

Snakemake for reproducible research

Making a more general-purpose Snakemake workflow



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Pop quiz

```
rule second_step:
   input:
      rules.first_step.output
   output:
      'results/first_step.txt'
   shell:
      'cp {input} {output}'
```

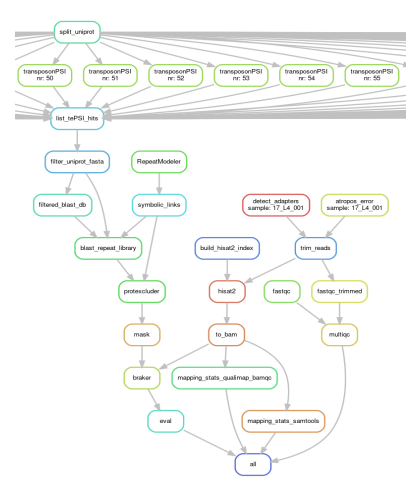
Pop quiz

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

```
rule second_step:
    input:
        rules.first_step.output
    output:
        'results/first_step.txt'
    shell:
        'cp {input} {output}'
```

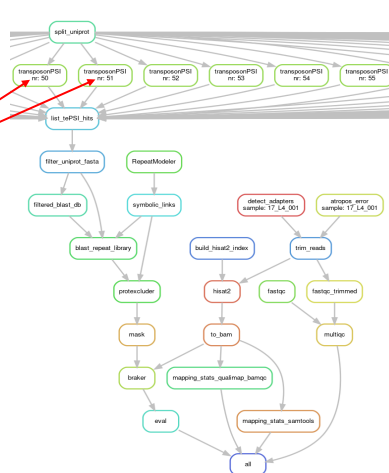
Building a Directed Acyclic Graph (DAG)

- Snakemake determines which jobs to run to produce desired outputs
 - DAG depends on Snakefile, requested target outputs, and files already present



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 - DAG depends on Snakefile, requested target outputs, and files already present
- 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
 - DAG depends on Snakefile, requested target outputs, and files already present
- 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 does not check shell directives
 - Shell commands are tested during execution (1. Works? 2. Produces expected outputs?)



DAG (re-)run policy

- Snakemake runs a job if:
 - Target file explicitly requested is missing
 - Intermediate file is missing and needed to create target file
 - Input file is newer than an output file (timestamps comparison)
 - Can skip parts of the DAG
- Allows to:
 - Change/add inputs to existing analysis without re-running everything
 - Resume running a workflow that failed part-way

DAG (re-)run policy

- Snakemake runs a job if:
 - Target file explicitly requested is missing
 - Intermediate file is missing and needed to create target file
 - Input file is newer than an output file (timestamps comparison... but not only)
 - Can skip parts of the DAG
- Allows to:
 - Change/add inputs to existing analysis without re-running everything
 - Resume running a workflow that failed part-way
- Altering DAG (re-)run policy:
 - o -f, --force <target_name>
 - o -F, --forceall
 - -R, --forcerun <rule_name>
 - --rerun-triggers {mtime,params,input,software-env,code}
 - --touch

Several problems...

Several problems...

- Hard-coded file paths
- Processing list of files
- Only one input/output per rule
- Resources are not optimised

... that can be solved!

- Hard-coded file paths

 Placeholders and wildcards
- Only one input/output per rule
 Numbered/named inputs/outputs

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 example:
    input:
        'data/first_step.tsv'
    output:
        'results/first_step.txt'
    shell:
        'cp data/first_step.tsv results/first_step.txt'
```

```
rule example:
    input:
        'data/first_step.tsv'
    output:
        'results/first_step.txt'
    shell:
        'cp data/first_step.tsv results/first_step.txt'
```

```
rule example:
    input:
        'data/first_step.tsv'
    output:
        'results/first_step.txt'
    shell:
        'cp {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 used in placeholders: {log}, {benchmark}, {params}...

```
rule example:
    input:
        'data/first_step.tsv'
    output:
        'results/first_step.txt'
    shell:
        'cp {input} {output}'
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```
rule example:
    input:
        'data/first_step.tsv'
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        'results/first_step.txt'
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        'cp {input} {output}'
```

"Hard-coded" input and output files

```
rule example:
    input:
        'data/first_step.tsv'
    output:
        'results/first_step.txt'
    shell:
        'cp {input} {output}'
```

"Hard-coded" input and output files

```
rule example:
    input:
        'data/{sample}.tsv'
    output:
        'results/{sample}.txt'
    shell:
        'cp {input} {output}'
```

"General" input/output files with wildcards

```
rule example:
    input:
        'data/first_step.tsv'
    output:
        'results/first_step.txt'
    shell:
        'cp {input} {output}'
```

"Hard-coded" input and output files

```
rule example:
    input:
        'data/{sample}.tsv'
    output:
        'results/{sample}.txt'
    shell:
        'cp {input} {output}'
```

"General" input/output files with wildcards

- Enclose wildcard name with curly brackets { }
- How does Snakemake execution work when there are wildcards?

- Wildcards ≈ Snakemake "variables"
- 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 example:
    input:
        'data/{sample}.tsv'
    output:
        'results/{sample}.txt'
    shell:
        'cp {input} {output}'
```

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

Snakemake interpretation:
{sample} = "first_step"
```

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

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

Snakemake interpretation:

```
{sample} = "first"
{treatment} = "step"
```

```
rule example:
    input:
        'data/{sample}_{treatment}.tsv'
    output:
        'results/{sample}_{treatment}.txt'
    shell:
        'echo {wildcards.sample};'
        'cp {input} {output}'
```

- Wildcards ≈ Snakemake "variables"
- 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 have to share the same wildcards
- All outputs/logs... created by a rule must have same wildcards!

```
rule example:
    input:
        'data/{sample}.tsv'
    output:
        'results/{sample}_{treatment}.txt'
    shell:
        'echo {wildcards.sample};'
        'cp {input} {output}'
```

```
Snakemake --cores 1 results/first_step.txt

Snakemake interpretation:

input = 'data/first.tsv'
```

Rules can use multiple inputs/outputs

- Rules can use multiple inputs/outputs
- Don't forget the comma!

```
rule example:
    input:
        'data/first_step1.tsv',
        'data/first_step2.tsv'
    output:
        'results/first_step.txt'
    shell:
        'cat {input} > {output}'
```

- Rules can use multiple inputs/outputs
- Don't forget the comma!

Input directive values are unpacked (replaced by a space-separated list)

```
rule example:
    input:
        'data/first_step1.tsv',
        'data/first_step2.tsv'
    output:
        'results/first_step.txt'
    shell:
        'cat {input} > {output}'
```

```
shell:
   cat data/first_step_1.tsv data/first_step_2.tsv > results/first_step.txt
```

- Rules can use multiple inputs/outputs
- Don't forget the comma/semicolon!
- Inputs can be accessed by their positional index: input[n]
 - Numbering starts at 0

```
rule example:
    input:
        'data/first_step1.tsv',,
        'data/first_step2.tsv'
    output:
        'results/first_step.txt'
    shell:
        'cat {input[0]} > {output};'
         cat {input[1]} >> {output}'
```

Commands are concatenated

- Rules can use multiple inputs/outputs
- Don't forget the comma!
- Inputs can be accessed by their positional index: input[n]
 - Numbering starts at 0

```
rule example:
    input:
        'data/first_step1.tsv',
         'data/first_step2.tsv'
    output:
        'results/first_step.txt'
    shell:
         1 1 1
        cat {input[0]} > {output}'
        cat {input[1]} >> {output}'
         1 1 1
```

- Rules can use multiple inputs/outputs
- Don't forget the comma!
- Inputs can be accessed by their positional index: input[n]
 - Numbering starts at 0
- Named input can be accessed by their names: input.input_name
 - You cannot mix named and unnamed inputs

```
rule example:
    input:
        input_1='data/first_step1.tsv',
        input_2='data/first_step2.tsv'
    output:
        'results/first_step.txt'
    shell:
         1 1 1
        cat {input.input_1} > {output}'
        cat {input.input_2} >> {output}'
         1 1 1
```

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

```
rule example:
    input:
        input_1='data/first_step1.tsv',
        input_2='data/first_step2.tsv'
    output:
        output_1='results/first_step1.txt',,
        output_2='results/first_step2.txt'
    shell:
        I = I = I
        cat {input.input_1} > {output.output_1}'
        cat {input.input_2} > {output.output_2}'
        I - I - I
```

snakemake --cores 1 results/first_step_1.txt

results/first_step_1.txt, results/first_step_2.txt

- expand(): Snakemake function to automatically expand a wildcard expression to several wildcard values
 - Useful to define multiple inputs or outputs with a common pattern

- expand(): Snakemake function to automatically expand a wildcard expression to several wildcard values
 - Useful to define multiple inputs or outputs with a common pattern
 - Syntax: expand('{wildcard_name}', wildcard_name=<values>)
 - <values>: iterable (i.e. list, tuple, set) containing the wildcard values

```
rule example:
    input:
        'data/A.tsv',
        'data/B.tsv',
        'data/C.tsv'
    output:
        'results/total.tsv'
    shell:
        'cat {input} > {output}'
```

```
rule example:
    input:
        expand('data/{sample}.tsv',
    sample=['A,'B','C'])
    output:
        'results/total.tsv'
    shell:
        'cat {input} > {output}'
```

➤ The rule example uses all three input files to generate a single output file. expand() does not apply the rule separately to the three inputs!

When there are several wildcards, expand() creates all possible combinations

When there are several wildcards, expand() creates all possible combinations

```
samples=['A,'B']
replicates = [1, 2]

rule example:
    input:
        expand('data/{sample}_{replicate}.tsv', sample=samples, replicate=replicates)
    output:
        'results/total.tsv'
    shell:
        'cat {input} > {output}'
```

input = 'data/A_1.tsv data/A_2.tsv data/B_1.tsv data/B_2.tsv'

The wildcards in expand are INDEPENDENT from any other wildcard in the rule

The wildcards in expand are INDEPENDENT from any other wildcard in the rule

```
samples=['A,'B']
replicates = [1, 2]

rule example:
    input:
        expand('data/{sample}_{replicate}.tsv', sample=samples, replicate=replicates)
    output:
        'results/{sample}.tsv'
    shell:
        'cat {input} > {output}'
```

In this case, the value of the {sample} wildcard will NOT be propagated to the input

Optimising workflow performances

- Producing log files
- Benchmarking rules
- Multi-threading and controlling resource usage

Optimising workflow performances: log files

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

Optimising workflow performances: log files

- 'log' is a directive; its value is a path to a log file for a rule
 - Can be accessed with a placeholder in 'shell': {log}
- Logs still need to be handled manually for each command, but Snakemake automatically creates the directory in the log file path

```
rule example:
    input:
        'data/first_step.tsv'
    output:
        'results/first_step.txt'
    log:
        'logs/first_step.log'
    shell:
        'cp {input} {output} 2> {log}'
```

Optimising workflow performances: log files

- 'log' is a directive; its value is a path to a log file for a rule
 - Can be accessed with a placeholder in 'shell': {log}
- Logs still need to be handled manually for each command, but Snakemake automatically creates the directory in the log file path
- Log files must have the same wildcards as the output!
- Best to regroup logs in a 'logs' folder

```
rule example:
    input:
        'data/first_step.tsv'
    output:
        'results/first_step.txt'
    log:
        'logs/first_step.log'
    shell:
        'cp {input} {output} 2> {log}'
```

Optimising workflow performances: benchmarks

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

Optimising workflow performances: benchmarks

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

```
rule example:
    input:
        'data/first_step.tsv'
    output:
        'results/first_step.txt'
    benchmark:
        'benchmarks/first_step.txt'
    shell:
        'cp {input} {output}'
```

Optimising workflow performances: benchmarks

- 'benchmark' is a directive; its value is a path to a benchmark results file for a rule
- Snakemake will automatically 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 to regroup benchmarks in a 'benchmarks folder

```
rule example:
    input:
        'data/first_step.tsv'
    output:
        'results/first_step.txt'
    benchmark:
        'benchmarks/first_step.txt'
    shell:
        'cp {input} {output}'
```

Optimising workflow performances: threads

- 'threads' is a directive; its value is the number of threads to allocate to each job spawned by a rule
 - New kind of directive value: numeric (integer)
 - Check whether software can actually multithread!

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 - New kind of directive value: numeric (integer)
 - Check whether software can actually multithread!

```
rule example:
    input:
        'data/first_step.tsv'
    output:
        'results/first_step.txt'
    threads: 4
    shell:
        'command --threads {threads} {input} > {output}'
```

 In local mode, total number of threads allocated to Snakemake is constrained by the execution parameter '--cores'

Optimising workflow performances: resources

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

Optimising workflow performances: resources

- 'resources' is a directive; its values aim to set the resources available for a job
 - New kind of directive value: pair of <key>=<value>
- mem_<unit>
 - Amount of memory needed by the job
 - <unit>: mb, gb, tb...
- runtime_<unit>
 - Amount of wall clock time a job needs to run
 - <unit>: s, m, h, d...

```
rule example:
    input:
        'data/first_step.tsv'
    output:
        'results/first_step.txt'
    resources:
        mem_gb=1
        runtime_s=3600
    shell:
        'command {input} > {output}'
```

Exercises

- Through the day:
 - Develop a simple RNAseq analysis workflow, from reads (fastq files) to Differentially Expressed Genes (DEG)
- For now:
 - Session 2:
 - Use multiple inputs and outputs
 - Use placeholders and wildcards
 - Optimise workflow performance
 - Visualise a DAG
 - Session 3:
 - Use non-file parameters
 - Manage non-conventional outputs
 - Process list of inputs
 - Modularise a workflow