Using Python and Jupyter on Perlmutter
Python/Jupyter users, welcome to Perlmutter!

What we’ll cover in this short 30 mins

• General Python advice
• Using Python on GPUs
• Using Jupyter on Perlmutter
• Open Q&A (5 mins at end)
Python on Perlmutter
What should I know moving from Cori to Perlmutter?

• More unified Python module + Jupyter configuration 👏
  o Jupyter Python 3 kernel is based on Python default module, currently `python/3.9-anaconda-2021.11`
  o Python and Jupyter Python 3 Kernel share `$_PYTHONUSERBASE`, which determines location and search path of pip installed packages

• Many “best practices” remain
  o Use our `/global/common/software/<your project>` filesystem for better performance
  o Use custom conda environments for customizable Python sandboxes, can be easily converted into a Jupyter kernel
  o Use a Shifter container, can be easily converted into a Jupyter kernel

• 6000+ GPUs!
More systems, more problems! How to avoid common problems:

• Cori and Perlmutter are two very different systems that are still sharing filesystems, including $HOME and dotfiles (\.bashrc, \.bash_profile, etc.)

• Don’t put system-specific modifications in your dotfiles
  o Actually, put as little in your dotfiles as you can and make sure to periodically review their contents

• Most likely code/environments you have built for Cori will not work on Perlmutter, and vice-versa. mpi4py will not work across systems.

• Tip– append system name to your custom conda environments, like dask-cori and dask-pm to help you keep track

• Be more careful using pip (we’ll cover this)
Use environments with conda activate

• You can now use **conda activate** on Perlmutter without using **conda init** and it will not make changes to your .bashrc.

• This is already possible on Perlmutter and will change on Cori at the AY rollover on Jan 19.

Old 😞

module load python
source activate myenv
conda deactivate

or

module load python
conda init
conda activate myenv
conda deactivate

New 👍

module load python
conda activate myenv
conda deactivate

The current default python module is python/3.9-anaconda-2021.11

Check out our pending updates to the Python docs for more information.
Building and using mpi4py

• mpi4py is available via `module load python`
• This mpi4py is CUDA-aware (can communicate GPU objects)
• To build your own CUDA-aware mpi4py, follow this recipe:

```bash
module load PrgEnv-gnu cudatoolkit python
conda create -n cudaaware python=3.9 -y
conda activate cudaaware
MPICC="cc -target-accel=nvidia80 -shared" pip install --force --no-cache-dir --no-binary=mpi4py mpi4py
```

• Be aware that with any CUDA-aware mpi4py, you must have `cudatoolkit` loaded, even for code that does not use the GPU
Use pip with caution⚠️

- Be careful with pip!!!! pip will try to be clever and find existing packages to save time, but most likely you don’t want this
- Best practices for pip
  - Use it inside of a conda environment, not outside (don’t use --user)
  - Always pip install `<package> --force` (Did you notice this in our mpi4py recipe?)
  - This will force a rebuild, which is important if tries to reuse a package from Cori
- If you use `pip --user`, it will install packages to the location specified by `PYTHONUSERBASE`, which is by default $HOME/.local/perlmutter/3.9-anaconda-2021.11
- It is safe to delete this directory if you want to clean up/save space
Python on GPUs
Getting started with GPUs in Python

• NumPy and SciPy do not utilize GPUs out of the box

• There are many Python GPU frameworks out there:
  o “drop in” replacements for numpy, scipy, pandas, scikit-learn, etc
    o CuPy, RAPIDS, cuNumeric (coming soon?)
  o “machine learning” libraries that also support general GPU computing
    o PyTorch, TensorFlow
  o “I want to write my own GPU kernels”
    o Numba, PyOpenCL, PyCUDA

• Many of these GPU libraries have adopted the CUDA Array Interface which makes it easier to share array-like objects stored in GPU memory between the libraries

• There is also some effort in the community to standardize around a common Python array API
Getting started with GPUs in Python (CuPy)

Numpy features support by CuPy:

- **Basic indexing** (indexing by ints, slices, newaxes, and Ellipsis)
- Most of **Advanced indexing** (except for some indexing patterns with boolean masks)
- Data types (dtypes): bool_, int8, int16, int32, int64, uint8, uint16, uint32, uint64, float16, float32, float64, complex64, complex128
- Most of the **array creation routines** (empty, ones_like, diag, etc.)
- Most of the **array manipulation routines** (reshape, rollaxis, concatenate, etc.)
- All operators with **broadcasting**
- All **universal functions** for elementwise operations (except those for complex numbers)
- Linear algebra functions, including product (dot, matmul, etc.) and decomposition (cholesky, svd, etc.), accelerated by cuBLAS and cuSOLVER
- Multi-dimensional **fast Fourier transform** (FFT), accelerated by cuFFT
- Reduction along axes (sum, max, argmax, etc.)
Getting started with GPUs in Python (CuPy)

> ssh perlmutter
> module load cudatoolkit python
> conda create -y --name cupy-demo python=3.9 numpy scipy
> source activate cupy-demo
> pip install cupy-cuda114
> python
>>> import cupy as cp
>>> print(cp.array([1, 2, 3]))
[1 2 3]

Current default version is: cudatoolkit/21.9_11.4

See documentation at https://docs.nersc.gov/development/languages/python/using-python-perlmutter/ or open a ticket at https://help.nersc.gov/
Getting started with GPUs in Python (CuPy)

```python
>>> import numpy as np
>>> import cupy as cp

# Create an array on GPU/device
>>> x_gpu = cp.array([1, 2, 3])
>>> isinstance(x_gpu, cp.ndarray)
True

# Data Transfer
>>> x_cpu = np.array([1, 2, 3])  # create an array on CPU/host
>>> x_gpu = cp.asarray(x_cpu)  # move the data to the GPU/device

>>> x_gpu = cp.array([1, 2, 3])  # create an array in the GPU/device
>>> x_cpu = cp.asnumpy(x_gpu)  # move the array to the CPU/host
```

In general, it is a good idea to minimize data movement between Host and Device.
import cupy
import numba.cuda
import numpy

# CUDA kernel
@numba.cuda.jit
def _cuda_addone(x):
i = numba.cuda.grid(1)
if i < x.size:
x[i] += 1

# convenience wrapper with thread/block configuration
def addone(x):
    # threads per block
tpb = 32
    # blocks per grid
bpg = (x.size + (tpb - 1)) // tpb
_cuda_addone[bpg, tpb](x)

# create array on device using cupy
x = cupy.zeros(1000)

# pass cupy ndarray to numba.cuda kernel
addone(x)

# Use numpy api with cupy ndarray
total = numpy.sum(x)

https://docs.cupy.dev/en/stable/user_guide/basic.html

- NumPy's __array_function__ protocol (NEP 18)
  - numpy.sum(x) -> cupy.sum(x)
- CPU and GPU execution paths can share same implementation (sometimes)
- Can also use helper functions to get the appropriate array module. For example:
  - xp = cupy.get_array_module(x)
Profiling using NVIDIA Nsight Systems

```python
import cupy as cp

cp.cuda.nvtx.RangePush(message)
...
cp.cuda.nvtx.RangePop()

@cp.prof.TimeRangeDecorator(message)
def function():
    pass

with cp.prof.time_range(message):
    pass
```

CuPy supports for NVIDIA Tools Extension (NVTX) markers and ranges

Or use decorator syntax without modifying function body

Can also use `with`-statement context blocks

Run your application with Nsight Systems:

```bash
> nsys profile --trace cuda,nvtx --stats=true python myapp.py
```
Is my code a good fit for a GPU?

GPUs are likely a good fit if the following are true for your application:

- Performs computation using large arrays, matrices, or images
- Dataset can fit in GPU memory
  - (40GB for Perlmutter’s A100 GPUs)
- IO is not a bottleneck

For more help choosing a GPU-accelerated Python framework:

https://docs.nersc.gov/development/languages/python/perlmutter-prep/

or open a ticket at https://help.nersc.gov/

```
a = xp.random.rand(size, size)
b = xp.random.rand(size, size)
def f(a, b):
    return xp.dot(a, b)
```
How do I run Jupyter notebooks on Perlmutter Phase I?

Making sure you can access Perlmutter with Jupyter
Available configurations and what they are for

How do I make use of Perlmutter GPUs from Jupyter?

Hardware you can use for each configuration
Monitoring GPU usage in JupyterLab

What should I know moving from Cori to Perlmutter?

What differences in the deployment matter to you
How do I run Jupyter notebooks on Perlmutter Phase I?

Making sure you can access Perlmutter with Jupyter

https://jupyter.nersc.gov

https://jupyter.nersc.gov/hub/home (home or “console”)

If this row does not show up, you need Perlmutter to be added to your list of “server logins” in Iris.

Note: Your console may look a little different if you don’t have Cori GPU access for instance.
How do I run Jupyter notebooks on Perlmutter Phase I?

Available configurations and what they are for

- **Notebooks on a shared login node**
  - Will not be charged
  - No CPU/GPU/memory limits yet
  - Debug/test/develop
  - Not as compute intensive

- **Whole compute node to yourself**
  - Will be charged to sensible default
  - 6 hour time limit
  - Interactive GPU work
  - Compute intensive

- **Up to 4 nodes, up to 6 hours**
  - Customize your Slurm allocation
  - Interactive GPU work
  - You need to scale, baby!
  - ... need more? Contact me!
How do I make use of Perlmutter GPUs from Jupyter?

Hardware you can use for each configuration:

- **Login Node**
  - 1 "shared" NVIDIA A100
  - Kind of free-for-all for now, future: NVIDIA Multi Instance GPU (MIG)?

- **1 Compute Node**
  - 4 NVIDIA A100’s

- **4 Compute Nodes**
  - 16 NVIDIA A100’s
  - Limits may change per demand
How do I make use of Perlmutter GPUs from Jupyter?

Notebook: `!nvidia-smi`

Terminal: `nvidia-smi`

**Monitoring GPU usage in JupyterLab**

- **NVDashboard**
- **LabExtension**

Monitor GPUs on the node (Compute or login node)

For multi-node jobs, only GPUs on the same node as the notebook
How do I make use of Perlmutter GPUs from Jupyter?

Monitoring GPU usage in JupyterLab

Like Dask?
Want to try Dask?

Monitor everything
Watch progress
Profile your workflow
Separate tab for now
(Demo on 100 GPUs)

dask-labextension needed a “fix” for us to deploy
Just merged so it’s coming soon…!

Example notebook with dashboard setup instructions:
https://gitlab.com/NERSC/nersc-notebooks/-/tree/master/dask
What should I know moving from Cori to Perlmutter?

## What differences in the deployment matter to you

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<thead>
<tr>
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<th>Cori</th>
<th>Perlmutter</th>
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<tbody>
<tr>
<td>Shared Node Notebooks</td>
<td>24 login nodes total&lt;br&gt;4 dedicated to Jupyter&lt;br&gt;No other usages allowed&lt;br&gt;CPU/memory limits in place</td>
<td>40 login nodes total&lt;br&gt;Jupyter <em>not restricted to a subset</em>&lt;br&gt;Runs alongside ssh-based logins&lt;br&gt;No resource limits in place yet</td>
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<tr>
<td>Exclusive Node Notebooks</td>
<td>Limited resources available&lt;br&gt;Requires access to special QOS&lt;br&gt;Kind of fussy setup</td>
<td>Jupyter jobs are first class&lt;br&gt;No plans for special QOS&lt;br&gt;Watching allocation success rate&lt;br&gt;Will make adjustments to queues</td>
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<tr>
<td>Configurable Notebooks</td>
<td>Limited to GPU partition</td>
<td>Same deal as on Cori&lt;br&gt;Your conda envs may be bigger!</td>
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<tr>
<td>Setting up Kernels</td>
<td>Conda envs, custom, helpers:&lt;br&gt;See NERSC <a href="https://jupyter.org">Jupyter docs</a></td>
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Wrap Up
Where to get Python/Jupyter information, help

- Have a question? Try our documentation (updated almost daily!)
  - Using Perlmutter
  - Python at NERSC
  - Python on Perlmutter
  - Jupyter at NERSC
  - Try the search bar at docs.nersc.gov, it’s pretty good!
- Can’t find the answer? Submit a ticket at help.nersc.gov
Summary

- Welcome to Perlmutter!
- We are here to help you use Python and Jupyter productively on Perlmutter
- If you have questions, please check our docs.nersc.gov or file a ticket at help.nersc.gov
- Don’t be shy—now is the time to ask us questions!