

Snakemake for reproducible research

Making a more general-purpose Snakemake workflow





Pop quiz

```
rule rename_file:
    input:
        rules.create_file.output
    output:
        'results/renamed_file.txt'
    shell:
        'mv {input} {output}'
```

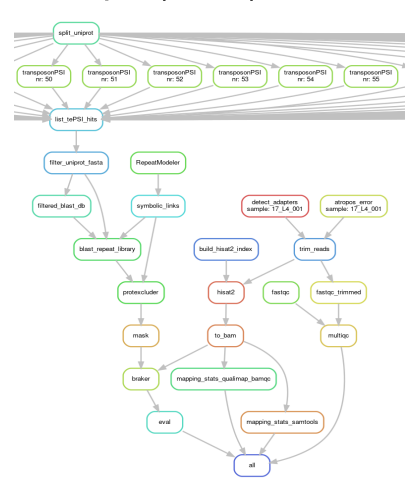
Pop quiz

- Snakemake keyword
- Rule name (user-defined)
- Snakemake directives
- Directives values:
 - Object
 - String (file path)
 - Instruction (command)
 - Numeric values (seen later)
- Mystery syntax?

```
rule rename_file:
    input:
        rules.create_file.output
    output:
        'results/renamed_file.txt'
    shell:
        'mv {input} {output}'
```

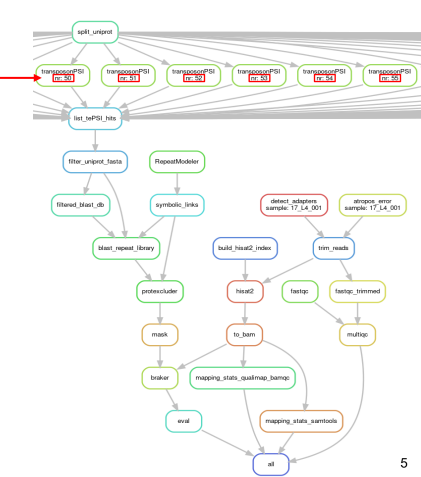
Building a Directed Acyclic Graph (DAG)

 Snakemake determines which jobs to run to produce desired outputs



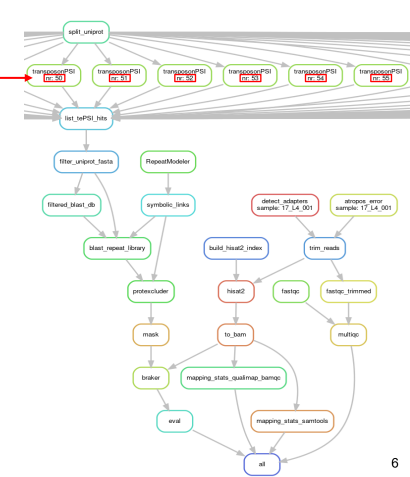
Building a Directed Acyclic Graph (DAG)

- Snakemake determines which jobs to run to produce desired outputs
- Rule can appear more than once, with different wildcards
 - 1 rule + 1 wildcard values = 1 job
- Arrows = dependency between jobs
 - Snakemake runs jobs in any order that doesn't break dependency



Building a Directed Acyclic Graph (DAG)

- Snakemake determines which jobs to run to produce desired outputs
- Rule can appear more than once, with different wildcards
 - 1 rule + 1 wildcard values = 1 job
- Arrows = dependency between jobs
 - Snakemake runs jobs in any order that doesn't break dependency
- DAG = work list, ≠ flowchart
 - No if/else decisions or loops
 - Snakemake runs every job in the DAG exactly once
- DAG ≠ checking shell directives
 - Shell commands are tested during execution
 - Works? Produces expected outputs?



What is a DAG useful?

- Skip parts of the DAG to avoid recomputing → Save time and resources (CPU, memory, energy, money)
- Change/add inputs to existing analyses without re-running everything
- Resume running a workflow that failed part-way

What could we improve?

What could we improve?

- Using hard-coded file paths
- Having multiple inputs/outputs per rule
- Checking Snakemake behaviour

What could we improve?

- Having multiple inputs/outputs per rule ——— Numbered/named inputs/outputs
- Checking Snakemake behaviour Log files, benchmarks

Placeholder:

- A person or thing that occupies the position or place of another person or thing
- A symbol in a mathematical or logical expression that may be replaced by the name of any element of a set

(From the Merriam-Webster dictionary)

```
rule rename_file:
    input:
        'data/test.txt'
    output:
        'results/renamed_test.txt'
    shell:
        'mv data/test.txt results/renamed_test.txt'
```

```
rule rename_file:
    input:
        'data/test.txt'
    output:
        'results/renamed_test.txt'
    shell:
        'mv data/test.txt results/renamed_test.txt'
```

```
rule rename_file:
    input:
        'data/test.txt'
    output:
        'results/renamed_test.txt'
    shell:
        'mv {input} {output}'
```

- {input} and {output} are placeholders
- Used in shell directive
- Similar to python f-string
- Snakemake will replace them with appropriate values before running the command
- Many directives can use placeholders: {log}, {benchmark}, {params}...

```
rule rename_file:
    input:
        'data/test.txt'
    output:
        'results/renamed_test.txt'
    shell:
        'mv {input} {output}'
```

 Wildcards ≈ "variables" automatically inferred by Snakemake

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```
rule rename_file:
    input:
        'data/test.txt'
    output:
        'results/renamed_test.txt'
    shell:
        'mv {input} {output}'
Defined paths
```

 Wildcards ≈ "variables" automatically inferred by Snakemake

```
rule rename file:
    input:
    output:
                                        Defined paths
    shell:
rule rename file:
    input:
                                        Adaptable paths
    output:
                                         with wildcards
    shell:
```

- Wildcards ≈ "variables" automatically inferred by Snakemake
- Enclose wildcard name with curly brackets {}

```
rule rename file:
    input:
    output:
                                         Defined paths
    shell:
rule rename file:
    input:
                                         Adaptable paths
    output:
                                         with wildcards
    shell:
```

- Wildcards are "resolved" from the target and propagated to other directives
 - Regular expression matching: .+
 - "1 or more occurrences of any character except newline"
 - Can be constrained
- Using wildcards forces to ask for output(s): Snakemake doesn't guess!
 - Target rules may not contain wildcards

```
rule rename_file:
    input:
        'data/{file}.txt'
    output:
        'results/renamed_{file}.txt'
    shell:
        'mv {input} {output}'
```

```
snakemake --cores 1 results/renamed_test.txt

{file} = "test"

input: 'data/test.txt'
```

- Wildcards are "resolved" from the target and propagated to other directives
 - Regular expression matching: .+
- Both a workflow and a rule can use multiple wildcards

```
rule rename_file:
    input:
        'data/{file}_{nb}.txt'
    output:
        'results/renamed_{file}_{nb}.txt'
    shell:
        'mv {input} {output}'
```

```
snakemake --cores 1 results/renamed_test_1.txt

{file} = "test"; {nb} = "1"

input: 'data/test_1.txt'
```

- Wildcards are "resolved" from the target and propagated to other directives
 - Regular expression matching: .+
- Both a workflow and a rule can use multiple wildcards
- Input and output files do not need to share the same wildcards
- All outputs, logs... created by a rule must have the same wildcards!

```
rule rename_file:
    input:
        'data/{file}.txt'
    output:
        'results/renamed_{file}_{nb}.txt'
    shell:
        'mv {input} {output}'
```

Rules can use multiple inputs/outputs

- Rules can use multiple inputs/outputs
 - Separated by a comma
 - Input values are unpacked (replaced by a space-separated list)

```
rule gather_files:
    input:
        'data/test1.txt',
        'data/test2.txt'
    output:
        'results/merged_test.txt'
    shell:
        'cat {input} > {output}'
```

```
shell:
    'cat data/test1.txt data/test2.txt > results/merged_test.txt
```

- Rules can use multiple inputs/outputs
 - Separated by a comma
 - Input values are unpacked (replaced by a space-separated list)
- Shell can have multiple commands
 - Separated by a semicolon
 - Commands are concatenated

```
rule gather_files:
    input:
        'data/test1.txt',
        'data/test2.txt'

output:
        'results/merged_test.txt'
shell:
        'cat {input} > {output}';
        'cat {input} >> {output}'
```

- Rules can use multiple inputs/outputs
 - Separated by a comma
 - Input values are unpacked (replaced by a space-separated list)
- Shell can have multiple commands
 - Separated by a semicolon
 - Commands are concatenated

```
rule gather files:
    input:
        'data/test1.txt',
    output:
    shell:
```

- Rules can use multiple inputs/outputs
 - Separated by a comma
 - Input values are unpacked (replaced by a space-separated list)
- Shell can have multiple commands
 - Separated by a semicolon
 - Commands are concatenated
- Inputs can be accessed by their positional index: input[n]
 - Numbering starts at 0

```
rule gather files:
    input:
        'data/test1.txt',
    output:
    shell:
```

- Rules can use multiple inputs/outputs
 - Separated by a comma
 - Input values are unpacked (replaced by a space-separated list)
- Shell can have multiple commands
 - Separated by a semicolon
 - Commands are concatenated
- Inputs can be accessed by their positional index: input[n]
 - Numbering starts at 0
- Named inputs can be accessed by their names: input.input_name
 - You cannot mix named and unnamed inputs

```
rule gather files:
    input:
        file 1='data/test1.txt',
        file 2='data/test2.txt'
    output:
    shell:
```

- Outputs work like inputs
 - Separated by ','
 - Can be named
 - Can be accessed by positional index or by name
- All outputs need to be created or the job will fail

```
rule gather files:
    input:
        file 1='data/test1.tsv',
        file 2='data/test2.tsv'
    output:
        copy 1='results/copied test1.txt',
        copy 2='results/copied test2.txt'
    shell:
```

snakemake --cores 1 results/first step 1.txt



Checking Snakemake behaviour

- Producing log files
- Benchmarking rules

Checking Snakemake behaviour: log files

- 'log' is a directive; its value is a path to a log file for one rule
 - Can be accessed with a placeholder in shell: {log}

```
rule rename_file:
    input:
        'data/test.txt'
    output:
        'results/renamed_file.txt'
    log:
        'logs/renaming.log'
    shell:
        'mv {input} {output} 2> {log}'
```

Checking Snakemake behaviour: log files

- 'log' is a directive; its value is a path to a log file for one rule
 - Can be accessed with a placeholder in shell: {log}
- You need to manually redirect messages to logs, but Snakemake automatically creates the folder path

```
rule rename_file:
    input:
        'data/test.txt'
    output:
        'results/renamed_file.txt'
    log:
        'logs/renaming.log'
    shell:
        'mv {input} {output} 2> {log}'
```

Checking Snakemake behaviour: log files

- 'log' is a directive; its value is a path to a log file for one rule
 - Can be accessed with a placeholder in shell: {log}
- You need to manually redirect messages to logs, but Snakemake automatically creates the folder path
- Log files must have the same wildcards as the output!
- Good practice: put all logs in same folder

```
rule rename_file:
    input:
        'data/test.txt'
    output:
        'results/renamed_file.txt'
    log:
        'logs/renaming.log'
    shell:
        'mv {input} {output} 2> {log}'
```

Checking Snakemake behaviour: 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}'
```

Checking Snakemake behaviour: benchmarks

- '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

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

Checking Snakemake behaviour: benchmarks

- '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
- Benchmark files must have the same wildcards as the output!
- Best practice: put all benchmarks in same folder

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

Exercises

- Through the day:
 - Develop a simple RNAseq analysis workflow, from reads (fastq files) to Differentially Expressed Genes
 (DEG)
- For this session:
 - Use placeholders and wildcards
 - Use multiple inputs and outputs
 - Check workflow behaviour
 - Visualise a DAG