Introduction to Snakemake

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Lesson 1

- 1. Why Snakemake?
- 2. Introducing Workflows
- 3. Workflow Syntax
- 4. Running Snakemake
- 5. Assignment 1

Preparation for Assignment 1:

```
sbatch -c 1 --mem 4G --wrap \
"conda create -n smk-lesson-1 -c conda-forge -c bioconda prokka pirate"
```

Why Snakemake?

Automation

Reproducibility

Others' Snakefiles

Front-loading your effort.

Modest investment at the beginning of a project yields a hands-off tool for performing routine analyses.

Why Snakemake?

Automation

Reproducibility

Others' Snakefiles

Guarantee that the same inputs will give the same outputs.

Altering any input will make Snakemake re-evalutate the outputs.

Built-in version tracking.

Why Snakemake?

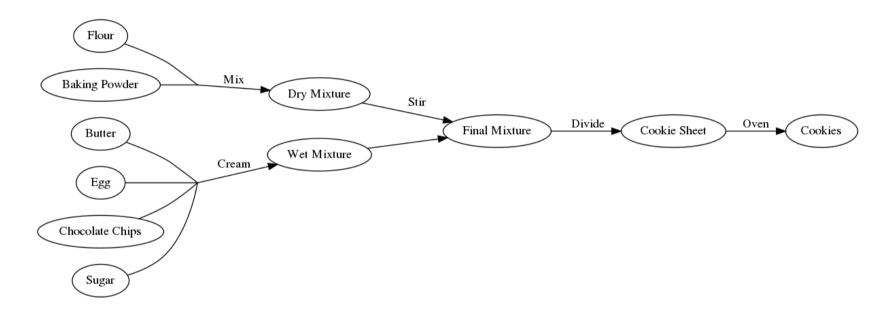
Automation

Understand and modify the tools others have created for you.

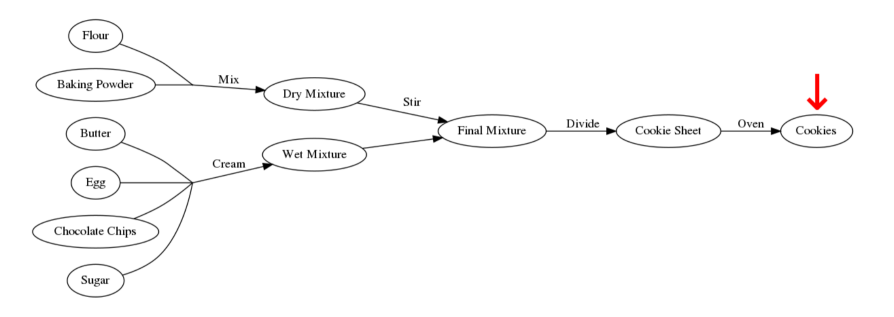
Reproducibility

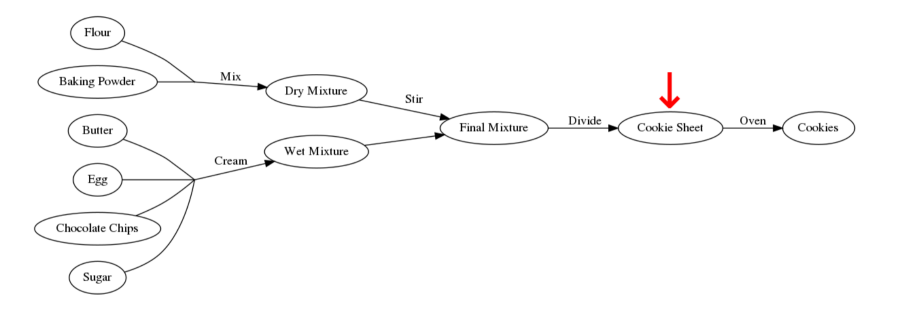
Others'
Snakefiles

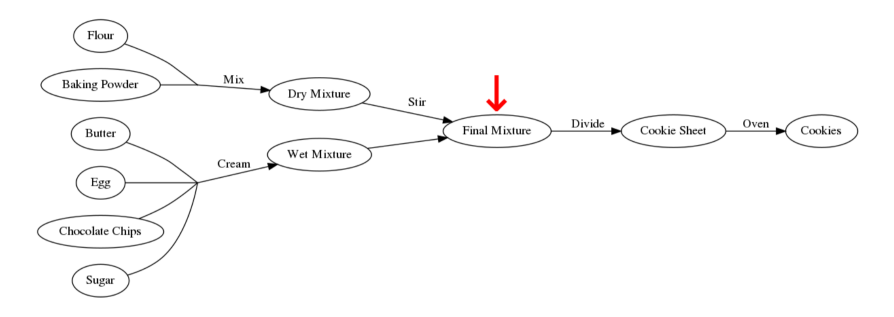
Baking with Graphs

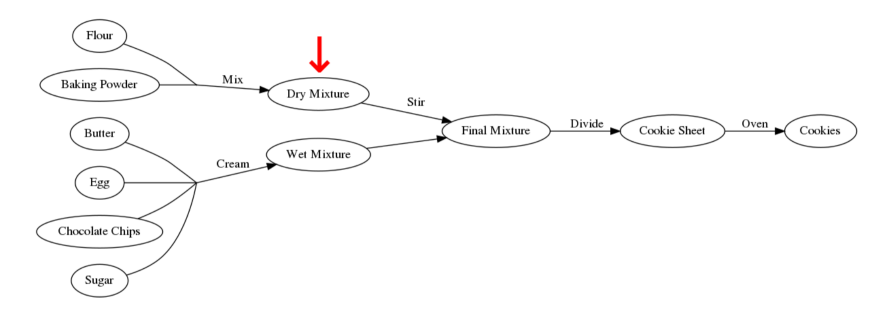


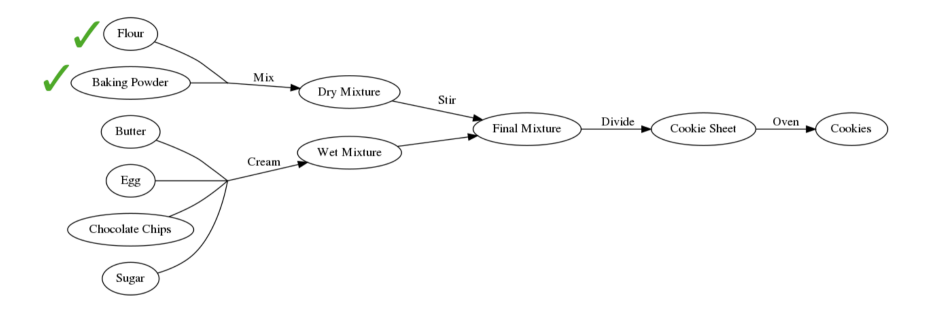
- Snakemake figures out how to achieve the desired result:
 - starts at the final product
 - works backwards until it finds what it needs
- A collection of dependencies not a sequence of instructions
- You tell it how to convert each input to each output

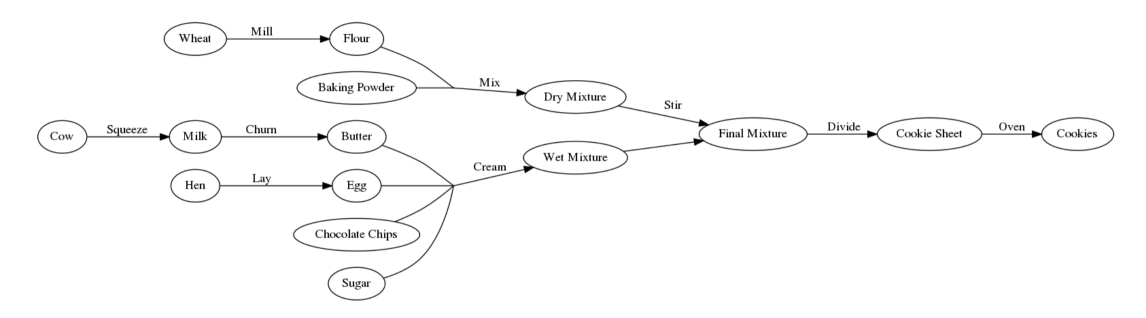








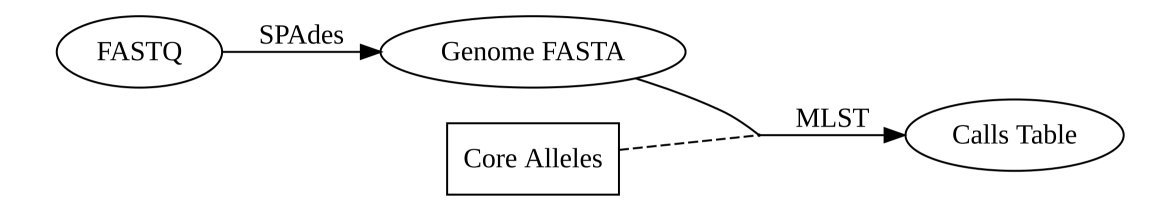




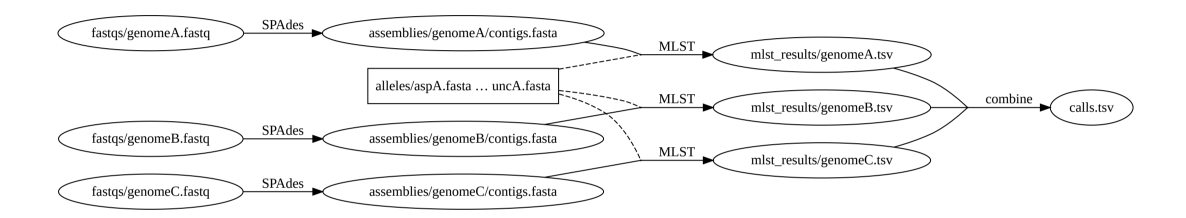
Rules for Baking

```
rule all:
    input: "cookies"
rule bake_cookies:
    input: "pan/dough.blobs"
    output: "cookies"
    shell: "oven -i {input} -o {output} --temp 350 --time 15"
rule apportion_dough_blobs:
    input: "bowls/final.mix"
    output: "pan/dough.blobs"
    shell: "scoop -n 24 {input} > {output}"
rule combine bowls:
    input: wet="bowls/wet.mix", dry="bowls/dry.mix"
    output: "bowls/final.mix"
    shell: "mixer {input.wet} {input.dry} > {output}"
```

Baking \rightarrow **Bioinformatics**



Multiple Samples



Wildcards

- We can match every file with particular naming pattern with wildcards
- In a rule, wrap a variable name with curly braces

```
• e.g. {sample}
```

- Rule will be applied in parallel to each file matching the rule
- In shell block, you can access these when preceded by wildcards

```
• e.g. {wildcards.sample}
```

Rules for Multiple Samples

```
from pathlib import Path
sample_names = [fq.stem for fq in Path("fastqs").glob("*")]
rule all:
    input: "calls.tsv"
rule assemble:
    input:
        fwd="fastqs/{sample}/{sample} 1.fastq", rev="fastqs/{sample}/{sample} 2.fastq"
    output: "assemblies/{sample}/contigs.fasta"
    shell: "spades -1 {input.fwd} -2 {input.rev} -o assemblies/{wildcards.sample}"
rule get sequence types:
    input: "assemblies/{sample}/contigs.fasta"
    output: "mlst results/{sample}.tsv"
    shell: "mlst --scheme campylobacter {input} > {output}"
rule combine mlst results:
    input: expand("mlst results/{sample}.tsv", sample=sample names)
    output: "calls.tsv"
    shell: "cat {input} > {output}"
```

Threads

- Many (but not all!) bioinformatics tools use multiple CPU threads
- threads directive defaults to 1
 - Accessible in the shell block, similar to input and output
 - {threads}

```
rule annotate_genome:
    input: "genomes/{sample}.fasta"
    output: "annotations/{sample}/{sample}.gff"
    threads: 8
    shell: "prokka --cpus {threads} -o annotations/{wildcards.sample} {input}"
```

Caveats and Assumptions

Directory Structure

- Snakefiles themselves can live anywhere
 - I keep mine in ~/snakefiles/
- Workflows tightly coupled to any directory structure described within
- Will implicitly create any directories it needs
 - No need for mkdir

Directory Structure

This rule...

```
rule annotate_genome:
    input: "genomes/{sample}.fasta"
    output: "annotations/{sample}/{sample}.gff"
    threads: 8
    shell: "prokka --cpus {threads} -o annotations/{wildcards.sample} {input}"
```

requires this structure:

```
analysis/
|— annotations/
| — isolateA.fasta
| — isolateB.fasta
| — isolateC.fasta
| — isolateA.fasta
| — isolateB.fasta
| — isolateB.fasta
| — isolateC.fasta
```

Caveats and Assumptions

Independent Jobs

- Failure of any job will abort all other jobs
 - Override with --keep-going
 - Dependent jobs will still await all inputs
 - *e.g.* if stiring the dry cookie mixture fails, the wet mixture still gets made, but nothing goes in the oven

Running Snakemake (Basic)

The **basic invocation** of Snakemake:

```
snakemake --jobs <number of parallel jobs> -s <path to your Snakefile> -d <work directory>
```

Example populated with real values:

```
snakemake --jobs 5 -s ~/snakefiles/assemble.smk -d ~/Projects/cj_population_study
```

Running Snakemake on Waffles

- Snakemake can be run on HPCs like Waffles
 - Must be combined with Slurm
 - Don't run it on the head node!
- Two parts:
 - 1. Tell Snakemake how to submit jobs with --cluster
 - 2. Submit Snakemake itself as a Slurm job

Running Snakemake on Waffles

The --cluster argument:

- Create a template command to pass to Slurm
- May access Snakemake special variables like {threads}
 - More on this later

```
--cluster 'sbatch -c {threads} --mem 12G --partition NMLResearch '
```

Running Snakemake on Waffles

Submitting the Snakemake job to Slurm:

```
sbatch -c 1 --mem 4G --wrap "snakemake --jobs 5 -s
~/snakefiles/assemble.smk -d ~/Projects/cj_population_study --cluster
'sbatch -c {threads} --mem 12G --partition NMLResearch '"
```

Assignment 1

Write a Snakemake workflow that does the following:

- 1. Run Prokka on each genome
- 2. Symlink GFF annotations into gffs/
- 3. Build a pangenome with PIRATE

conda activate smk-lesson-1

Lesson 2

Lesson 2

- 1. Assignment 1 Answers
- 2. Conda Integration
- 3. Params and Threads
- 4. Mixing in Python
- 5. Assignment 2

Assignment 1

Conda Integration

- Snakemake can manage conda directly
- No need to manually build or activate conda environments

Conda directive

```
rule annotate_genome:
    input: "genomes/{sample}.fasta"
    output: "annotations/{sample}/{sample}.gff"
    conda: "envs/prokka.yaml"
    shell: "prokka --cpus {threads} -o annotations/{wildcards.sample} {input}"
```

Conda YAML files

• Placed **relative to the Snakefile**, *not* the project directory

```
# annotate.smk
rule annotate_genome:
    input: "genomes/{sample}.fasta"
    output: "annotations/{sample}/{sample}.gff"
    conda: "envs/prokka.yaml"
    shell: "prokka --cpus {threads} -o annotations/{wildcards.sample} {input}"
```

The above will look for the following directory structure:

```
snakefiles/
— annotate.smk
— envs
— prokka.yaml
```

Conda YAML files

This YAML file ...

```
name: prokka
channels:
    - conda-forge
    - bioconda
    - defaults
dependencies:
    - prokka
```

... is equivalent to this conda command:

```
conda create -n prokka -c conda-forge -c bioconda -c defaults prokka
```

Using Conda Directives with Snakemake

Must explicitly tell Snakemake to use Conda

```
snakemake --use-conda <...>
```

Automatic installation and activation

Config

Configuration is possible through config

- Python dict available within the Snakefile
- Available through two methods
 - --config passes arguments directly via command line
 - --configfile points to a YAML file that provides values

--config "key=value" is equivalent to --configfile config.yaml where...

```
# config.yaml
key: "value"
```

Configuration via:

--config flag:

- ↓ effort
- ↑ flexible
- \upsilon reproducible

YAML file:

- ↑ effort
- | flexible
- ↑ reproducible

Params

• Non-file parameters may be provided in the params directive

Abusing Params to Fine-tune Resources

```
snakemake <...> --cluster 'sbatch -c {threads} --mem {params.mem} --time {params.time} '
rule annotate_genome:
    input: "genomes/{sample}.fasta"
    output: "annotations/{sample}/{sample}.gff"
    threads: 8
    params:
        time="45:00",
        mem="16G"
    shell: "prokka --cpus {threads} -o annotations/{wildcards.sample} {input}"
rule symlink_gff:
    input: "annotations/{sample}/{sample}.gff"
    output: "gffs/{sample}.gff"
    threads: 1
    params:
        time="01:00",
        mem="100M"
    shell: "In -sr {input} {output}"
```

Config vs Params

- Params are fairly "fixed"
 - Used primarily to simplify shell block
- Config for run-specific information
 - e.g. providing a particular host database to kat or training file to chewBBACA

Mixing in Python

- Python may be mixed in arbitrarily into Snakemake
 - *i.e.* All Python is valid Snakemake
- Two main ways of using Python in Snakemake
 - o run blocks
 - Python used directly in the Snakemake file

Run blocks

- run blocks can be used in place of shell blocks
- Write Python inside the run block, rather than Bash in a shell block
- May access snakemake values like input and output

```
rule transpose_table:
    input: "data/results_table.csv"
    output: "data/results_table_transposed.csv"
    run:
        import pandas as pd
        original = pd.read_csv(input[0], header=0)
        transposed = original.transpose()
        transposed.to_csv(output[0], header=False)
```

Directly Using Python in Snakemake

- You can directly use Python in Snakemake
- Particularly useful for handling cases where a rule generates variable output
 - *e.g.* The number of gene FASTAs generated by a pangenome analysis
- Can provide a Python function to input instead of a file pattern

- select_high_quality_genomes takes a list of FASTAs, then symlinks high-quality ones into ./good_genomes/ and writes a report called quality_report.txt
- We don't know in advance which genomes will pass QC, so we need an input function

```
rule quality filter genomes:
    input: expand("genomes/{sample}.fasta", sample=samples)
    output: "quality report.txt"
    shell: "select high quality genomes {input} > {output}"
# input functions need to take parameter `wildcards`
def collect_good_genome_sample_names(wildcards):
    good genomes = Path("good genomes/").glob("*.fasta")
    return list(good genomes)
# use the report as a dummy input to make sure quality_filter_genomes executes
rule run abricate:
    input: report="quality report.txt", fastas=collect good genome sample names
    output: "amr results.tsv"
    shell: "abricate {input.fastas} > {output}"
```

Assignment 2

- Build upon Assignment 1
- 1. Create conda YAMLs for prokka and pirate
- 2. Give appropriate resources to each rule with params
- 3. Write a rule with a run block that reads PIRATE.gene_families.tsv, finds loci present in 100% of genomes, and writes their names to a text file
 - columns of interest: gene_family & number_genomes
- 4. A rule that:
 - Uses an input function reads selected genes from the text file in Part 3
 - Symlinks these into a directory called loci
 - Either shell or run at your preference

Assignment 2 Hints

pandas for easily reading tabular files

```
import pandas as pd
data_table = pd.read_csv(input[0], sep = "\t")
# select rows from columnA where columnC is greater than 42
selected_rows = data_table["columnC"] > 42
selected_columnA = data_table["columnA"].loc[selected_rows]
```

Creating symlinks from a list of file basenames

```
# list_of_names = ["larry", "moe", "curly"]
import os
for name in list_of_names:
    src = f"originals/{name}.txt"
    dst = f"filtered/{name}.txt"
    os.symlink(src, dst)
```

Assignment 2 Hints

Reading a text file into a list with Python

Consider combining functions like this with expand()

```
o expand("path/to/{sample}.txt", sample=read_list_to_list())
```

```
def read_lines_to_list(path: str):
    lines = []
    with open(path, "r") as f:
        for line in f:
            trimmed_line = line.strip()
                  lines.append(trimmed_line)
    return lines
```

Reading a text file, but using at as an input function

```
def aggregate_files(path: str):
    lines = []
    with open(path, "r") as f:
        for line in f:
            trimmed_line = line.strip()
            lines.append(trimmed_line)
        lines_with_paths = [f"path/to/{sample}.txt" for sample in lines]
    return lines_with_paths
```

```
rule do_something:
   input: aggregate_files
   output: "somefile.txt"
   shell: "my_program {input} -o {output}"
```

Lesson 3

Lesson 3

- 1. Assignment 2 Answer
- 2. Fixing when things go wrong
- 3. Priorities
- 4. Script files (Python, R, and Julia)
- 5. Visualizing Workflows
- 6. Clinic

Assignment 2

When Things Go Wrong

- Snakemake locks its working directory when running
 - Prevents other snakemake instances from running in the same place
- Snakemake removes the lock when it completes (success or failure)
- Lockfile may not be removed when if snakemake crashes or is killed by slurm

```
o scancel --user your_username
```

Unable to lock working directory.

Unlocking a Stale Lock

The Correct Way: --unlock

```
sbatch -c 1 --mem 2G --wrap "snakemake --unlock -s path/to/your/workflow.smk -j 1"
```



```
rm -r ./.snakemake
```

Priorities

- Rule priorities may be set
- These are not the same as slurm partitions
- Only determine the priority of execution within the workflow
- Rules default to priority 0
- Bigger numbers are higher priority

```
rule my_example:
  input: ...
  output: ...
  priority: 50
  shell: ...
```

Script Files