

Snakemake for reproducible research

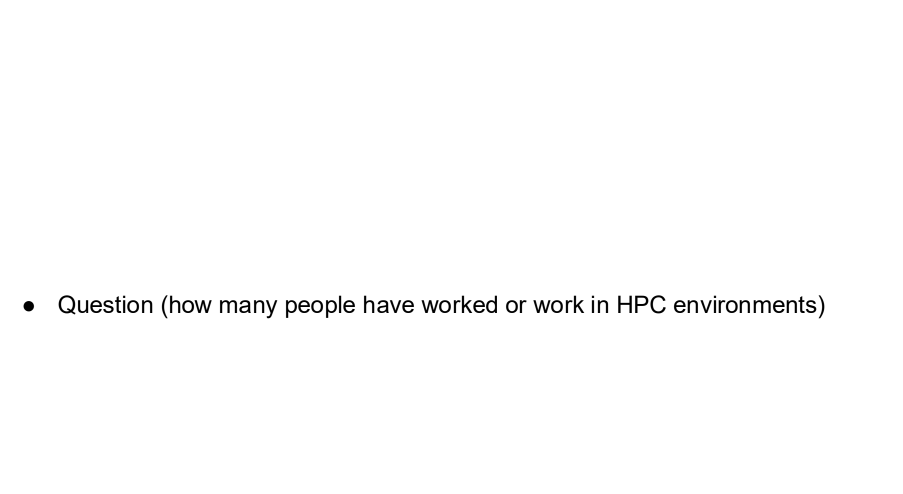
Running Snakemake in an HPC environment







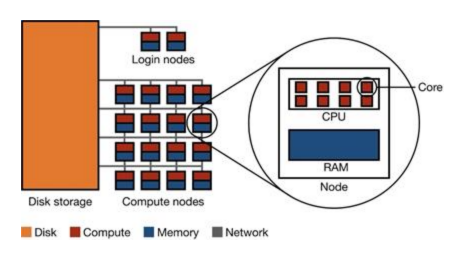




Question (are you familiar with slurm)

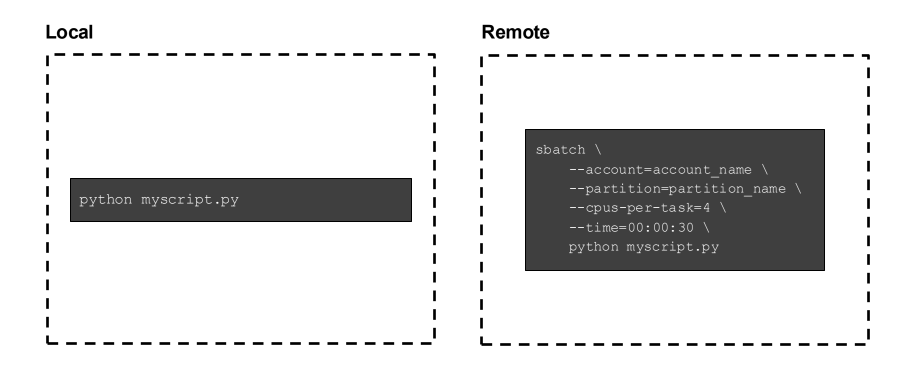
HPC environments

- Aggregated computing resources to gain performance greater than that of a single workstation, server, or computer.
- Used to run computationally heavy processes.
- Commonly used simultaneously by multiple users
- Job schedulers (i.e. SLURM) manage jobs sent by all users to ensure a safe and efficient use of the resources.
- Extra configuration required:
 - O RAM usage, CPUs, run time, ...

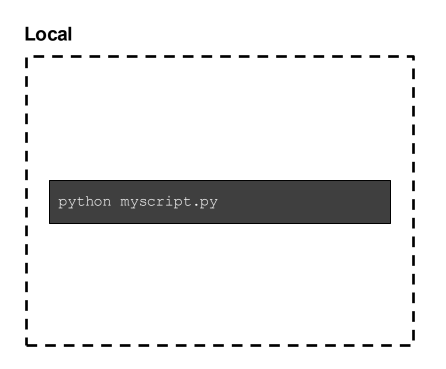


Statistical Computing and Communication https://ekatsevi.github.io/statistical-computing/hpc-basics.html

Local versus remote execution



Local versus remote execution



Remote

```
jobscript.sh
#!/bin/bash
#SBATCH --job-name=test
#SBATCH --account=account name
#SBATCH --partition=partition name
#SBATCH --cpus-per-task=4
#SBATCH --time=00:00:30
python myscript.py
sbatch jobscript.sh
```

Running Snakemake in HPC environments

- Snakemake can interact with multiple schedulers to run on clusters and cloud:
 - AWS
 - Azure
 - Flux
 - Google Batch
 - HTCondor
 - Kubernetes
 - LSF
 - Slurm
- Almost no changes required to the rules
 - Scheduler command can take job information from rule definition
 - One key parameter: maximum number of jobs running in parallel: -j / --jobs
- Implemented with:
 - v7 and before: --cluster "<scheduler_name>" in the Snakemake command
 - v8+: install <u>plugins</u> then --executor "<scheduler_name>" in the Snakemake command

Towards HPC execution

- Checking rule resource requirements ———— Benchmark directive
- Resource optimisation → Resources directive

Checking rule resource requirements: benchmarks

 'benchmark' is a directive; its value is a path to a benchmark results file for a rule

```
rule rename_file:
    input:
        'data/test.txt'
    output:
        'results/renamed_file.txt'
    benchmark:
        'benchmarks/renaming.txt'
    shell:
        'mv {input} {output}'
```

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- 'benchmark' is a directive; its value is a path to a benchmark results file for a rule
- Snakemake will measure runtime and memory usage for the rule and save it to the file

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- Snakemake will measure runtime and memory usage for the rule and save it to the file
- Benchmark files must have the same wildcards as the output!
- Best practice: put all benchmarks in same folder

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    output:
        'results/renamed_file.txt'
    benchmark:
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    shell:
        'mv {input} {output}'
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Towards HPC execution

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Optimising resource usage: memory and runtime

- 'resources' is a directive; its values set the resources available for a job
 - New kind of directive value: pair of <key>=<value>

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mem <unit>

- Amount of memory needed by the job
- o <unit>: mb, gb, tb...

runtime <unit>

- Amount of wall clock time a job needs to run
- <unit>: s, m, h, d...

```
rule example:
    input:
    output:
    resources:
        mem gb = 1,
        runtime h = 1
    shell:
```

Specifying job resources in Snakemake

- Job resources are determined as follows:
 - Specifying them by using the resources directive.
 - Using default values when no resources specified:
 - RAM: max(2*input.size mb, 1000)
 - Disk space: max(2*input.size_mb, 1000)
 - Temporary directory: system's tempdir
- Default resources can also be extended when calling Snakemake (i.e. slurm account)

```
rule myrule:
    input:
        'input_{file}.txt'
    output:
        'output_{file}.txt'
    resources:
        mem_mb: 100
    shell:
        'cat {input} > {output}'
```

Rule-specific settings in the Snakefile

- Some jobs are so small that it would be wasteful (and would take longer) to execute on an HPC
- You can define local execution rules using:
 - localrules keyword

```
localrules: light
rule light:
    input: 'input.txt'
    output: 'light output.txt'
    resources:
       mem mb: 100
    shell:
rule heavy:
    input: light.output
    output: 'heavy output.txt'
    resources:
    shell:
```

Rule-specific settings in the Snakefile

- Some jobs are so small that it would be wasteful (and would take longer) to execute on an HPC
- You can define local execution rules using:
 - localrules keyword
 - localrule directive

```
rule light:
    input: 'input.txt'
    output: 'light output.txt'
    resources:
                 100
   localrule: True
    shell:
rule heavy:
    input: light.output
    output: 'heavy output.txt'
    resources:
    shell:
```

Configuration profiles

Configuration profiles

- Preconfigured configuration parameters: resources, executor, sdm...
 - Can manage executor parameters as well:
 - Scripts to submit jobs
 - Scripts to check job status
 - Advanced customisation
- Currently, there are two types of profile:
 - Global: directory stored in ~/.config/snakemake/<profile name>/
 - Workflow-specific: directory named profile_name> and containing a config.yaml file.
- The directory contains config files in YAML format.
- Official list of Snakemake profiles <u>here</u>

Configuration profiles

```
./
├─ input_data/
├─ Snakefile
├─ config.yaml
└─ myprofile/
└─ config.yaml
```

```
executor: cluster-generic
cluster-generic-submit-cmd: \sbatch -job-name={rule} -cpus-per-task={threads}'
jobs: 10
```

- executor: used to indicate how to communicate with the scheduler
 - o **cluster-generic** is a Snakemake plugin that handles communication with the scheduler
- cluster-generic-submit-cmd: command to use to run the jobs. In the case of SLURM, this command is sbatch followed by the arguments you want to use
- jobs: used to indicate the maximum amount of jobs to run simultaneously

Running Snakemake using a profile

Once set up, running Snakemake using a profile is as simple as:

```
snakemake --profile <path_to_profile_folder>
```