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Intended Audience

This document is meant for PARAM Pravega users.

Typographic Conventions

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Meaning</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blue underlined text</td>
<td>A hyperlink or link you can click to go to a related section in this document or to a URL in your web browser.</td>
</tr>
<tr>
<td>Bold</td>
<td>The names of menus, menu items, headings, and buttons.</td>
</tr>
<tr>
<td>Italics</td>
<td>Variables or placeholders or special terms in the document.</td>
</tr>
<tr>
<td>Console text</td>
<td>Console commands</td>
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</table>

Getting help

For technical assistance, use ticketing tool.
Give us your feedback

We value your feedback. Kindly send your comments on content of this document to samirs@cdac.in. Please include the page number of the document along with your feedback.

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Spack

Introduction

Spack automates the download-build-install process for software - including dependencies - and provides convenient management of versions and build configurations. It is designed to support multiple versions and configurations of software on a wide variety of platforms and environments. It is designed for large supercomputing centers, where many users and application teams share common installations of software on clusters with exotic architectures, using libraries that do not have a standard ABI. Spack is non-destructive: installing a new version does not break existing installations, so many configurations can coexist on the same system.

Getting Started

On your login node command prompt execute below commands:

```
$ module load spack
```

To load SPACK module and setting up environment for SPACK.

Kindly see the above screenshot and source below line including initial dot.

```
$ . /home/apps/spack/share/spack/setup-env.sh
```
To Use Pre-installed Applications from Spack

**Spack find**

The `spack find` command is used to query installed packages on Param Pravega. Note that some packages appear identical with the default output. The `-l` flag shows the hash of each package, and the `-f` flag shows any non-empty compiler flags of those packages.

**Spack load application name**

The easiest way is to use `spack load <application name@version>`
To Know the Pre-Loaded Application/Compliers

$ spack find --loaded
=> 6 installed packages
-- linux-ubuntu18.04-x86_64 / gcc@7.5.0 ---------------
gcc@8.3.0  gmp@6.1.2  isl@0.18  mpc@1.1.0  mpfr@3.1.6  zlib@1.2.11
To install new application

First check the available compilers in Spack with below command:

**spack compilers**

Spack manages a list of available compilers on the system, detected automatically from the user’s PATH variable. The `spack compilers` command is an alias for the command `spack compiler list`.

![Compilation Result](image)

**To Check the Compilers Available in the System**

```
$ spack compiler list
==> Available compilers
-- clang ubuntu18.04-x86_64 -------------------------------------
  clang@6.0.0
-- gcc ubuntu18.04-x86_64 ---------------------------------------
  gcc@8.3.0  gcc@7.5.0  gcc@6.5.0
```

Check if application is available in Spack repo with command-

**spack list**

The `spack list` command shows available packages.

The `spack list` command can also take a query string. Spack automatically adds wildcards to both ends of the string, or you can add your own wildcards.
Before installing application check its spec with command

```bash
spack spec zlib
```

Show what would be installed, given a spec. The spec syntax also includes compiler flags.

Spack accepts `cppflags`, `cflags`, `cxxflags`, `fflags`, `ldflags`, and `ldlibs` parameters. The values of these fields must be quoted on the command line if they include spaces. These values are injected into the compile line automatically by the Spack compiler wrappers.

To change default compiler for zlib installation to oneapi
spack spec zlib %oneapi

Below is an example of installation of package using spack:

```bash
spack install gromacs@2020.5 +cuda-mpi+blas %intel ^intel-mkl
```

Above command will install gromacs version 2020.5 with blas and cuda support and without MPI support. For blas there are multiple providers like OpenBLAS, Intel MKL, amdblis, and essl, ^intel-mkl will tell spack to use intel-mkl for blas routines.

Operators in Spack

- `%` to select compiler out of available compilers
- `^` to use variant of package
- `@` to define the version number of packages to be installed.
- `+` to enable variant for package
- `~` to disable variant for package
Uninstalling Packages

Earlier we installed many configurations each of zlib. Now we will go through and uninstall some of those packages that we didn’t really need.

```
$ spack uninstall zlib %gcc@6.5.0
(type : y)
```
Using Environments

Spack has an environment feature in which you can group installed software. You can install software with different versions and dependencies in each environment and can change software to use at once by changing environments. You can create a Spack environment by `spack env create` command. You can create multiple environments by specifying different environment names here.

```
spack env create myenv
```

To activate the created environment, type `spack env activate`. Adding `-p` option will display the current activated environment on your console. Then, install software you need to the activated environment.

```
spack env activate -p myenv
[myenv] [username@es1 ~]$ spack install xxxxx
```

You can deactivate the environment by `spack env deactivate`. To switch to another environment, type `spack env activate` to activate it.

```
[myenv] [username@es1 ~]$ spack env deactivate
[username@es1 ~]$ 
```

Use `spack env list` to display the list of created Spack environments.

```
[username@es1 ~]$ spack env list
==> 2 environments
    myenv
    another_env
```

```
spack env
```

Refer below screenshot to activate an environment, add a package and install it in that environment:
The `spack env activate` will load the view associated with the Environment into the user environment.
Packaging (For Application developers)

Spack packages are installation scripts, which are essentially recipes for building the software.

They define properties and behaviour of the build, such as:

- where to find and how to retrieve the software.
- its dependencies.
- options for building the software from source; and
- build commands.

Once we’ve specified a package’s recipe, users of our recipe can ask Spack to build the software with different features on any of the supported systems. Please refer Packaging Guide — Spack 0.17.0 documentation for detailed understanding of the Spack packaging.

Example Creating Own Package:

In below spec file we have used Linewidth an IISc developed code. Please see the bold lines for comments related to preceding lines in the spec file of spack package named lisclinewidth:

```python
# Copyright 2013-2021 Lawrence Livermore National Security, LLC and other # Spack Project Developers. See the top-level COPYRIGHT file for details. # # SPDX-License-Identifier: (Apache-2.0 OR MIT)
import os
import platform
import sys
import llnl.util.tty as tty
from spack import *
class IiscLinewidth(MakefilePackage):
    ""
    Linewidth developed by IISC Banglore.
    ""
    homepage = ""
    
    #Url for homepage
    url = "file:///{0}/linewidth.tar.gz".format(os.getcwd())
    
    #Url for source code
    manual_download = True
    
    #If source code is not available in public domain
    version('1',
        sha256='7215f6765e5f5eddfde5f0c67a5bbdef5960607f3e199a609ef5619278ec8a66',
        preferred=True)
    
    #You can add different versions for you package.
    variant('mpi', default=True, description='Install with MPI support')
    variant('openmp', default=True, description='Install with OpenMP support')
    
    #Variant gives flexibility to users for changing parameter before
    compilation.
    depends_on('gmake', type='build')
    depends_on('mpi', when='+mpi')
    depends_on('hdf5+fortran+hl+mpi')
    depends_on('intel-mkl')
```
depends_on('py-h5py')
depends_on('py-matplotlib', type=('build', 'run'))

#Depend clause used to specify dependencies for your code.

@property
def build_targets(self):
targets = [
    '#--directory=SRC',
    '--file=Makefile',
    'LIBS={0} {1} '.format(self.spec['intel-mkl'].libs.ld_flags,
                          self.spec['hdf5'].libs.ld_flags),
    'HDFINCFLAGS={0}'.format(self.spec['hdf5'].prefix.include),
    'HDF5_HOME={0}'.format(self.spec['hdf5'].prefix),
    'FC={0}'.format(self.spec['mpi'].mpifc)
]
    return targets
def install(self, spec, prefix):
    mkdirp(prefix.bin)
    install('linewidth', prefix.bin)

###
#This code uses Makefile for building application. We can define some properties
# to make changes in Makefile, changing parameter in Makefile at compile time.

Sample Steps taken for Creating Linewidth application recipe for Spack

1. Source code
   Source code of Linewidth was not available through public repo like github, so needed to import OS package.
   - os.getcwd() - expects the source tar present in current working directory.
   - cha256- to check for sha256 checksum we added same in version clause and for place holder we have give version as 1.
   - manual download = True refers to spack will not try to download source code for the package.
   - name- make sure that name of tar file is same as used inside package recipe

2. Variant- User can control behavior of application being built through this clause.
   - Ex- To enable MPI support we have define it to be true by default.

3. depends_on() - This clause defines all dependencies required to build the given application.
   - Ex- In linewidth example we have used Intel-mkl and HDF5.

4. @property - With this decorator we can define some properties for build system like edit, build, install.

5. property build_targets - Defines logic of building source for native platform.

6. property install - Defines install procedure to be used after building source code.
   - Ex- In our example we define prefix path
Sample SLURM script for OpenMP applications/programs. to use spack

```bash
#!/bin/bash
#SBATCH --nodes=1
#SBATCH -p cpu ## gpu/standard
#SBATCH --exclusive
#SBATCH -t 1:00:00

echo "SLURM_JOBID="$SLURM_JOBID
echo "SLURM_JOB_Nodelist"=$SLURM_JOB_NODELIST
echo "SLURM_Nnodes"=$SLURM_NNODES
echo "SLURM_Ntasks"=$SLURM_NTASKS
ulimit -s unlimited
ulimit -c unlimited
export OMP_NUM_THREADS=4  ### Maximum number of threads= Number of physical core

#To load necessary application/compiler through spack
module load spack
export SPACK_ROOT=/home/apps/spack
$SPACK_ROOT/share/spack/setup-env.sh
spack load intel-mpi@2019.10.317 /6icwzn3
spack load intel-mkl@2020.4.304
spack load intel-oneapi-compilers@2021.4.0
spack load gcc@11.2.0
(time <executable_path> )
```

Sample SLURM script for MPI applications/programs. to use spack

```bash
#!/bin/bash
#SBATCH --nodes=2
#SBATCH -p cpu ## gpu/standard
#SBATCH --exclusive
#SBATCH -t 1:00:00

echo "SLURM_JOBID="$SLURM_JOBID
echo "SLURM_JOB_Nodelist"=$SLURM_JOB_NODELIST
echo "SLURM_Nnodes"=$SLURM_NNODES
echo "SLURM_Ntasks"=$SLURM_NTASKS
ulimit -s unlimited
ulimit -c unlimited

#To load necessary application/compiler through spack
module load spack
export SPACK_ROOT=/home/apps/spack
$SPACK_ROOT/share/spack/setup-env.sh
spack load intel-mpi@2019.10.317 /6icwzn3
spack load intel-mkl@2020.4.304
spack load intel-oneapi-compilers@2021.4.0
spack load gcc@11.2.0
(time mpirun -np $SLURM_Ntasks <executable_path>
```

For more information, related to SLURM, please refer the USER MANUAL.
References

2. https://github.com/spack/spack
3. Getting started
4. Basic usage
5. Packaging guide
6. Build system