK/OpenCL directory.

8.4 Compiling Code

Both CUDA and OpenCL involve running code on different platforms:

- host: with one or more CPUs
- device: with one or more CUDA enabled GPUs

Accordingly, both the host and device manage their own memory space, and it is possible to copy data between them. The CUDA and OpenCL Best Practices Guides in the doc directory, provided by the CUDA toolkit package, have more information on how to handle both platforms and their limitations.

The nvcc command by default compiles code and links the objects for both the host system and the GPU. The nvcc command distinguishes between the two and it can hide the details from the developer.

To compile the host code, nvcc will use gcc automatically.

nvcc [options] <inputfile>

A simple example to compile CUDA code to an executable is:

nvcc testcode.cu -o testcode

The most used options are:

- -g or -debug <level>: This generates debuggable code for the host
- -G or -device-debug <level>: This generates debuggable code for the GPU
- -o or -output-file <file>: This creates an executable with the name <file>
- -arch=sm_13: This can be enabled if the CUDA device supports compute capability 1.3, which includes double-precision

If double-precision floating-point is not supported or the flag is not set, warnings such as the following will come up:

warning: Double is not supported. Demoting to float

The nvcc documentation manual, “The CUDA Compiler Driver NVCC” has more information on compiler options.

The CUDA SDK has more programming examples and information accessible from the file $CUDA_SDK/C/Samples.html.

For OpenCL, code compilation can be done by linking against the OpenCL library:

gcc test.c -lOpenCL
g++ test.cpp -lOpenCL
nvcc test.c -lOpenCL
8.5 Available Tools

8.5.1 CUDA gdb
The CUDA debugger can be started using: cuda-gdb. Details of how to use it are available in the "CUDA-GDB (NVIDIA CUDA Debugger)" manual, in the doc directory. It is based on GDB, the GNU Project debugger, and requires the use of the "-g" or "-G" options compiling.

Example
nvcc -g -G testcode.cu -o testcode

8.5.2 The nvidia-smi Utility
The NVIDIA System Management Interface command, nvidia-smi, can be used to allow exclusive access to the GPU. This means only one application can run on a GPU. By default, a GPU will allow multiple running applications.

Syntax:

nvidia-smi [OPTION1 [ARG1]] [OPTION2 [ARG2]] ...

The steps for making a GPU exclusive:

- List GPUs
- Select a GPU
- Lock GPU to a compute mode
- After use, release the GPU

After setting the compute rule on the GPU, the first application that executes on the GPU blocks out any of the others attempting to run. The first application does not have to be from the user that set the exclusivity lock on the GPU.

To list the GPUs, the -L argument can be used:

$ nvidia-smi -L
GPU 0: (05E710DE:068F10DE) Tesla T10 Processor (S/N: 706539258209)
GPU 1: (05E710DE:068F10DE) Tesla T10 Processor (S/N: 2486719292433)

To set the ruleset on the GPU, the -c|--compute-mode option can be used:

$ nvidia-smi -i 0 -c 1

The ruleset may be one of the following:

- 0 - Default mode (multiple applications allowed on the GPU)
- 1 - Exclusive thread mode (only one compute context is allowed to run on the GPU, usable from one thread at a time)
- 2 - Prohibited mode (no compute contexts are allowed to run on the GPU)
- 3 - Exclusive process mode (only one compute context is allowed to run on the GPU, usable from multiple threads at a time)

To check the state of the GPU, the -q|--query option can be used:

$ nvidia-smi -i 0 -q

In this example, GPU0 is locked, and there is a running application using GPU0. A second application attempting to run on this GPU will not be able to run on this GPU.
Using GPUs

$ histogram --device=0
main.cpp(101) : cudaSafeCall() Runtime API error :
o no CUDA-capable device is available.

After use, the GPU can be unlocked to allow multiple users:

nvidia-smi -i 0 -c 0

8.5.3 CUDA Utility Library

CUTIL is a simple utility library designed for use in the CUDA SDK samples. There are 2 parts for CUDA and OpenCL. The locations are:

- $CUDA_SDK/C/lib
- $CUDA_SDK/OpenCL/common/lib

Other applications may also refer to them, and the toolkit libraries have already been pre-configured accordingly. However, they need to be compiled prior to use. Depending on the cluster, this might have already been done.

[fred@demo ~] cd
[fred@demo ~] cp -r $CUDA_SDK
[fred@demo ~] cd $(basename $CUDA_SDK); cd C
[fred@demo C] make
[fred@demo C] cd $(basename $CUDA_SDK); cd OpenCL
[fred@demo OpenCL] make

CUTIL provides functions for:

- parsing command line arguments
- read and writing binary files and PPM format images
- comparing data arrays (typically used for comparing GPU results with CPU results)
- timers
- macros for checking error codes
- checking for shared memory bank conflicts

8.5.4 CUDA “Hello world” Example

A hello world example code using CUDA is:

Example

/*
CUDA example
"Hello World" using shift13, a rot13-like function.
Encoded on CPU, decoded on GPU.

shift13 cycles between 26 normal alphabet characters.
shift13 shifts 13 steps along the normal alphabet characters
So it translates half the alphabet into non-alphabet characters
shift13 is used because it is simpler than rot13 in c
so we can focus on the point

(c) NVIDIA
Taras Shapovalov <tshapovalov@nvidia.com>
*
#include <cuda.h>
#include <stdio.h>

// CUDA kernel definition: undo shift13
__global__ void helloWorld(char* str)
{
    int idx = blockIdx.x * blockDim.x + threadIdx.x;
    str[idx] -= 13;
}

t void checkCudaError(cudaError_t err, const char *msg) {
    if (err != cudaSuccess) {
        fprintf(stderr, "CUDA error: %s: %s\n", msg, cudaGetErrorString(err));
        exit(EXIT_FAILURE);
    }
}

int main(int argc, char **argv) {
    char s[] = "Hello World!"
    printf("String for encode/decode: %s\n", s);

    // CPU shift13
    int len = sizeof(s);
    for (int i = 0; i < len; i++) {
        s[i] += 13;
    }
    printf("String encoded on CPU as: %s\n", s);

    // Allocate memory on the CUDA device
    char *cuda_s;
    checkCudaError(cudaMalloc((void**)&cuda_s, len), "cudaMalloc");

    // Copy the string to the CUDA device
    checkCudaError(cudaMemcpy(cuda_s, s, len, cudaMemcpyHostToDevice), "cudaMemcpy");

    // Set the grid and block sizes (dim3 is a type)
    // and "Hello World!" is 12 characters, say 3x4
    dim3 dimGrid(3);
    dim3 dimBlock(4);

    // Invoke the kernel to undo shift13 in GPU
    helloWorld<<< dimGrid, dimBlock >>>(cuda_s);
    checkCudaError(cudaGetLastError(), "kernel execution");

    // Retrieve the results from the CUDA device
    checkCudaError(cudaMemcpy(s, cuda_s, len, cudaMemcpyDeviceToHost), "cudaMemcpy");

    // Free up the allocated memory on the CUDA device
    checkCudaError(cudaFree(cuda_s), "cudaFree");
    printf("String decoded on GPU as: %s\n", s);
Using GPUs

The hello world code example may be compiled and run on a node with a GPU and CUDA 12.1 with:

```
[fred@node001 ~]$ module load shared
[fred@node001 ~]$ module load cuda12.1
[fred@node001 ~]$ nvcc hello.cu -o hellocuda
[fred@node001 ~]$ ./hellocuda
```

String for encode/decode: Hello World!
String encoded on CPU as: Uryy|-d|yq.
String decoded on GPU as: Hello World!

The number of characters displayed in the encoded string are less than the number in the unencoded string. This is because there are unprintable characters generated by the encoding, because the cipher used is not exactly rot13.

To make it run from a head node via a workload manager such as Slurm, on a compute node with a GPU, the following batch file could be built and run:

```
[fred@basecm10 ~]$ cat helloslurm.sh
#!/bin/sh
#SBATCH -o my.stdout
#SBATCH --ntasks-per-node=1
#SBATCH -p defq #assuming node in defq has a GPU
#SBATCH --gpus=1
module clear -f
./hellocuda
```

```
[fred@basecm10 ~]$ sbatch helloslurm.sh
[fred@basecm10 ~]$ cat my.stdout
String for encode/decode: Hello World!
String encoded on CPU as: Uryy|-d|yq.
String decoded on GPU as: Hello World!
```

8.5.5 OpenACC

OpenACC ([http://www.openacc-standard.org](http://www.openacc-standard.org)) is a new open parallel programming standard aiming at simplifying the programmability of heterogeneous CPU/GPU computing systems. OpenACC allows parallel programmers to provide OpenACC directives to the compiler, identifying which areas of code to accelerate. This frees the programmer from carrying out time-consuming modifications to the original code itself. By pointing out parallelism to the compiler, directives get the compiler to carry out the details of mapping the computation onto the accelerator.

Using OpenACC directives requires a compiler that supports the OpenACC standard.

In the following example, where $\pi$ is calculated, adding the `#pragma` directive is sufficient for the compiler to produce code for the loop that can run on either the GPU or CPU:

```
#include <stdio.h>
#define N 1000000

int main(void) {
    double pi = 0.0f; long i;
    #pragma acc parallel loop reduction(+:pi)
    for (i=0; i<N; i++) {
```
double t = (double)((i+0.5)/N);
pi += 4.0/(1.0+t*t);
}
printf("pi=%16.15f\n", pi/N);
return 0;
}
Using Kubernetes

9.1 Introduction To Kubernetes Running Via NVIDIA Base Command Manager

Kubernetes is a system for managing containerized applications across multiple hosts in a cluster.

- A container is an extremely lightweight virtualized operating system that runs without the unneeded extra emulated hardware components of a regular virtualized operating system.
- A containerized application runs within a container, and it only accesses files, environment variables, and libraries within the container, unless volumes are mounted and used.
- A containerized application provides services to other software or users. Kubernetes thus manages containerized applications as a service, and is aware of the container states and resources used.

Kubernetes provides mechanisms for application deployment, scheduling, updating, maintenance, and scaling. It actively manages the containers to ensure that the state of the cluster continually matches the user’s intentions. The user’s desired state is communicated to the Kubernetes API server, typically in the form of a YAML file. Kubernetes stores the YAML file in Etcd, and ensures it is reflected by the current state of the containers.

This chapter describes how Kubernetes works with BCM, which currently supports Kubernetes 1.27. For details on Kubernetes that are outside the scope of its use with BCM, the official Kubernetes documentation at [https://kubernetes.io/docs/](https://kubernetes.io/docs/) can be consulted.

By default, in BCM the user is given access to containers only via Kubernetes. The administrator can however configure direct access if required. In this chapter, only container access via Kubernetes is described.

The `kubectl` utility is normally used to communicate with Kubernetes, although using the API directly instead of using `kubectl` is also possible. The `kubectl` utility can be used to get information about Kubernetes runtime, creation and management of resources, as well for other tasks. Resources are items such as pods ([https://kubernetes.io/docs/concepts/workloads/pods/](https://kubernetes.io/docs/concepts/workloads/pods/)) and volumes ([https://kubernetes.io/docs/concepts/storage/volumes/](https://kubernetes.io/docs/concepts/storage/volumes/)), that are consumed while containers are in use.

The official Kubernetes documentation has some introductory tutorials, including:
[https://kubernetes.io/docs/tutorials/kubernetes-basics/](https://kubernetes.io/docs/tutorials/kubernetes-basics/)

Familiarity with the concepts in those tutorials is recommended before continuing with the rest of this chapter.

9.2 Kubernetes User Privileges

The privileges (view-only, edit or admin) that a user has depends on the permissions granted to the user by the Kubernetes Administrator. Since BCM version 9.0 (known earlier as Bright Cluster manager 9.0), a custom namespace is created for each user, `<user>`, according to the following format:
This is intended to be a sandbox for the user.

The default setting in NVIDIA Base Command Manager version 9.0 is to have the PodSecurityPolicies (PSP) feature inactive. Users can check with their BCM administrator if that is indeed the case. If PSP is not active, then users have access to the default namespace.

In future versions of BCM, the default namespace will no longer be available for users by default.

If the Administrator enables the PSP features, then permissions to the default namespace are also not added by default for new users. Role bindings to the default namespace for existing users are also removed.

This “restricted” namespace is always created by default, and users are encouraged to use this namespace.

9.3 Kubernetes Quickstarts

This section provides quickstart recipes on some common tasks for an end user using Kubernetes.

Requirements:

- A Kubernetes cluster on BCM. The administrator should have set this up as described in Chapter 4 of the Containerization Manual.

- a dedicated user on the Kubernetes cluster

The simplest way for a user to use a BCM-managed Kubernetes cluster is if the cluster administrator has configured it for use via the head node. A user can then simply connect to the Kubernetes cluster via an ssh connection to the head node, and then load the environment via module load kubernetes. The remainder of this section can then be skipped.

An alternative way for a user to use a BCM-managed Kubernetes cluster avoids going via the head node, and uses a local PC.

9.3.1 Quickstart: Accessing The Kubernetes Dashboard

The Kubernetes Dashboard is a web browser-based way to manage Kubernetes tasks.

Requirements:

- from the local PC it should be possible to access the Kubernetes Dashboard URL at: https://dashboard.<kubernetes cluster name>:30443

If that does not work, then the cluster administrator has probably modified the standard configuration, and should be consulted on how to access the Kubernetes Dashboard.

- If the Dashboard is accessible, then the user needs a token to authenticate with the Kubernetes Dashboard.

Obtaining And Using The Token

A user test was created according to the procedure in section 4.11 of the Containerization Manual. For the user test, a token can be obtained from the head node with the following procedure, using the namespace and username of the user:

Example

```
$ module load kubernetes
$ USERNAME=test
$ NS=${USERNAME}-restricted
$ SECRET_NAME=$(kubectl get serviceaccount -n $NS $USERNAME -o jsonpath='{.secrets[0].name}')
$ kubectl get secret -n $NS $SECRET_NAME -o jsonpath='{.data.token}' | base64 -d ; echo
```
If the username and related namespace are not known to the user, then the cluster administrator should be asked to provide these.

The output of the base64 command is the decoded token. This can be pasted from the terminal into the web browser, at the place where the token is asked for (figure 9.1):

![Kubernetes Dashboard](image)

**Figure 9.1: Kubernetes User Dashboard Authentication, Using Token Copy-Paste From A Terminal**

After a successful authentication, the Dashboard displayed should appear as in figure 9.2:
The user can now deploy a job using the previously-defined default namespace, as in the following example. The task can be submitted as a YAML file, via the Dashboard URL

https://dashboard.<kubernetes cluster name>:30443#!/deploy?namespace=default

The task could be the pi-job.yml job of section 9.3.3. The result—π to 4000 places—can be seen in the log associated with the jobs, accessible from the Dashboard URL

https://dashboard.<kubernetes cluster name>:30443#!/job?namespace=default

9.3.2 Quickstart: Using kubectl From A Local Machine

The advantage of connecting from a local PC is that there is no need to connect to the head node via SSH.

Requirements:
• the local PC should be able to access the Kubernetes API server at https://dashboard.<kubernetes cluster name>:30443. The URL that is actually used is set up by the cluster administrator, who should be contacted for details.

• The local PC should be Linux-based and run on an amd64 architecture.

Steps:

• On the PC, kubectl for Kubernetes 1.27 should be downloaded from the head node <headnode>. It can be downloaded to a directory in the user path, such as /usr/bin

Example

```
$ rsync <username>@<headnode>:/cm/local/apps/kubernetes/current/bin/kubectl \\
<directory in the user path>
```

• The user can make a .kube directory on the PC. The Kubernetes configuration for the user <username> can then be picked up from <headnode>. This includes the keys and the certificates:

Example

```
$ mkdir ~/.kube
$ rsync <username>@<headnode>:.kube/config-<cluster name> ~/.kube/config
```

• <cluster name> must be replaced with the fully qualified domain name of the Kubernetes cluster

• The user can check if kubectl is able to connect to the cluster by running the following commands:

Example

```
$ kubectl cluster-info
$ kubectl get nodes
$ kubectl get all
```

9.3.3 Quickstart: Submitting Batch Jobs With kubectl

A batch job can be created in Kubernetes. It is simply referred to as a “job”. It is basically made up of non-persistent pods that run a one-off task.

A simple job to calculate $\pi$ can be created by building a pi-job.yml file with the following content:

Example

```
apiVersion: batch/v1
kind: Job
metadata:
  name: pi
spec:
  completions: 8
  parallelism: 1
  template:
    metadata:
      name: pi
    spec:
      containers:
        - name: pi
          image: perl
          command: ["perl", "-Mbignum=bpi", "-wle", "print bpi(4000)"]
          restartPolicy: Never
```
This carries out a Perl-based calculation of π to 4000 places. Running it from the command line in Bash directly could be done with:

**Example**

```bash
$ perl -Mbignum=bpi -wle "print bpi(4000)"
```

However, the idea here is to demonstrate Kubernetes batch jobs firing up and scaling pods for this task instead, which is described next:

If the administrator has allowed the user access via Kubernetes policies, and has made the user a Kubernetes user, then the job can be submitted with:

```bash
$ kubectl apply -f pi-job.yml
job "pi" created
```

If the job is horizontally scalable, then the number of replicas can be scaled with:

```bash
$ kubectl scale job/pi --replicas=4
job "pi" scaled
```

Information about the job can be obtained with (output truncated):

```bash
$ kubectl get job/pi
NAME  DESIRED  SUCCESSFUL  AGE
pi    8        8           6m
```

```bash
$ kubectl describe job/pi
Name: pi
Namespace: default
...
```

The jobs can be followed with (output truncated):

```bash
$ kubectl get pods -aw
NAME  READY STATUS   RESTARTS AGE
pi-74gnd 0/1 Completed 0 6m
...
```

The logs of a pod can be viewed with:

```bash
$ kubectl logs -f <pod name>
```

An output is shown that starts with:

```
3.141592653589793238462643383279502884197169399375105820974...
```

Further information on the following job topics can be found at the associated links:

- job: https://kubernetes.io/docs/concepts/workloads/controllers/job/
- job parallelism: https://kubernetes.io/docs/tasks/job/parallel-processing-expansion/

### 9.3.4 Quickstart: Persistent Storage For Kubernetes Using Ceph

Pods running on Kubernetes can use a distributed storage system such as Ceph for persistent data storage.

The following are assumed:

- a working Ceph cluster
- a Kubernetes StorageClass for Ceph RBD
A `redis-persistent-storage.yml` file can be created for the Persistent Volume Claim (PVC), with the following content:

Example

```yaml
apiVersion: v1
classKind: PersistentVolumeClaim
metadata:
  name: redis-persistent-storage
spec:
  accessModes:
  - ReadWriteOnce
  resources:
    requests:
      storage: 1Gi
    storageClassName: fast
apiVersion: extensions/v1beta1
kind: Deployment
metadata:
  name: redis-master
spec:
  replicas: 1
  template:
    metadata:
      labels:
        app: redis
        role: master
    spec:
      containers:
      - name: redis-master
        image: gcr.io/google_containers/redis:e2e
        args: ['/usr/local/bin/redis-server', '--appendonly', 'yes', '--appendfsync', 'always']
      resources:
        requests:
          cpu: 100m
          memory: 100Mi
      ports:
      - containerPort: 6379
      volumeMounts:
      - name: redis-storage
        mountPath: /data
      volumes:
      - name: redis-storage
        persistentVolumeClaim:
          claimName: redis-persistent-storage
```

The Redis master deployment with PVC can then be created on Ceph persistent storage using:

```
$ kubectl apply -f redis-persistent-storage.yml
deployment "redis-master" created
```

Data can now be stored on Redis. After storing data on Redis, if all the pods are then deleted, they can be recreated without any data loss.
Helm is installed and properly configured by default as a Kubernetes add-on. It is initialized for every Kubernetes user when the Kubernetes module is loaded—there is no `helm init` or similar that needs to be carried out first. For example (some text elided):

**Example**

```bash
$ module load kubernetes
$ helm version
version.BuildInfo{Version:"v3.0.0-beta.4", GitCommit:"7ffcd879f137bd3a69eea53349b01f05e3d1d2385",
GitTreeState:"dirty"}
```

Choices can be made from among the charts at the official repository at https://github.com/kubernetes/charts. For example, GitLab and WordPress can be installed with:

```bash
$ helm install stable/gitlab --name my-gitlab
$ helm install stable/wordpress --name my-wordpress
```

A tutorial on using Helm is available at https://docs.helm.sh/using_helm/#using-helm
Apache Spark is “a lightning-fast unified analytics engine for big data and machine learning”.

Since NVIDIA Base Command Manager version 9.0, the recommended way to run Spark workloads inside BCM is within Kubernetes.

Documentation for Spark is available at https://spark.apache.org/docs/.

10.1 Important Requirements

By default only the root user of the cluster can access Kubernetes. Since that is not very useful, the cluster administrator can grant access to regular users by using cm-kubernetes-setup with the --add-user flag.

The regular user should check that Kubernetes can be accessed via the user’s account (sections 9.2–9.3), or Spark workloads will fail to run. Being able to load the kubernetes module and being able to run the kubectl commands suggests that Kubernetes is properly accessible to the user:

Example

```
[test@cluster ~]$ module load kubernetes/default
[test@cluster ~]$ kubectl get all
```

10.2 Running Spark Jobs Via The Kubernetes Spark Operator

Spark jobs can be run within Kubernetes if the cluster administrator has installed the cm-kubernetes-spark-operator package. The installation can be done

- as part of cm-kubernetes-setup (page 31 of the Containerization Manual) or
- as part of the Base View Kubernetes Wizard using the clickpath:
  Containers→Kubernetes→Kubernetes Wizard.

The following session assumes that the cluster administrator has configured Spark to run as a Kubernetes operator (Chapter 6 of the Containerization Manual). In the session, a user alice carries out a Spark way of calculating pi. This can be regarded as a "Hello world!" type of demonstration for Spark users.

10.2.1 Example Spark Operator Run: Calculating Pi

For alice, a YAML file based on the specification at https://github.com/GoogleCloudPlatform/spark-on-k8s-operator/blob/master/examples/spark-py-pi.yaml can be used:

Example
$ module load kubernetes
$ cat <<EOF > pi-spark.yaml
apiVersion: "sparkoperator.k8s.io/v1beta2"
kind: SparkApplication
metadata:
  name: pyspark-pi
spec:
  type: Python
  pythonVersion: "3"
  mode: cluster
  image: "gcr.io/spark-operator/spark-py:v3.1.1"
  imagePullPolicy: Always
  mainApplicationFile: local:///opt/spark/examples/src/main/python/pi.py
  sparkVersion: "3.1.1"
  restartPolicy:
  type: OnFailure
  onFailureRetries: 3
  onFailureRetryInterval: 10
  onSubmissionFailureRetries: 5
  onSubmissionFailureRetryInterval: 20
  driver:
    cores: 1
    coreLimit: "1200m"
    memory: "512m"
    labels:
      version: 3.1.1
      serviceAccount: spark
  executor:
    cores: 1
    instances: 1
    memory: "512m"
    labels:
      version: 3.1.1
EOF
$ kubectl apply -f pi-spark.yaml
sparkapplication.sparkoperator.k8s.io/pyspark-pi created
$ kubectl get pods
NAME     READY   STATUS      RESTARTS   AGE
pyspark-pi-driver 0/1   ContainerCreating 0 1s
$ kubectl get pods
NAME     READY   STATUS      RESTARTS   AGE
pyspark-pi-driver 1/1   Running   0 3s
$ kubectl get sparkapplications
NAME   AGE
pyspark-pi 7s
$ kubectl get pods
NAME     READY   STATUS      RESTARTS   AGE
pyspark-pi-driver 1/1   Running   0 14s
pythonpi-e768128383a881b3-exec-1 0/1   ContainerCreating 0 0s
$ kubectl get pods
NAME     READY   STATUS      RESTARTS   AGE
pyspark-pi-driver 0/1   Completed 0 34s
pythonpi-e768128383a881b3-exec-1 0/1   Terminating 0 20s
$ kubectl get pods
NAME     READY   STATUS      RESTARTS   AGE
Instead of tracking the pod with:

```
kubectl get pods
```
as in the preceding session, or with the more convenient:

```
watch kubectl get pods
```
the pod could be tracked with the `--follow` option to stream the driver logs:

**Example**

```
[alice@basecm10 ~]$ kubectl logs pyspark-pi-driver -f
```

To get intended output of the pi run—the calculated value of pi—it is sufficient to grep the log as follows:

**Example**

```
[alice@basecm10 ~]$ kubectl logs pyspark-pi-driver | grep 'Pi
Pi is roughly 3.148800
```

After the pi run has completed, the resources can be removed from the namespace:

```
[alice@basecm10 ~]$ kubectl delete -f pi-spark.yaml
sparkapplication.sparkoperator.k8s.io "pyspark-pi" deleted
```

10.3 Running Spark Jobs Directly Via `spark-submit`

The Spark documentation covers a pi run via `spark-submit`. The BCM knowledge base explains how to use `spark-submit` to run a job with older BCM versions (https://kb.brightcomputing.com/knowledge-base/how-to-deploy-spark-with-kubernetes-on-bright-9-0-9-1-9-2/). However, using the Kubernetes operator submission method (section 10.2) is recommended instead in more recent versions.

10.4 Accessing The Spark User Interface

If a job is run via the Kubernetes Operator or `spark-submit`, then the Spark User Interface (Spark UI) can be used to monitor the job. The Spark UI is shut down once the job has finished, so it only makes sense to access the Spark UI for longer-running jobs.

The `kubectl port-forward` command can be used to allow access to the Spark UI:

```
[test@cluster ~]$ module load kubernetes
[test@cluster ~]$ kubectl get pod
NAME               READY STATUS    RESTARTS AGE
spark-pi-1540840660830-driver 0/1 Completed 0 4m34s
spark-pi-1540840938874-driver 0/1 Running 0 2s
[test@cluster ~]$ kubectl port-forward spark-pi-1540840938874-driver 3000:4040
Forwarding from 127.0.0.1:3000 -> 4040
Forwarding from [::1]:3000 -> 4040
Handling connection for 3000
```
The preceding example makes the dashboard available via local port 3000 on the machine where the `port-forward` command is executed. The Spark UI runs on port 4040 inside the pod, and displays something like in figure 10.1:

![Spark Jobs](image)

---

### 10.5 Mounting Volumes Into Containers

The official documentation at [https://spark.apache.org/docs/latest/running-on-kubernetes.html#using-kubernetes-volumes](https://spark.apache.org/docs/latest/running-on-kubernetes.html#using-kubernetes-volumes) gives an outline of how volumes can be mounted into containers.

In summary, there are 3 ways a Kubernetes volume ([https://kubernetes.io/docs/concepts/storage/volumes](https://kubernetes.io/docs/concepts/storage/volumes)) can be mounted:

1. **hostPath**: mounts a file or directory from the host node’s filesystem into a pod.
2. **emptyDir**: an initially empty volume, created when a pod is assigned to a node.
3. **persistentVolumeClaim**: used to mount a PersistentVolume into a pod.

#### Mounting A hostPath

The official documentation mentions adding the following two flags to specify how the volume is mounted inside the container:

```
--conf spark.kubernetes.driver.volumes.[VolumeType].[VolumeName].mount.path=<mount path>
--conf spark.kubernetes.driver.volumes.[VolumeType].[VolumeName].mount.readOnly=<true|false>
```

The following flag should also be added, to specify the path on the host:

```
--conf spark.kubernetes.driver.volumes.[VolumeType].[VolumeName].options.path=<mount path>
```
The volumes can be specified for each executor as well as for the driver. For an executor, the specification should use `spark.kubernetes.executor` instead of `spark.kubernetes.driver`.

For example, the NFS share `/cm/shared` of a typical BCM setup could be made available on executor pods by adding the following 3 flags to the `spark-submit` command:

**Example**

```MODULE
module load kubernetes/default
module load spark
spark-submit \
--master k8s://https://localhost:10443 \
--deploy-mode cluster \
--name spark-pi \
--class org.apache.spark.examples.SparkPi \
--conf spark.kubernetes.namespace=default \
--conf spark.kubernetes.executor.instances=2 \
--conf spark.kubernetes.executor.container.image=docker.io/brightcomputing/spark:2.4.0 \
--conf spark.kubernetes.executor.volumes.hostPath.cmshared.options.path=/cm/shared \
--conf spark.kubernetes.executor.volumes.hostPath.cmshared.mount.path=/data \
--conf spark.kubernetes.executor.volumes.hostPath.cmshared.mount.readOnly=false \
local:///opt/spark/examples/jars/spark-examples_2.11-2.4.0.jar 10000
```

This results in `/cm/shared` being mounted in read-write mode on the mount path `/data`:

```
[root@cluster ~]# kubectl get pod
NAME       READY STATUS              RESTARTS AGE
spark-pi-1552656323319-driver 0/1 Completed 0 23h
spark-pi-1552648760369-driver 0/1 Completed 0 5m40s
spark-pi-1552648961734-driver 0/1 Completed 0 2m26s
spark-pi-1552649048516-driver 1/1 Running   0 59s
spark-pi-1552649048516-exec-1 1/1 Running   0 53s
spark-pi-1552649048516-exec-2 1/1 Running   0 53s
[root@cluster ~]# kubectl exec -it spark-pi-1552649048516-exec-1 /bin/bash
bash-4.4# mount | grep shared
master:/cm/shared on /data type nfs (rw,relatime,vers=3,...)
bash-4.4# ls -l /data
```

**Using Persistent Volume Claims**

Spark in this case assumes that the Kubernetes cluster being managed by the user has persistent volumes available. A list of persistent volumes types can be found at: https://kubernetes.io/docs/concepts/storage/persistent-volumes/#types-of-persistent-volumes.

Claims to specific types of storage can be created for use with Spark. If `spark-submit` tries to use a persistent volume claim, then it assumes the claim already exists. It does not initiate a claim by itself.

In the following specification, for demonstration purposes, a claim `my-claim` is created:

```YAML
apiVersion: v1
kind: PersistentVolumeClaim
```
This claim should bind to a persistent volume that meets the criteria. If none are configured, then the user can create a local volume for `/cm/shared`, for example with:

```
kind: PersistentVolume
apiVersion: v1
metadata:
  name: kube-pv-volume
labels:
  type: local
spec:
  capacity:
    storage: 1000Gi
  accessModes:
  - ReadWriteOnce
  hostPath:
    path: /cm/shared
```

Applying the above two YAML configurations should result in the persistent volume claim object `my-claim`, and the persistent volume `kube-pv-volume` in Kubernetes:

```
[root@cluster ~]# kubectl get pv
NAME CAPACITY ACCESS MODES RECLAIM POLICY STATUS CLAIM STORAGECLASS
kube-pv-volume 1000Gi RWO Retain Bound default/my-claim

[root@cluster ~]# kubectl get pvc
NAME STATUS VOLUME CAPACITY ACCESS MODES STORAGECLASS AGE
my-claim Bound kube-pv-volume 1000Gi RWO 5m35s
```

Example

Invoking `spark-submit` as follows will then try to find the claim with name `my-claim`.

```
module load kubernetes/default
module load spark
spark-submit \
   --master k8s://https://localhost:10443 \n   --deploy-mode cluster \n   --name spark-pi \n   --class org.apache.spark.examples.SparkPi \n   --conf spark.kubernetes.namespace=default \n   --conf spark.kubernetes.executor.instances=2 \n   --conf spark.kubernetes.container.image=docker.io/brightcomputing/spark:2.4.0 \n   --conf spark.kubernetes.authenticate.driver.serviceAccountName=spark \n   --conf spark.kubernetes.executor.volumes.persistentVolumeClaim.cmshared.options.claimName=my-claim \n   --conf spark.kubernetes.executor.volumes.persistentVolumeClaim.cmshared.mount.path=/data \n   --conf spark.kubernetes.executor.volumes.persistentVolumeClaim.cmshared.mount.readOnly=false \n   local:///opt/spark/examples/jars/spark-examples_2.11-2.4.0.jar 10000
```
[root@cluster ~]# kubectl get pod
NAME                  READY STATUS    RESTARTS AGE
spark-pi-1552563523319-driver 0/1 Completed 0 24h
spark-pi-1552648768369-driver 0/1 Completed 0 22m
spark-pi-1552648961734-driver 0/1 Completed 0 18m
spark-pi-1552649048516-driver 0/1 Completed 0 17m
spark-pi-1552649606812-driver 0/1 Completed 0 8m9s
spark-pi-1552650084516-driver 1/1 Running 0 11s
spark-pi-1552650084516-exec-1 1/1 Running 0 4s
spark-pi-1552650084516-exec-2 1/1 Running 0 3s
```

The claim that was specified can be found in the pod description:

```
[root@cluster ~]# kubectl describe pod spark-pi-1552650084516-exec-1 | grep my-claim -C 2
  Type: PersistentVolumeClaim (reference to a PersistentVolumeClaim in the same namespace)
  ClaimName: my-claim
  ReadOnly: false
  default-token-rvrf8:

Inside the executor pods /data is then found to be available, and presents the contents of /cm/shared from the host OS.
Using Singularity

Singularity is a tool that allows applications to be packaged and run in containers. Singularity containers can include a simple binary and library stack, or a complicated work flow. The packaged applications are then called Singularity images. These images are completely portable, with their only dependency requirement being that Singularity must be running on the target system.

NVIDIA Base Command Manager provides the cm-singularity package. This allows users to create container images, which are run in containers. The containers can also be used with MPI applications and with different workload managers. The containers run within user space and are always executed as the run-time user (non-root). The application within the container may have access to the filesystem inside or outside the container. The package also provides a local module file called singularity that must be loaded, before using the singularity command line utility or running a Singularity container.

In this chapter several examples are given of how to create container images and start Singularity containers.

11.1 How To Build A Simple Container Image

The Singularity image is a single file which physically contains the container filesystem. Singularity container images can be bootstrapped with a definition file. The definition file describes how the container is to be built. There are several Singularity keywords (parameters) in the definition file, laid out in lines. If a line does not contain a Singularity keyword, then the line is treated as a shell script line.

The main keywords in the definition file are:

• **BootStrap**: The name of the Linux distribution type module. This informs Singularity which distribution module should be used to parse the commands in the definition file. At present the following 4 modules are supported:
  - **yum**: Bootstraps distributions such as Red Hat, Centos, and Scientific Linux.
  - **debootstrap**: Bootstraps Debian- and Ubuntu-based distributions.
  - **arch**: Bootstraps Arch Linux.
  - **docker**: Bootstraps Docker. It creates a core operating system image based on an image hosted on a particular Docker Registry server. For the docker module, several other keywords may also be defined:
    * **From**: this keyword defines the string of the registry name used for this image in the format [name]:[version].
    * **IncludeCmd**: use the Docker-defined Cmd as the %runscript, if the Cmd is defined,
    * **From**: sets the docker registry name,
    * **Token**: sets the docker authorization token.

• **MirrorURL**: the URL to get packages from.
• **OSVersion**: this keyword must be defined as the alphabet-character string associated with the version of the distribution you wish to use. For example: `trusty` or `stable`.

• **Include**: install additional packages.

• **%setup**: this section blob is a bash scriptlet that is executed on the host outside the container, during bootstrapping.

• **%post**: this scriptlet section is executed once from inside the container, during bootstrapping.

• **%runscript**: the scriptlet that is executed inside the container when it is started.

• **%test**: this section is run at the very end of the bootstrapping process and validates the container during the bootstrap process.

A very simple image definition file for CentOS can look like this:

**Example**

```bash
[root@basecm10 ~]# cat centos7.def
BootStrap: yum
OSVersion: 7
MirrorURL: http://mirror.centos.org/centos-%{OSVERSION}/%{OSVERSION}/os/$basearch/
Include: yum
%post
echo "Installing extra packages..."
yum install vim util-linux -y
%runscript
echo "Hello from container!"
cat /etc/os-release
```

Once the definition file is ready then a user can run the Singularity command `build` to create and install the defined operating system into the container image.

**Example**

```bash
[root@basecm10 ~]# mkdir /cm/shared/sing-images
[root@basecm10 ~]# singularity build /cm/shared/sing-images/centos7.img centos7.def
INFO: Starting build...
INFO: Skipping GPG Key Import
INFO: Running post scriptlet
+ echo 'Installing extra packages...'
Installing extra packages...
+ yum install vim util-linux -y

dependencies resolved and packages installed

INFO: Adding runscript
INFO: Creating SIF file...
INFO: Build complete: /cm/shared/sing-images/centos7.img
[root@basecm10 ~]#
```

The image must be built by a privileged user, even if it is always supposed to be executed with regular user permissions. If, as is the usual case, users are not allowed to use root permissions on a cluster, then they can build the new image on their own computers. The image can then be transferred to the cluster.

The image can be placed in any directory, but it makes sense to share it among compute nodes. The directory `/cm/shared/sing-images` is therefore a sensible location for keeping the container images.

The image created can then be executed as a regular binary:
11.1 How To Build A Simple Container Image

Example

```bash
[user@basecm10 ~]$ module load singularity
[user@basecm10 ~]$ /cm/shared/sing-images/centos7.img
Hello from container!
NAME="CentOS Linux"
VERSION="7 (Core)"
ID="centos"
ID_LIKE="rhel fedora"
VERSION_ID="7"
PRETTY_NAME="CentOS Linux 7 (Core)"
ANSI_COLOR="0;31"
CPE_NAME="cpe:/o:centos:centos:7"
HOME_URL="https://www.centos.org/"
BUG_REPORT_URL="https://bugs.centos.org/"
CENTOS_MANTISBT_PROJECT="CentOS-7"
CENTOS_MANTISBT_PROJECT_VERSION="7"
REDHAT_SUPPORT_PRODUCT="centos"
REDHAT_SUPPORT_PRODUCT_VERSION="7"

The following example shows an image definition file that adds the /etc/services file and /bin/grep binary from the filesystem of the host, to the container image. When a user runs the image that is created, it greps the services file for arguments passed to the image:

Example

[root@basecm10 ~]# cat grep.def
BootStrap: yum
OSVersion: 7
MirrorURL: http://mirror.centos.org/centos-%{OSVERSION}/%{OSVERSION}/os/$basearch/
Include: grep
%setup
  cp /etc/services ${SINGULARITY_ROOTFS}/etc/services
%runscript
  exec /bin/grep $@ /etc/services

[root@basecm10 ~]# singularity build /cm/shared/sing-images/grep.img grep.def
INFO: Starting build...
INFO: Skipping GPG Key Import
INFO: Running setup scriptlet
+ cp /etc/services /tmp/sbuild-102714887/fs/etc/services
INFO: Adding runscript
INFO: Creating SIF file...
INFO: Build complete: /cm/shared/sing-images/grep.img

The container can now be run as if it were a simple script. If a string is passed as an argument, the /etc/services file that is packaged inside the container is searched for the string:

Example

[ user@basecm10 ~]$ /cm/shared/sing-images/grep.img telnets
telnets 992/tcp
telnets 992/udp
[ user@basecm10 ~]$ 
```
11.2 Using MPI

MPI applications can be run from a Singularity container according to two models:

1. the Hybrid model: this relies on both the host and the container to provide MPI implementations for the application that is to run. (This is sometimes also called the Host MPI model).

2. the Bind model: this relies on only the host MPI implementation. The container itself has no MPI implementation, but bind mounts the host MPI implementation into the container, which tends to keep the container lighter.

This is discussed in the upstream Singularity MPI documentation at https://sylabs.io/guides/3.5/user-guide/mpi.html.

BCM administrators typically keep the system software synchronized at the same version across all nodes, and use /cm/shared/ and /home as a shared directory. Containers placed on the nodes therefore usually make use of the shared, synchronized software, which means that in BCM, bind mounting is mostly not needed.

A container can be defined and built on the head node for testing purposes. After adding an MPI implementation, the container can be placed onto the node image, and execution can be carried out via a workload manager such as Slurm.

11.2.1 Running An MPI Example

Singularity at the time of writing (November 2020) supports the two main MPI implementations: OpenMPI (https://www.open-mpi.org/), and MPICH (https://www.mpich.org/).

This section shows a simple example of how an MPICH and OpenMPI run can be carried out.

Container Definitions

**MPICH container definition:** An example of a container definition for MPICH is:

```
[root@basecm10 ~]# cat mpich.def
Bootstrap: yum
OSVersion: 7
MirrorURL: http://mirror.centos.org/centos-%{OSVERSION}/%{OSVERSION}/os/$basearch/
Include: yum
%post
  echo "Installing extra packages..."
  yum install tcl -y
  yum install less -y
  yum install which -y
  yum install cm-config-yum -y
  yum install env-modules -y
  yum install gcc-recent -y
  yum install libgfortran -y
  yum install mpich-ge-gcc-64 -y
%files
  # <source> [destination]
  /etc/yum.repos.d/cm.repo
  /etc/pki/rpm-gpg/RPM-GPG-KEY-cm
%runscript
  #run when container runs
  # takes arguments after container name when container is run
  echo "Hello from container!"
  cat /etc/os-release
  echo "loaded modules are:"
  module list
  echo "available modules are:"
```
module avail
module load gcc
module load mpich/ge/gcc/
%setup
# runs on host, as root
# touch $SINGULARITY_ROOTFS/var/log/yum.log
[root@basecm10 ~]#

The %files section takes care of some binding that is needed for the container to install the extra YUM packages in the %post section.

OpenMPI container definition: Similarly, an example of a container definition for OpenMPI could be:

Bootstrap: yum
OSVersion: 7
MirrorURL: http://mirror.centos.org/centos-%{OSVERSION}/%{OSVERSION}/os/$basearch/
Include: yum
%post
  echo "Installing extra packages..."
  yum install tcl -y
  yum install cm-config-yum -y
  yum install env-modules -y
  # yum install openmpi -y
  yum install openmpi-geib-gcc-64 -y
  module load gcc
  module load openmpi
%files
  # <source> [<destination>]
  /etc/yum.repos.d/cm.repo
  /etc/pki/rpm-gpg/RPM-GPG-KEY-cm
%runscript
  echo "Hello from container!"
  cat /etc/os-release
  echo "loaded modules are:"
  module list
  echo "available modules are:"
  module avail
  echo "repos are:"

Building The Container
If the directory /cm/shared/sing-images was created earlier on, then:

- For MPICH, a Singularity image can then be built by the root user with:

  singularity build /cm/shared/sing-images/mpich.img mpich.def

- Similarly, for OpenMPI, a Singularity image can be built by the root user with:

  singularity build /cm/shared/sing-images/openmpi.img openmpi.def

Execution With A Container On The Head Node For Testing
A hello world binary can be run.

The hello world example from Chapter 3 was:
#include <stdio.h>
#include <mpi.h>

int main(int argc, char *argv[]) {
    int id, np, i;
    char processor_name[MPI_MAX_PROCESSOR_NAME];
    int processor_name_len;
    MPI_Init(&argc, &argv);
    MPI_Comm_size(MPI_COMM_WORLD, &np);
    MPI_Comm_rank(MPI_COMM_WORLD, &id);
    MPI_Get_processor_name(processor_name, &processor_name_len);
    for(i=1;i<2;i++) {
        printf("Hello world from process %03d out of %03d, processor name %s\n", id, np, processor_name);
    }
    MPI_Finalize();
    return 0;
}

This can be compiled by a user (fred in this case), after loading the appropriate MPICH or OpenMPI module:

Example

[fred@basecm10 ~]$ module load mpich/ge/gcc/64/  # or module load openmpi/gcc/64/ as appropriate
[fred@basecm10 ~]$ mpicc hello.c -o hello

For an MPICH implemented by the administrator on the regular nodes, the user can run the binary
with a machine file (containing node001 and node002 here) with:

Example

[fred@basecm10 ~]$ module rm openmpi
[fred@basecm10 ~]$ module load mpich/ge/gcc/64/
[fred@basecm10 ~]$ mpirun --map-by slot -machinefile mpirun.hosts /home/fred/hello
Hello world from process 000 out of 002, processor name node001
Hello world from process 001 out of 002, processor name node002

For an OpenMPI implemented by the administrator, the corresponding run is:

Example

[fred@basecm10 ~]$ module rm mpich/ge/gcc/64/3.3.2
[fred@basecm10 ~]$ module load openmpi/gcc/64/1.10.7
[fred@basecm10 ~]$ /cm/shared/apps/openmpi/gcc/64/1.10.7/bin/mpirun --mca btl tcp,self --map-by slot
    --machinefile mpirun.hosts /home/fred/hello
Hello world from process 001 out of 002, processor name node002
Hello world from process 000 out of 002, processor name node001

The --mca btl tcp,self option avoids warnings being displayed when Ethernet is used for MPI. The
option can be dropped if InfiniBand has been configured by the cluster administrator.

**Running A Binary Via MPI From A Singularity Image**

The MPICH or OpenMPI binary can be run using Singularity with the MPICH or OpenMPI container
image on the head node:
Running a binary via MPI from an MPICH Singularity image:

Example

[fred@basecm10 ~]$ module rm openmpi/gcc/64/1.10.7
[fred@basecm10 ~]$ module load mpich/ge/gcc/64/
[fred@basecm10 ~]$ mpirun -np 4 -machinefile mpirun.hosts singularity exec /cm/shared/sing-images/mpich.img /home/fred/hello
Hello world from process 001 out of 002, processor name node002
Hello world from process 000 out of 002, processor name node001

Running a binary via MPI from an OpenMPI Singularity image:

Example

[fred@basecm10 ~]$ module load openmpi/gcc/64/1.10.7
[fred@basecm10 ~]$/cm/shared/apps/openmpi/gcc/64/1.10.7/bin/mpirun --mca btl tcp,self --map-by slot -machinefile mpirun.hosts singularity exec /cm/shared/sing-images/openmpi.img /home/fred/hello
Hello world from process 001 out of 002, processor name node002
Hello world from process 000 out of 002, processor name node001
[fred@basecm10 ~]$

Execution Via Workload Manager On Regular Nodes With Slurm
If the user fred is meant to submit MPI jobs via a workload manager, such as Slurm, then a standard way to execute them is to create a job script (Chapter 4). With Slurm, srun is generally recommended as an optimal way to run MPI tasks, in preference to mpirun.

A simple job script for the MPICH image is:

Example

[fred@basecm10 ~]$ cat slurm.job
#!/bin/bash
#SBATCH --ntasks=2
module load slurm
module load mpich/ge/gcc
srun singularity exec /cm/shared/sing-images/mpich.img /home/fred/hello
[fred@basecm10 ~]$ module load slurm
[fred@basecm10 ~]$ sbatch slurm.job
Submitted batch job 2
[fred@basecm10 ~]$ cat slurm-2.out
Hello world from process 001 out of 002, processor name node002
Hello world from process 000 out of 002, processor name node001

A similar run with OpenMPI is:

[fred@basecm10 ~]$ cat slurm.job
#!/bin/bash
#SBATCH --ntasks=2
module load slurm
module load openmpi
module load singularity
srun singularity exec /cm/shared/sing-images/openmpi.img /home/fred/hello
[fred@basecm10 ~]$ module load slurm
[fred@basecm10 ~]$ sbatch slurm.job
Submitted batch job 8
[fred@basecm10 ~]$ cat slurm-8.out
Hello world from process 001 out of 002, processor name node002
Hello world from process 000 out of 002, processor name node001
User Portal

The user portal allows users to log in via a browser and view the state of the cluster themselves. The interface does not allow administration, but presents data about the system. The presentation of the data can be adjusted in many cases.

The user portal is accessible at a URL with the format of https://<head node host name, or IP address>:8081/userportal, unless the administrator has changed it.

The first time a browser is used to log in to the cluster portal, a warning about the site certificate being untrusted appears in a default NVIDIA Base Command Manager configuration. This can safely be accepted.

The user portal has several modes.

- The Overview mode is opened by default, and allows a user to access the following pages via links in the left hand column:
  - Overview (section 12.1)
  - Workload (section 12.2)
  - Nodes (section 12.3)
  - OpenStack (section 12.4)
  - Kubernetes (section 12.5)

- The Monitoring mode (section 12.6) allows a user to plot device measurables

- The Accounting and reporting mode (section 12.7) allows a user to plot job-based resource consumption.

12.1 Overview Page

The default Overview page allows a quick glance to convey the most important cluster-related information for users (figure 12.1):
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Figure 12.1: User Portal: Overview Page

The following items are displayed on a default home page:

- a Message Of The Day. The administrator may put up important messages for users here
- links to the documentation for the cluster
- contact information. This typically shows how to contact technical support
- an overview of the cluster state, displaying some cluster parameters

12.2 Workload Page

The Workload page allows a user to see workload-related information for the cluster (figure 12.2). The columns are sortable.

By default, only the cluster administrator can see information for all users. The administrator can adjust the profile for a user to allow that user to view information from other users.
12.3 Nodes Page

The workload jobs are organized in tables according to:

- Queues
- Parent Jobs
- Jobs

12.3 Nodes Page

The Nodes page shows nodes on the cluster (figure 12.3), along with some of their properties. Nodes and their properties are arranged in sortable columns.
The following information about the head or regular nodes is presented:

- **HOSTNAME**: the node name
- **STATE**: For example, UP, DOWN, INSTALLING, along with some other information
- **MEMORY**: RAM on the node
- **CORES**: Number of cores on the node
- **CPU**: Type of CPU, for example, Dual-Core AMD Opteron™
- **SPEED**: Processor speed
- **GPU**: GPUs on the node, if any
- **NICS**: Number of network interface cards on the node, if any
- **IB**: Number of InfiniBand interconnects on the node, if any
- **CATEGORY**: The node category that the node has been allocated by the administrator (by default it is default)

12.4 OpenStack Page

The OpenStack page (figure 12.4) shows an overview of the virtual machines, compute hosts, network nodes, the resources used by these and their properties.
Items shown are:

- **Total VMs up**: total number of virtual machines up
- **Total VMs with errors**: total number of virtual machines that are not running
- **Total projects**: Total number of projects
- **Total networks**: Total number of networks
- **Total subnets**: Total number of subnets
- **Total routers**: Total number of routers. These are usually interconnecting networks
- **Total users**: Total number of users
- **Compute hosts**: Compute hosts
- **Network nodes**: Network nodes
- **Projects**: The projects are listed in sortable columns, by name and UUID,
- **Cloud status text**
- **Dashboard URLs**: URLs to the clusters
- **Default project name**: by default set to `tenant-${username}`, unless the administrator has changed this.
12.5 Kubernetes Page

The Kubernetes page (figure 12.5) shows an overview of the resources available in clusters running Kubernetes.

![Kubernetes Page](image)

Figure 12.5: User Portal: Kubernetes Page

The Kubernetes cluster is subset of a BCM cluster, and is the part of the BCM cluster that runs and controls pods. The items shown are:

- **NAME**: The Kubernetes Cluster name
- **VERSION**: The Kubernetes version
- **NODES**: The number of nodes in the Kubernetes cluster
- **NAMESPACE**: The number of namespaces defined for the Kubernetes cluster
- **SERVICES**: The number of services that are served by the Kubernetes cluster
- **REPLICATION CONTROLLERS**: The number of replication controllers that run on the Kubernetes cluster
- **PERSISTENT VOLUMES**: The number of persistent volumes created for the pods of the Kubernetes cluster
- **PERSISTENT VOLUMES CLAIMS**: The number of persistent volumes claims created on the Kubernetes cluster
12.6 Monitoring Mode

By default the Monitoring mode page displays two empty plot panels within the dashboard section of the page.

The panels can have measurables drag-and-dropped into them from the measurables navigation tree on the left hand side of the page (figure 12.6). The tree can be partly or fully expanded.

A filter can be used to select from visible measurables. For example, after expanding the tree, it is possible to find a measurable related to the cluster occupation rate (Appendix G of the Administrator Manual) by using the key word “occupation” in the filter. The measurable can then be dragged from the options that remain visible.

Extra plot panel widgets can be added to the page by clicking on the Add new widget option (the + button) in the panels section of the dashboard page.

Figure 12.6: User Portal: Monitoring Page Plots

A new dashboard tab can be added to the page by clicking on the Add new dashboard option (the + button) in the dashboard tabs section of the page.

12.7 Accounting And Reporting Mode

The Accounting and reporting mode page allows resource use to be displayed and reported by running PromQL queries. Further background on this can be found in Chapter 14 of the Administrator Manual.

A wizard allows resource reports to be created per user and per account. The reports can be gathered in tabs (figure 12.7).
The browser configuration that generates the reports can be saved, and the report data values can be exported as a download in CSV or Microsoft Excel format (figure 12.8).
Figure 12.8: User Portal: Simple Accounts
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Using Jupyter

13.1 Introduction

Jupyter is a general name given to a system of software components integrated with Jupyter Notebook. Chapter 18 of the Administrator Manual describes Jupyter for NVIDIA Base Command Manager, but is meant for cluster administrators rather than end users.

For the orientation of end users, excerpts from that chapter that give a short overview of the Jupyter components are repeated in this introductory section here. How a user can use the components is described in greater detail in the remaining sections of this chapter.

What Is Jupyter Notebook?

Jupyter Notebook (https://jupyter-notebook.readthedocs.io/), or Jupyter, is a client-server open-source application that provides a convenient way for a cluster user to write and execute notebook documents in an interactive environment.

In Jupyter, a notebook document, or notebook, is content that can be managed by the application. Notebooks are organized in units called cells and can contain both executable code, as well as items that are not meant for execution.

Items not meant for execution can be, for example: explanatory text, figures, formulas, or tables. Notebooks can also store the inputs and outputs of an interactive session.

These documents are encoded as JSON files and saved with the .ipynb extension. Since JSON is a plain text format, notebooks can be version-controlled, shared with other users and exported to other formats, such as HTML, LaTeX, PDF, and slide shows.

What Is A Notebook Kernel?

A notebook kernel (often shortened to kernel) is a computational engine that handles the various types of requests in a notebook (e.g. code execution, code completions, inspection) and provides replies to the user (https://jupyter.readthedocs.io/en/latest/projects/kernels.html). Usually kernels only allow execution of a single language. There are kernels available for many languages, of varying quality and features.

What Is JupyterHub?

Jupyter on its own provides a single user service. JupyterHub (https://jupyterhub.readthedocs.io/) allows Jupyter to provide a multi-user service, and is therefore commonly installed with it. JupyterHub is an open-source project that supports a number of authentication protocols, and can be configured in order to provide access to a subset of users.
What Is JupyterLab?

JupyterLab (https://jupyterlab.readthedocs.io/) is a modern and powerful interface for Jupyter. It enables users to work with notebooks and other applications, such as terminals or file browsers. It is open-source, flexible, integrated, and extensible.

JupyterLab works out of the box with JupyterHub. It can be used to arrange the user interface to support a wide range of workflows in data science, scientific computing, and machine learning.

JupyterLab is extensible with plugins that can customize or enhance any part of the interface. Plugins exist for themes, file editors, keyboard shortcuts, as well as for other components.

What Is A Jupyter Extension?

Several components of the Jupyter environment can be customized in different ways with extensions. Some types of extensions are:


Extensions are usually developed, bundled, released, installed, and enabled in different ways.

Each extension provides a new functionality for a specific component. For example, JupyterLab extensions can customize or enhance any part of the JupyterLab user interface. Extensions can provide new themes, file viewers, editors and renderers for rich output in notebooks. They can also add settings, add keyboard shortcuts, or add items to the menu or command palette.

What Is Jupyter Enterprise Gateway?

By default, Jupyter runs kernels locally, which can exhaust server resources. A resource manager can be used to deal with this issue. Jupyter Enterprise Gateway (https://jupyter-enterprise-gateway.readthedocs.io/) (JEG) is a pluggable open-source framework that leverages local underlying resource managers, such as Slurm or Kubernetes, to distribute kernels across the compute cluster. In addition to scalability, JEG also provides an improved multi-user support, and more granular security for Jupyter. This makes Jupyter more useful for enterprise, scientific, and academic implementations.

In BCM, all the technologies mentioned in these sections are combined to provide a powerful, customizable and user-friendly Jupyter Notebook web interface running on a lightweight, multi-tenant, multi-language, scalable and secure environment, ready for a wide range of enterprise scenarios.

For convenience, in the following sections, Jupyter is generally used to collectively refer to Jupyter Notebook, JupyterHub, JupyterLab and Jupyter Enterprise Gateway.

BCM Jupyter Extensions

For a default deployment of Jupyter, BCM installs and enables the following extensions to the Jupyter environment:

- Jupyter Addons: A Jupyter Notebook server extension that performs API calls to CMDaemon and manages other server extensions;
- Jupyter Kernel Creator (section 13.4): A Jupyter Notebook server extension that provides a new interactive and user-friendly way to create kernels;
- Jupyter VNC (section 13.5): A Jupyter Notebook server extension that enables remote desktops with VNC from notebooks;
13.2 Jupyter Notebook Examples

- JupyterLab Tools: A JupyterLab extension that exposes BCM server extensions functionalities to the users and shows the Cluster View section;

- Jupyter WLM Magic (section 13.6): An IPython extension that simplifies scheduling of workload manager jobs from the notebook;

- Jupyter Kubernetes Operators Manager (section 18.9 of the Administrator Manual): An extension that integrates with Kubernetes clusters, and for which it provides basic overview and management features.

The default setup for the user is described in the next sections. It may however be the case that the cluster administrator has customized Jupyter for the needs of the organization, in which case the description that follows may differ from the real life situation.

For example, if the Jupyter login host is at an IP address of 10.2.75.147 then the URL the user uses to log in from with a browser (figure 13.1) is by default:

https://10.2.75.147:8000

Figure 13.1: JupyterHub User Login Screen

13.2 Jupyter Notebook Examples

The default BCM implementation of Jupyter provides a number of machine learning notebook examples that can be executed with Jupyter.

The notebooks include some applications developed with TensorFlow, PyTorch, MXNet, and other frameworks. The applications can be found in the /cm/shared/examples/jupyter/notebooks/ directory:

```
[jupyterhubuser@basecm10 ]$ ls /cm/shared/examples/jupyter/notebooks/
Keras+TensorFlow2-addition.ipynb R-iris.ipynb TensorFlow-minigo.ipynb
MXNet-superresolution.ipynb Spark-pipeline.ipynb
Pytorch-cartpole.ipynb Spark+XGBoost-mortgage.ipynb
```

The datasets needed to execute these notebooks can be found in the /cm/shared/examples/jupyter/datasets/ directory:
Users can copy these examples to their home directories, create or choose appropriate kernels to execute them, and interactively run them from Jupyter. In order to edit notebooks, the write permissions must be kept during the copy.

It is the responsibility of users to make sure that the required modules are loaded by their Jupyter kernels. The list of frameworks and libraries required to run an example is usually available at the beginning of each notebook.

### 13.3 Jupyter Kernels

In Jupyter, kernels are defined as JSON files.

Any user that the cluster administrator has registered in the Linux-PAM system can list the available Jupyter kernels via the command line. The following example is run in the initial Jupyter environment:

**Example**

```
[jupyterhubuser@basecm10 ~]$ module load jupyter
[jupyterhubuser@basecm10 ~]$ jupyter kernelspec list
Available kernels:
  python3 /cm/shared/apps/jupyter/14.0.0/share/jupyter/kernels/python3
```

Each kernel directory contains a `kernel.json` file describing how Jupyter spawns that kernel:

**Example**

```
[jupyterhubuser@basecm10 ~]$ ls /cm/shared/apps/jupyter/current/share/jupyter/kernels/*/kernel.json
/cm/shared/apps/jupyter/current/share/jupyter/kernels/python3/kernel.json
```

In addition to specifications for shared kernels, each user can define new personal ones in the home directory. By default, the Jupyter data directory for a user is located at `$HOME/.local/share/jupyter`.

This path can be verified with Jupyter by using the `--paths` option:

**Example**

```
[jupyterhubuser@basecm10 ~]$ jupyter --paths
```

```
config:
  /home/jupyterhubuser/.jupyter
  /home/jupyterhubuser/.local/etc/jupyter
  /cm/local/apps/python39/etc/jupyter
  /cm/shared/apps/jupyter/14.0.0/etc/jupyter
  /etc/jupyter
data:
  /home/jupyterhubuser/.local/share/jupyter
  /cm/local/apps/python39/share/jupyter
  /cm/shared/apps/jupyter/14.0.0/share/jupyter
  /usr/share/jupyter
runtime:
  /home/jupyterhubuser/.local/share/jupyter/runtime
```

The simplest definition for a Python3 kernel designed to run on the login node is:

```json
{
  "argv": ["python", 
            "-m", 
            "ipykernel_launcher".
```
In the preceding kernel definition:

- **argv**: is the command to be executed to locally spawn the kernel
- **"display_name"**: is the name to be displayed in the JupyterLab interface
- **"language"**: is the supported programming language ("language")
- **"{connection_file}"** is a placeholder, and is replaced by Jupyter with the actual path to the connection file before starting the kernel.

The following kernel is Jupyter’s default Python 3 kernel distributed by BCM in the initial environment:

**Example**

```
[jupyterhubuser@basecm10 ~]$ cat /cm/shared/apps/jupyter/current/share/jupyter/kernels/python3/kernel.json
{
    "argv": [
        "/cm/local/apps/python37/bin/python3.9",
        "-m",
        "ipykernel_launcher",
        "--InteractiveShellApp.extra_extension=cm_jupyter_wlm_magic",
        "--TerminalIPythonApp.extra_extension=cm_jupyter_wlm_magic",
        "-f",
        "{connection_file}"],
    "display_name": "Python 3",
    "language": "python",
    "env": {
        "PYTHONPATH": "/cm/shared/apps/jupyter/current/lib64/python3.9/site-packages:/cm/shared/apps/jupyter/current/lib/python3.9/site-packages"
    }
}
```

The two kernels are not very different. They differ from each other in the Python 3 binary path, the IPython extension (Jupyter WLM Magic), and the exported `PYTHONPATH` environment variable ("env").

### 13.3.1 Jupyter Enterprise Gateway Kernels

Jupyter is designed to run both the kernel processes, as well as the user interface (JupyterLab or Jupyter Notebook) on the same host. The kernel `{connection_file}` is therefore stored in the `~/local/share/jupyter/runtime` directory, or in the `/run` directory.

JupyterLab can delegate the task of spawning kernels to another component. During the BCM installation it is Jupyter Enterprise Gateway (JEG) that takes on this role, performing a complete lifecycle management of the kernel. That is: start, status monitoring, and termination.

JEG requires an extended `kernel.json` definition to describe a particular process-proxy module to handle the kernel.

A simple definition for a Python3 kernel designed to be scheduled via JEG is:
In this example, the "metadata" entry has been added. It includes "class_name", which defines a path to a Python class compatible with the JEG API; and "config", an initial arguments dictionary passed to the JEG process.

JEG can use several pluggable modules that are included in the BCM environment to interact with a wide range of resource managers, such as Kubernetes or Slurm. By doing so, kernels can be scheduled across compute nodes.

BCM recommends that JEG kernels are created and used with the Jupyter Kernel Creator (section 13.4) extension.

13.4 Jupyter Kernel Creator Extension

Creating or editing kernels can be cumbersome and error-prone for users, depending on the features of the execution context desired for their notebooks.

To provide a more user-friendly experience, BCM includes the Jupyter Kernel Creator extension in JupyterLab. This extension is accessed from the navigation pane in the JupyterLab interface, by clicking on the BCM icon.

Jupyter Kernel Creator allows users to create kernels using the JupyterLab interface, without the need to directly edit JSON files. With this interface users can create kernels by customizing an available template according to their needs.

A template can be considered to be the skeleton of a kernel, with several preconfigured options, and others options that are yet to be specified. Common customizations for templates include environment modules to be loaded, workload manager queues to be used, number and type of GPUs to acquire, and so on.

Templates are usually defined by administrators according to cluster capabilities, programming languages and user requirements. Each template can provide different options for customizations.

Administrators often create different templates to take advantage of different workload managers, programming languages and hardware resources. For example, an administrator may define a template for scheduling Python kernels via Kubernetes, another one for R kernels via Slurm, and yet another one for Bash kernels via Platform LSF.

13.4.1 BCM Predefined Kernel Templates

To simplify Jupyter configuration for administrators, BCM distributes a number of pre-defined templates with Jupyter Kernel Creator. These templates can be used for default configurations of BCM workload managers, and can be customized and extended for more advanced use. Kernel templates defined by BCM can be found in the Jupyter Kernel Creator installation directory, under the kerneltemplates directory:
Filter yaml

<table>
<thead>
<tr>
<th>Template Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>jupyter-eg-kernel-k8s-bash</td>
<td>Bash via OpenPBS</td>
</tr>
<tr>
<td>jupyter-eg-kernel-k8s-r</td>
<td>R via OpenPBS</td>
</tr>
<tr>
<td>jupyter-eg-kernel-k8s-spark</td>
<td>Spark via OpenPBS</td>
</tr>
<tr>
<td>jupyter-eg-kernel-ks-cmjkop-bash</td>
<td>Bash via Kubernetes</td>
</tr>
<tr>
<td>jupyter-eg-kernel-ks-cmjkop-julia</td>
<td>Julia via Kubernetes</td>
</tr>
<tr>
<td>jupyter-eg-kernel-ks-cmjkop-nc-py</td>
<td>Python via Kubernetes</td>
</tr>
<tr>
<td>jupyter-eg-kernel-ks-cmjkop-py</td>
<td>Python via Kubernetes</td>
</tr>
</tbody>
</table>

**BCM Predefined Kernel Templates Seen By Users**

Users can view the available predefined kernel templates in the Jupyter web browser interface, within the **KERNEL TEMPLATES** section of the dedicated BCM extensions panel (figure 13.2):

![Figure 13.2: JupyterLab BCM extensions section with kernel templates](image)

However, the templates provided by BCM are listed in the panel only if they can be used on the cluster. This means that the templates are listed only after the associated workload manager instance, or associated Kubernetes configuration (such as a Kubernetes operator), have been deployed by cluster manager utilities. For example:

- After running the `cm-wlm-setup` cluster manager utility to deploy an OpenPBS workload manager, the `jupyter-eg-kernel-openpbs-bash` and `jupyter-eg-kernel-openpbs-py39` templates become available. The templates are listed as:
  - Bash via OpenPBS
  - Python 3.9 via OpenPBS

and accessed via navigation menu→dedicated BCM extension panel→kernel templates section.
After running the `cm-kubernetes-setup` cluster manager utility to deploy a Kubernetes cluster, the Kubernetes cluster instance is displayed (navigation menu → dedicated BCM extension panel → Kubernetes clusters section)

Then, after running the `cm-jupyter-kernel-operator` cluster manager utility to deploy a Jupyter kernel operator package, and configuring a user (section 6.2 of the Containerization Manual), the templates `jupyter-eg-kernel-k8s-cmjkop-julia`, `jupyter-eg-kernel-k8s-cmjkop-py`, and `jupyter-eg-kernel-k8s-cmjkop-py-spark` become available.

The templates are listed as:

- Julia on Kubernetes Operator
- Python on Kubernetes Operator
- Python+Spark on Kubernetes Operator

and accessed via navigation menu → dedicated BCM extension panel → kernel templates section.

Users can instantiate a kernel template to create an actual kernel from the dedicated BCM extensions section using the + button of the template. A dialog is dynamically generated for the template being instantiated, and users are asked to fill a number of customization options defined by administrators (figure 13.3):

![Figure 13.3: Jupyter kernel template customization screen](image)

Once the template is completely customized, the kernel can be created. It automatically appears in the JupyterLab Launcher screen (figure 13.4) and can be used to run notebooks or a console session (figure 13.4):
A user who lists available Jupyter kernels via the command line now sees the newly-created kernel:

```
[alice@basecm10 ~]$ module load jupyter
[alice@basecm10 ~]$ jupyter kernelspec list
Available kernels:
jupyter-eg-kernel-k8s-cmjkop-py-1gii7nm01 /home/alice/.local/share/jupyter/kernels/jupyter-eg-kernel-
k8s-cmjkop-py-1gii7nm01
python3 /cm/shared/apps/jupyter/14.0.0/share/jupyter/kernels/python3
```

The new kernel directory will contain the JSON definition generated by the Jupyter Kernel Creator:

```
[alice@basecm10 ~]$ cat .local/share/jupyter/kernels/jupyter-eg-kernel-k8s-cmjkop-py-
lgi7nm01/kernel.json
{
  "language": "python",
  "display_name": "Python on Kubernetes Operator 221123133450",
  "metadata": {
    "process_proxy": {
      "class_name": "cm_jupyter_kernel_creator.eg_processproxies.k8scmjkop.KubernetesCMJupyterKernelOperator",
      "config": {
        "k8s_env_module": "kubernetes",
        "cmjk_template": "templates/cmjk.yaml.j2",
        "image": "jupyter/datascience-notebook",
        "image_pull_policy": "IfNotPresent",
        "namespace": "alice-restricted",
        "gpu_limit": 0
      }
    }
  }
}```
Jupyter kernel names need not be unique. Users should therefore choose meaningful and distinguishable display names for their kernels. Doing so makes the JupyterLab Launcher screen easier to use.

For convenience, a summary of the available kernel templates and their requirements is shown in Table 13.1:

<table>
<thead>
<tr>
<th>Template name</th>
<th>Requirement</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>jupyter-eg-kernel-k8s-bash</td>
<td>Kubernetes 1</td>
<td>Bash via Kubernetes</td>
</tr>
<tr>
<td>jupyter-eg-kernel-k8s-py</td>
<td>Kubernetes 1</td>
<td>Python 3.9 via Kubernetes</td>
</tr>
<tr>
<td>jupyter-eg-kernel-k8s-py-spark</td>
<td>Kubernetes 2</td>
<td>Python 3.9 + Spark 3 via Kubernetes</td>
</tr>
<tr>
<td>jupyter-eg-kernel-k8s-r</td>
<td>Kubernetes 3</td>
<td>R 3.4 via Kubernetes</td>
</tr>
<tr>
<td>jupyter-eg-kernel-k8s-cmjkop-ngc-py</td>
<td>Kubernetes</td>
<td>NGC images via Jupyter Kernel Operator</td>
</tr>
<tr>
<td>jupyter-eg-kernel-k8s-cmjkop-julia</td>
<td>Kubernetes</td>
<td>Jupyter official image via Jupyter Kernel Operator (using Julia)</td>
</tr>
<tr>
<td>jupyter-eg-kernel-k8s-cmjkop-py</td>
<td>Kubernetes</td>
<td>Jupyter official image via Jupyter Kernel Operator (Python)</td>
</tr>
<tr>
<td>jupyter-eg-kernel-k8s-cmjkop-py-spark</td>
<td>Kubernetes 3</td>
<td>Python + Spark via Jupyter Kernel Operator (using Python and Spark)</td>
</tr>
<tr>
<td>jupyter-eg-kernel-1sf-bash</td>
<td>Platform LSF</td>
<td>Bash via Platform LSF</td>
</tr>
<tr>
<td>jupyter-eg-kernel-1sf-py39</td>
<td>Platform LSF</td>
<td>Python 3.9 via Platform LSF</td>
</tr>
<tr>
<td>jupyter-eg-kernel-openpbs-bash</td>
<td>Open PBS</td>
<td>Bash via Open PBS</td>
</tr>
<tr>
<td>jupyter-eg-kernel-openpbs-py39</td>
<td>Open PBS</td>
<td>Python 3.9 via Open PBS</td>
</tr>
<tr>
<td>jupyter-eg-kernel-pbspro-bash</td>
<td>PBS Professional</td>
<td>Bash via PBS Professional</td>
</tr>
<tr>
<td>jupyter-eg-kernel-pbspro-py39</td>
<td>PBS Professional</td>
<td>Python 3.9 via PBS Professional</td>
</tr>
</tbody>
</table>

...continues
Table 13.1: Available Jupyter kernel templates...continued

<table>
<thead>
<tr>
<th>Template name</th>
<th>Requirement</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>jupyter-eg-kernel-slurm-bash</td>
<td>Slurm</td>
<td>Bash via Slurm</td>
</tr>
<tr>
<td>jupyter-eg-kernel-slurm-py39</td>
<td>Slurm</td>
<td>Python 3.9 via Slurm</td>
</tr>
<tr>
<td>jupyter-eg-kernel-uge-bash</td>
<td>GE</td>
<td>Bash via GE</td>
</tr>
<tr>
<td>jupyter-eg-kernel-uge-py39</td>
<td>GE</td>
<td>Python 3.9 via GE</td>
</tr>
</tbody>
</table>

1 Docker image: brightcomputing/jupyter-kernel-sample:k8s-py36-1.1.1
2 Docker image: brightcomputing/jupyter-kernel-sample:k8s-spark-py37-1.2.1
3 Docker image: brightcomputing/jupyter-kernel-sample:k8s-r-1.0.0
DockerHub kernels page: https://hub.docker.com/r/brightcomputing/jupyter-kernel-sample/tags

13.5 Jupyter VNC Extension

13.5.1 What Is Jupyter VNC Extension About?
VNC (Virtual Network Computing) is a screen sharing service that can work in a browser.
If VNC is allowed by the cluster administrator, then the Jupyter environment configured by BCM can be used to start and control remote desktops via VNC with the Jupyter VNC extension.

13.5.2 Enabling User Lingering
User lingering is a systemd setting that sets a user manager for a user at boot and keeps it around after logout. This allows that user to run long-running sessions despite not being logged in. Enabling user lingering may be required for Jupyter VNC extension to run for a relatively complicated desktop such as KDE or GNOME.

For each user on each machine where these environments are installed, the following command must be run:

```
loginctl enable-linger <username>
```

The command may also be carried out using prolog/epilog scripts in the chosen WLM.

13.5.3 Starting A VNC Session With The Jupyter VNC Extension
Users can start a VNC session with the button added by Jupyter VNC (figure 13.5). Additional VNC parameters can be optionally specified.

![Figure 13.5: Starting Jupyter VNC session from kernel](image)

If VNC is available and correctly configured on the node where the kernel is running, then a new tab
is automatically created by Jupyter VNC containing the new session (figure 13.6). A user can now freely interact in JupyterLab both with the notebook and with the desktop environment.

To provide a user-friendly experience, Jupyter VNC also allows the graphical viewport to be resized, so that the desktop application can run full-screen (figure 13.7).
13.5.4 Running Examples And Applications In The VNC Session With The Jupyter VNC Extension

Once the VNC session is correctly started and the new JupyterLab tab has been created, Jupyter VNC automatically exports the DISPLAY environment variable to the running notebook (figure 13.8). Doing so means that any application or library running in the notebook can make use of the freshly created desktop environment. An example of such a library is OpenAI Gym, a toolkit for developing and comparing reinforcement learning algorithms, that is distributed by BCM.

Among the examples distributed by BCM (section 13.2), a notebook running PyTorch in the OpenAI Gym CartPole environment can be found. If executed after a VNC session has been started, a user can then observe the model being trained in real time in the graphical environment.
13.6 Jupyter WLM Magic Extension

In the Jupyter environment configured by BCM, the Jupyter WLM Magic extension can be used to schedule workload manager jobs from notebooks.

The Jupyter WLM Magic extension is an IPython extension. It is designed to improve the capabilities of Jupyter's default Python 3 kernel, which runs on the login node.

The Jupyter WLM Magic extension should therefore not be used from kernels running on compute nodes, such as those typically created with BCM's Jupyter Kernel Creator extension (section 13.4), and submitted via Jupyter Enterprise Gateway. Indeed, compute nodes running these kernels are often incapable of starting workload manager jobs in many default WLM configurations.

Jupyter WLM Magic extension makes it possible for users to programmatically submit WLM jobs, and then interact with their results. This can be done while using the Python programming language and its libraries, which are available in the notebook.

Users submit jobs and check their progress from the login node. The actual computation is distributed by the underlying workload manager across compute nodes, which means that server resources are spared.

Jupyter WLM Magic commands are available in the IPython kernel as magic functions (https://ipython.readthedocs.io/en/stable/interactive/tutorial.html#magic-functions). A new line magic (%) and a new cell magic (%%) are now added in the kernel, according to the workload manager:

- Platform LSF: %lsf_job and %%lsf_job
- PBS Professional: %pbspro_job and %%pbspro_job
- Slurm: %slurm_job and %%slurm_job
- GE: %uge_job and %%uge_job

A user can list the magic functions in the kernel to see if they are available, with Jupyter's built-in command %lsmagic (https://ipython.readthedocs.io/en/stable/interactive/magics.html#magic-lsmagic):

**Example**

```plaintext
In [1]: %lsmagic
Out [1]: root:
    line:
    ...
    slurm_job:"SLURMMagic"
pbspro_job:"PBSProMagic"
lsf_job:"LSFMagic"
uge_job:"UGEMagic"
    call:
    ...
    slurm_job:"SLURMMagic"
pbspro_job:"PBSProMagic"
lsf_job:"LSFMagic"
uge_job:"UGEMagic"
```

The magic functions introduced by this BCM extension share a similar syntax. For convenience, Slurm is used as an example in this section. However, the same instructions are valid for the other WLMs.

Users can check which options are available for a WLM function with the line magic helper:
Example

In []: %slurm_job --help
Out []: usage: %slurm_job [-h] [--module MODULE] [--module-load-cmd MODULE_LOAD_CMD]
         [--shell SHELL] [--submit-command SUBMIT_COMMAND]
         [--cancel-command CANCEL_COMMAND]
         [--control-command CONTROL_COMMAND]
         [--stdout-file STDOUT_FILE] [--stderr-file STDERR_FILE]
         [--preamble PREAMBLE] [--timeout TIMEOUT]
         [--check-condition-var CHECK_CONDITION_VAR]
         [--job-id-var JOB_ID_VAR]
         [--stdout-file-var STDOUT_FILE_VAR]
         [--stderr-file-var STDERR_FILE_VAR] [--dont-wait]
         [--write-updates WRITE_UPDATES]
         [--check-status-every CHECK_STATUS_EVERY]

optional arguments:
-h, --help            show this help message and exit
[...]

Line magic functions are typically used to set options with a global scope in the notebook. By doing so, a user will not need to specify the same option every time a job will be submitted via cell magic. For example, if two Slurm instances are deployed on the cluster and their associated environment modules are `slurm-primary` and `slurm-secondary`, a user could run the following line magic once to configure the Jupyter WLM Magic extension to always use the second deployment:

Example

In []: %slurm_job --module slurm-secondary
Out []: 

Now, jobs will always be submitted to `slurm-secondary`. This is more convenient than repeatedly defining the same module option for every cell magic upon scheduling a job:

Example

In []: %%slurm_job --module slurm-secondary
Out []: <WLM JOB DEFINITION>

In []: %%slurm_job --module slurm-secondary
Out []: <WLM JOB OUTPUT>

It should be noted that line magic functions cannot be used to submit WLM jobs. Cell magic functions have to be used instead.

A well-defined cell contains the WLM cell magic function provided by the extension, followed by the traditional job definition. For example, a simple MPI job running on two nodes can be submitted to Slurm by defining and running this cell:

Example

In []: %%slurm_job
   #SBATCH -J mpi-job-example
   #SBATCH -N 2
   module load openmpi
   mpirun hostname
Out []: COMPLETED
Users can take advantage of the Jupyter WLM Magic extension to store some information into Python variables about the job being submitted. The information could be the ID or the output file name, for example. Users can then later programmatically interact with them in Python. This feature is convenient when a user wants to, for example, programmatically carry out new actions depending on the job output:

**Example**

In []: %%slurm_job --job-id-var my_job_id --stdout-file-var my_job_out

#SBATCH -J mpi-job-example
#SBATCH -N 2
module load openmpi
mpirun hostname

Out []: COMPLETED

STDOUT file content: /home/demo/.jupyter/wlm_magic/slurm-2.out
node001
node001
node002
node002

In []: print(f"Job id {my_job_id} was written to {my_job_out}")
print(f"Output lines: {open(my_job_out).readlines()}")

Out []: Job id 2 was written to /home/demo/.jupyter/wlm_magic/slurm-2.out
Output lines: ['node001
node002

Users can also exploit Python variables to define the behavior of the Jupyter WLM Magic extension. For example, they can define a Python boolean variable to submit a WLM job only if a condition is true:

**Example**

In []: run_job = 1 == 2

Out []:

In []: %%slurm_job --check-condition-var run_job

#SBATCH -J mpi-job-example
#SBATCH -N 2
module load openmpi
mpirun hostname

Out []: Variable run_job is 'False'. Skipping submit.
A

MPI Examples

A.1 “Hello world”

A quick application to test the MPI libraries and the network.

/*
 * "Hello World' Type MPI Test Program
 */
#include <mpi.h>
#include <stdio.h>
#include <string.h>
#define BUFSIZE 128
#define TAG 0

int main(int argc, char *argv[])
{
    char idstr[32];
    char buff[BUFSIZE];
    int numprocs;
    int myid;
    int i;
    MPI_Status stat;

    /* all MPI programs start with MPI_Init; all 'N' processes exist thereafter */
    MPI_Init(&argc,&argv);
    MPI_Comm_size(MPI_COMM_WORLD,&numprocs); /* find out how big the SPMD world is */
    MPI_Comm_rank(MPI_COMM_WORLD,&myid); /* and this processes' rank is */

    /* At this point, all the programs are running equivalently, the rank is used to
       distinguish the roles of the programs in the SPMD model, with rank 0 often used
       specially... */
    if(myid == 0)
    {
        printf("%d: We have %d processors\n", myid, numprocs);
        for(i=1;i<numprocs;i++)
        {
            sprintf(buff, "Hello %d! ", i);
            MPI_Send(buff, BUFSIZE, MPI_CHAR, i, TAG, MPI_COMM_WORLD);
        }
    for(i=1;i<numprocs;i++)
    {
A.2 MPI Skeleton

The sample code below contains the complete communications skeleton for a dynamically load balanced head/compute node application. Following the code is a description of some of the functions necessary for writing typical parallel applications.

```c
#include <mpi.h>
#define WORKTAG 1
#define DIETAG 2

main(argc, argv) int argc; char *argv[];
{
    int myrank;
    MPI_Init(&argc, &argv); /* initialize MPI */
    MPI_Comm_rank(
        MPI_COMM_WORLD, /* always use this */
        &myrank); /* process rank, 0 thru N-1 */
    if (myrank == 0) {
        head();
    } else {
        computenode();
    }
    MPI_Finalize(); /* cleanup MPI */
}

head()
{
    int ntasks, rank, work;
    double result;
    MPI_Status status;
    MPI_Comm_size(
        MPI_COMM_WORLD, /* always use this */
        &ntasks); /* #processes in application */
    /* receive from rank 0: */
    MPI_Recv(buff, BUFSIZE, MPI_CHAR, 0, TAG, MPI_COMM_WORLD, &stat);
    sprintf(idstr, "Processor %d ", myid);
    strcat(buff, idstr);
    strcat(buff, "reporting for duty\n");
    /* send to rank 0: */
    MPI_Send(buff, BUFSIZE, MPI_CHAR, 0, TAG, MPI_COMM_WORLD);
    printf("%d: %s\n", myid, buff);
}
else
{
    MPI_Recv(buff, BUFSIZE, MPI_CHAR, i, TAG, MPI_COMM_WORLD, &stat);
    printf("%d: %s\n", myid, buff);
}

/* MPI Programs end with MPI Finalize; this is a weak synchronization point */
MPI_Finalize();
return 0;
}
```
A.2 MPI Skeleton

```c
/*
 * Seed the compute nodes.
 */
for (rank = 1; rank < ntasks; ++rank) {
    work = /* get_next_work_request */;
    MPI_Send(&work, /* message buffer */
             1, /* one data item */
             MPI_INT, /* data item is an integer */
             rank, /* destination process rank */
             WORKTAG, /* user chosen message tag */
             MPI_COMM_WORLD); /* always use this */
}

/*
 * Receive a result from any compute node and dispatch a new work
 * request work requests have been exhausted.
 */
work = /* get_next_work_request */;
while (/* valid new work request */) {
    MPI_Recv(&result, /* message buffer */
              1, /* one data item */
              MPI_DOUBLE, /* of type double real */
              MPI_ANY_SOURCE, /* receive from any sender */
              MPI_ANY_TAG, /* any type of message */
              MPI_COMM_WORLD, /* always use this */
              &status); /* received message info */
    MPI_Send(&work, 1, MPI_INT, status.MPI_SOURCE,
             WORKTAG, MPI_COMM_WORLD);
    work = /* get_next_work_request */;
}

/*
 * Receive results for outstanding work requests.
 */
for (rank = 1; rank < ntasks; ++rank) {
    MPI_Recv(&result, 1, MPI_DOUBLE, MPI_ANY_SOURCE,
              MPI_ANY_TAG, MPI_COMM_WORLD, &status);
}

/*
 * Tell all the compute nodes to exit.
 */
for (rank = 1; rank < ntasks; ++rank) {
    MPI_Send(0, 0, MPI_INT, rank, DIETAG, MPI_COMM_WORLD);
}
}
```

computenode()
{
    double result;
    int work;
    MPI_Status status;
    for (;;) {
        MPI_Recv(&work, 1, MPI_INT, 0, MPI_ANY_TAG,
                  MPI_COMM_WORLD, &status);
        /*
```
* Check the tag of the received message. */
  if (status.MPI_TAG == DIETAG) {
      return;
  }
  result = /* do the work */;
  MPI_Send(&result, 1, MPI_DOUBLE, 0, 0, MPI_COMM_WORLD);
}

Processes are represented by a unique rank (integer) and ranks are numbered 0, 1, 2, ..., N-1. MPI_COMM_WORLD means all the processes in the MPI application. It is called a communicator and it provides all information necessary to do message passing. Portable libraries do more with communicators to provide synchronisation protection that most other systems cannot handle.

A.3 MPI Initialization And Finalization

As with other systems, two functions are provided to initialize and clean up an MPI process:

MPI_Init(&argc, &argv);
MPI_Finalize();


Typically, a process in a parallel application needs to know who it is (its rank) and how many other processes exist.

A process finds out its own rank by calling:

MPI_Comm_rank( );
Int myrank;
MPI_Comm_rank(MPI_COMM_WORLD, &myrank);

The total number of processes is returned by MPI_Comm_size( ):

int nprocs;
MPI_Comm_size(MPI_COMM_WORLD, &nprocs);

A.5 Sending Messages

A message is an array of elements of a given data type. MPI supports all the basic data types and allows a more elaborate application to construct new data types at runtime. A message is sent to a specific process and is marked by a tag (integer value) specified by the user. Tags are used to distinguish between different message types a process might send/receive. In the sample code above, the tag is used to distinguish between work and termination messages.

MPI_Send(buffer, count, datatype, destination, tag, MPI_COMM_WORLD);

A.6 Receiving Messages

A receiving process specifies the tag and the rank of the sending process. MPI_ANY_TAG and MPI_ANY_SOURCE may be used optionally to receive a message of any tag and from any sending process.

MPI_Recv(buffer, maxcount, datatype, source, tag, MPI_COMM_WORLD, &status);
Information about the received message is returned in a status variable. The received message tag is `status.MPI_TAG` and the rank of the sending process is `status.MPI_SOURCE`. Another function, not used in the sample code, returns the number of data type elements received. It is used when the number of elements received might be smaller than `maxcount`.

```c
MPI_Get_count(&status, datatype, &nelements);
```

### A.7 Blocking, Non-Blocking, And Persistent Messages

`MPI_Send` and `MPI_Receive` cause the running program to wait for non-local communication from a network. Most communication networks function at least an order of magnitude slower than local computations. When an MPI process has to wait for non-local communication CPU cycles are lost because the operating system has to block the process, then has to wait for communication, and then resume the process.

An optimal efficiency is usually best achieved by overlapping communication and computation. **Blocking** messaging functions only allow one communication to occur at a time. **Non-blocking** messaging functions allow the application to initiate multiple communication operations, enabling the MPI implementation to proceed simultaneously. **Persistent** non-blocking messaging functions allow a communication state to persist, so that the MPI implementation does not waste time on initializing or terminating a communication.

#### A.7.1 Blocking Messages

In the following example, the communication implementation executes in a sequential fashion causing each process, `MPI_Recv`, then `MPI_Send`, to block while waiting for its neighbor:

**Example**

```c
while (looping) {
    if (i_have_a_left_neighbor)
        MPI_Recv(inbuf, count, dtype, left, tag, comm, &status);
    if (i_have_a_right_neighbor)
        MPI_Send(outbuf, count, dtype, right, tag, comm);
    do_other_work();
}
```

MPI also has the potential to allow both communications to occur simultaneously, as in the following communication implementation example:

#### A.7.2 Non-Blocking Messages

**Example**

```c
while (looping) {
    count = 0;
    if (i_have_a_left_neighbor)
        MPI_Irecv(inbuf, count, dtype, left, tag, comm, &req[count++]);
    if (i_have_a_right_neighbor)
        MPI_Isend(outbuf, count, dtype, right, tag, comm, &req[count++]);
    MPI_Waitall(count, req, &statuses);
    do_other_work();
}
```

In the example, `MPI_Waitall` potentially allows both communications to occur simultaneously. However, the process as show is blocked until both communications are complete.
A.7.3 Persistent, Non-Blocking Messages

A more efficient use of the waiting time means to carry out some other work in the meantime that does not depend on that communication. If the same buffers and communication parameters are to be used in each iteration, then a further optimization is to use the MPI persistent mode. The following code instructs MPI to set up the communications once, and communicate similar messages every time:

Example

```c
int count = 0;
if (i_have_a_left_neighbor)
    MPI_Recv_init(inbuf, count, dtype, left, tag, comm, &req[count++]);
if (i_have_a_right_neighbor)
    MPI_Send_init(outbuf, count, dtype, right, tag, comm, &req[count++]);
while (looping) {
    MPI_Startall(count, req);
    do_some_work();
    MPI_Waitall(count, req, &statuses);
    do_rest_of_work();
}
```

In the example, `MPI_Send_init` and `MPI_Recv_init` perform a persistent communication initialization.
## Compiler Flag Equivalence

The following table is an overview of some of the compiler flags that are equivalent or almost equivalent.

<table>
<thead>
<tr>
<th>Cray</th>
<th>Intel</th>
<th>GCC</th>
<th>Explanation</th>
</tr>
</thead>
<tbody>
<tr>
<td>default</td>
<td>default</td>
<td>-O3 -ffast-math</td>
<td>Produce high level of optimization</td>
</tr>
<tr>
<td>-Oomp (default)</td>
<td>-openmp</td>
<td>-f openmp</td>
<td>Activate OpenMP directives and pragmas in the code</td>
</tr>
<tr>
<td>-h byteswapio</td>
<td>-convert</td>
<td>-f convert=swap</td>
<td>Read and write Fortran unformatted data files as big-endian</td>
</tr>
<tr>
<td></td>
<td>big_endian</td>
<td></td>
<td></td>
</tr>
<tr>
<td>-f fixed</td>
<td>-fixed</td>
<td>-ffixed-form</td>
<td>Process Fortran source using fixed form specifications.</td>
</tr>
<tr>
<td>-f free</td>
<td>-free</td>
<td>-ffree-form</td>
<td>Process Fortran source using free form specifications.</td>
</tr>
<tr>
<td>-V</td>
<td>--version</td>
<td>--version</td>
<td>Dump version.</td>
</tr>
<tr>
<td>-h zero</td>
<td>N/A</td>
<td>-finit-local-zero</td>
<td>Zero fill all uninitialized variables.</td>
</tr>
<tr>
<td>-e m</td>
<td></td>
<td></td>
<td>Creates .mod files to hold Fortran90 module information for future compiles.</td>
</tr>
<tr>
<td>-j &lt;dir_name&gt;</td>
<td></td>
<td></td>
<td>Specifies the directory &lt;dir_name&gt; to which .mod files are written when the -e m option is specified</td>
</tr>
</tbody>
</table>