





#### Python Basics on the FASRC Clusters





### Learning objectives

- Python using CLI (Command Line Interface)
  - Interactive
  - Sbatch
- Python Package installation
  - Interactive
  - Sbatch
- Python using OOD (Open On Demand)
- Jupyter Notebook
  - Create conda environment (i.e., jupyter kernel)





#### Python Programming Language Python Programming Language – FASRC DOCS

- High-level, general-purpose, and object-oriented programming language with emphasis on code readability and use of significant indentation.
- Ideal for scripting and rapid application development given its dynamic typing, elegant syntax, and automatic memory management (garbage collection).
- Has a comprehensive standard library. Also known as "batteries-included" language.
- Python's implementation is mostly in C.
  - Python's core interpreter, CPython, written in C.
- Interpreted language, hence slower than compiled languages, like C and Fortran.
  - Compiled generates executable
  - Interpreted executes instructions directly on the fly without compiling a program into machine language





#### Python using CLI - Interactive

```
# Login to Cannon
ssh <username>@login.rc.fas.harvard.edu
```

```
# Go to a compute node on the test partition:
salloc --partition test --nodes=1 --cpus-per-task=2 --mem=12GB --time=00:30:00
```

```
# Check Python modules available on Cannon:
module spider python
```

```
# Get detailed information on specific module, e.g.:
module spider python/3.10.13-fasrc01
```

```
# Load the latest (usually also the default) Python module:
module load python
```

#### **CLI Training:**

https://docs.rc.fas.harvard.edu/wp-content/uploads/2013/10/Getting-started-on-FASRC-clusters-with-CLI.pptx.pdf





### **Training Material**

https://docs.rc.fas.harvard.edu/kb/training-materials/

# Check current location & change if desired for this training: pwd
cd <desired-location>

# Clone FASRC User Codes repository: https://github.com/fasrc/User Codes/tree/master

**SSH** - git clone <u>git@github.com</u>:fasrc/User Codes.git

HTTPS - git clone <u>https://github.com/fasrc/User\_Codes.git</u>

# Create a training folder & go to that folder:

mkdir python-training

cd python-training

# Copy Python folders from the User Codes directory:

cp -r ../User\_Codes/Languages/Python .

cp -r ../User\_Codes/Parallel\_Computing/Python/Python-Multiprocessing-Tutorial .





#### Python using CLI - Interactive

- Check Python version: python --version
- Invoke Python interpreter: python
- Execute Python programming interactively:

```
def square(x):
    """square a number"""
    return x ** 2
for N in range(1, 4):
    print(N, "squared is", square(N))
```

- o Exit Python: exit()
- Or run a python script interactively: python myscript.py





# Python using CLI - sbatch; Example 1

#### https://github.com/fasrc/User\_Codes/tree/master/Languages/Python/Example1

# Go to Example1 folder

cd Python/Example1

# Submit job

sbatch run.sbatch

**run.sbatch**: Batch-job submission script for queuing the job

**mc\_pi.py**: Source code for calculating Pi using Monte-Carlo method

#!/bin/bash				
#SBATCH	-J mc_pi			
#SBATCH	-o mc_pi.out			
#SBATCH	-e mc_pi.err			
#SBATCH	nodes=1			
#SBATCH	cpus-per-task=1			
#SBATCH	<pre>partition=serial_requeue</pre>			
#SBATCH	time=0-00:30			
#SBATCH	mem=4000			

# job name
# standard output file
# standard error file
# number of nodes
# number of cores
# partition
# time in D-HH:MM
# memory in MB

# Load required modules
module load python

# Run program
python mc\_pi.py





#### Python Package Installation - Interactive

• Go to a compute node on the test partition:

salloc -p test --nodes=1 --cpus-per-task=2 --mem=12GB --time=01:00:00

• Create a vanilla mamba/conda environment (for multiprocessing exercise):

```
module load python
mamba create --prefix=/n/holylabs/LABS/<desired-folder>/multiproc_env
python=3.11 -y
```

• Alternatively, if default *\$HOME* is desired, then do following instead:

```
module load python
conda create --name multiproc_env python=3.11 -y
```

See <u>Python Package Installation</u>





# Python Package Installation

• Activate conda/mamba environment:

mamba activate /n/holylabs/LABS/<desired-folder>/multiproc\_env

- Or if \$HOME used, then: mamba activate multiproc\_env
- Install relevant python packages (Mamba recommended):

mamba install numpy pandas matplotlib -y
pip install jupyterlab swifter

- Always pip install inside a conda environment to avoid package conflicts
- o <a href="https://docs.rc.fas.harvard.edu/kb/python-package-installation/#Pip\_Installs">https://docs.rc.fas.harvard.edu/kb/python-package-installation/#Pip\_Installs</a>
- Deactivate the environment: mamba deactivate





### Python Package Installation - sbatch

https://github.com/fasrc/User\_Codes/tree/master/Languages/Python/Example2

# Go to Example2 folder

cd ../Python/Example2

# Submit job

sbatch run.sbatch

**numpy\_pandas\_ex.py**: source code for generating a dataframe utilizing a mamba environment

#!/bin/bash					
#SBATCH -J	np_pandas	#	job name		
#SBATCH -o	np_pandas.out	#	standard output file		
#SBATCH -e	np_pandas.err	#	standard error file		
#SBATCH	cpus-per-task=1	#	number of cores		
#SBATCH	partition=test	#	partition		
#SBATCH	time=0-01:00	#	time in D-HH:MM		
#SBATCHr	nem=10G	#	memory in GB		

# Load required modules
module load python

# Build the environment
sh build\_env.sh

# Activate the environment
mamba activate my\_env

# Run program
python numpy\_pandas\_ex.py





# Python Using Open OnDemand (OOD)

- Open-source web portal to access clusters
- Web-based, no software needs be installed on your local laptop/desktop (except for a modern browser like Google Chrome, Mozilla Firefox)
- Easy to learn and simple to use
- Very similar to desktop applications
- $\circ$  ~ The easiest way to run GUI applications remotely on a cluster
- $_{\odot}$   $\,$  Safari is not recommended for OOD  $\,$
- OOD Training: <u>https://docs.rc.fas.harvard.edu/wp-content/uploads/2013/10/Getting-started-on-FASRC-clusters-with-OOD-May20</u> <u>24.pdf</u>





#### How to access OOD on FASRC Clusters

- Accessing OOD from Cannon
  - Connect to FASRC VPN <u>Virtual Desktop (VDI) through Open OnDemand FASRC</u> <u>DOCS</u>
  - Then go to <u>https://rcood.rc.fas.harvard.edu</u>
- Accessing OOD from FASSE
  - Connect to FASSE VPN FASSE VDI Apps FASRC DOCS
  - Then go to <u>https://fasseood.rc.fas.harvard.edu</u>





# FASSE proxy

Documentation: FASSE Proxy Settings – FASRC DOCS

- $\circ$   $\,$  You may need to set FASSE proxy on
  - Firefox (web browsing)
  - Jupyter Notebook
  - Access Github
  - (Basically, anything outside of FASSE)





# Filling a form to launch an app

- Request the resources that you need
   (If you don't know for a first trial run, use similar resources as your laptop/desktop)
  - Partition (Name): depends on <u>Cannon</u> (URL) vs <u>FASSE</u> (URL)
  - Memory (RAM): amount of memory in GB
  - Number of cores: recommended at least 2
  - Number of GPUs: if >= 1, make sure you **select** a gpu partition
  - Allocated time: time you would like your session to run
  - Email for status notification: to know when job starts, ends
  - Reservation: if you have a special reservation (this requires approval from FASRC)
  - Account: use this if you have more than one PI\_lab affiliation

the minimum and/or maximum values of each field depends on the selected partition





# Jupyter Notebook

- Launch new Jupyter Notebook session (existing session will not work!)
- $\circ$   $\,$  Select newly created conda environment as the kernel
  - a. Open a notebook
  - b. On the top menu, click Kernel -> Select Kernel -> Click on OOD\_env
  - c. Note: kernels is the same as conda, python, mamba environment







# Closing running OOD windows/tabs

- In most OOD apps, you can close the browser tab while the code is running, and the code will continue to run on the background
- Jupyter Notebook will not! The cell that is running will lose the data and output files will not be written
  - Solution: run Remote Desktop app and launch Jupyter Notebook from within Remote Desktop
  - Documentation: <u>Open OnDemand (OOD/VDI) Remote Desktop: How to open</u> <u>software – FASRC DOCS</u>





# FASRC documentation

- FASRC docs: <u>https://docs.rc.fas.harvard.edu/</u>
- FASRC Python docs:
  - <u>https://docs.rc.fas.harvard.edu/kb/python/</u>
  - <u>https://docs.rc.fas.harvard.edu/kb/python-package-installation/</u>
- GitHub User\_codes: <u>https://github.com/fasrc/User\_Codes/</u>
- Getting help
  - Office hours: <a href="https://www.rc.fas.harvard.edu/training/office-hours/">https://www.rc.fas.harvard.edu/training/office-hours/</a>
  - Ticket
    - Portal: <u>http://portal.rc.fas.harvard.edu/rcrt/submit\_ticket</u> (requires login)
    - Email: <u>rchelp@rc.fas.harvard.edu</u>





# FASRC Upcoming Trainings

Training calendar: <a href="https://www.rc.fas.harvard.edu/upcoming-training/">https://www.rc.fas.harvard.edu/upcoming-training/</a>

#### Python Multiprocessing on the FASRC cluster

Training is focused on some of the techniques to accelerate Python programming with emphasis on utilizing multiprocessing with numpy arrays.

**Audience**: Users who are familiar with basic Python, command line, HPC systems, and have attended our Python Basics on FASRC clusters training.

Note: All topics below are a brief overview to utilizing multiprocessing on FASRC clusters.

#### **Objectives**:

- 1. Understanding Multiprocessing
- 2. Executing Multiprocessing on FASRC clusters





#### Training session evaluation

Please, fill out our training session evaluation. Your feedback is essential for us to improve our trainings!!

https://tinyurl.com/FASRC-training









#### **Thank you :)** FAS Research Computing