

Snakemake overview

Thomas Cokelaer

Institut Pasteur

Dec. 12th 2016 Journée Snakemake

S = M + P

Think Makefile, think DAG

Snakemake is a workflow manager

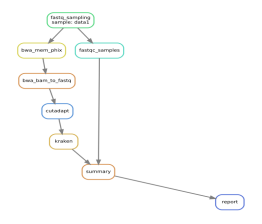


Figure:

Think Makefile, think DAG

Snakemake is a workflow manager

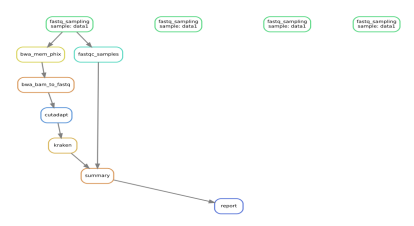


Figure: Ideal for embarassingly parallel problem

Introduction Toy example Snakemake tutorial Advanced topics Conclusions and discussions

Think Makefile, think DAG

Snakemake is a workflow manager

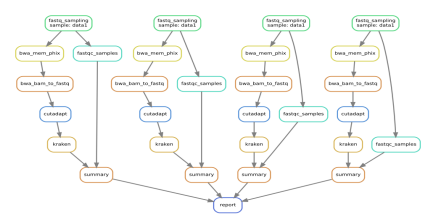


Figure: Ideal for embarassingly parallel problem

Introduction Toy example Snakemake tutorial Advanced topics Conclusions and discussions

Think Makefile, think DAG

Snakemake is a workflow manager

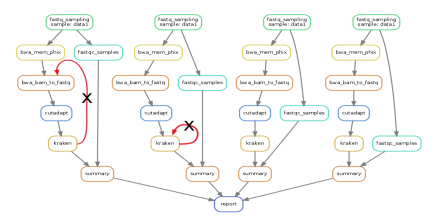


Figure: Requires a DAG (no self loop of feedback loop allowed !)

Growing community



Growing community

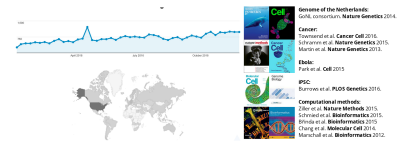


Figure: References: slides.com/johanneskoester/snakemake-tutorial-2016

Introduction Toy example Snakemake tutorial Advanced topics Conclusions and discussions

Why is it successful

Python is a batteries included language.

Snakemake overview 5 / 39

Why is it successful

Python is a batteries included language.

Snakemake as well!!

Why is it successful

Python is a batteries included language.

Snakemake as well!!

- Clusters can be used with minimum efforts (no intrusive code)
- Workflows can be run from or up to a given rule
- Data provenance
- Nice logging system to follows the status
- Suspend / Resume
- Various code can be integrated: R, bash, and of course Python

From sequential commands to

dependent rule graph:

a toy example

The problem

Let us consider two FastQ files (independent samples) and let us map them on a reference (phiX174). The two sample files are named sample_A.fastq.gz and sample_B.fastq.gz



The minimalist solution



Shell commands: pretty simple

```
# Create a directory
mkdir -p mapped_sample

# Build the index of the reference
bwa index phiX174.fa

# Do the mapping twice on the two input FastQ files
bwa mem phiX174.fa A.fastq.gz | samtools view -Sb - > A.bam
bwa mem phiX174.fa B.fastq.gz | samtools view -Sb - > B.bam
```

The minimalist solution



Shell commands: pretty simple

```
# Create a directory
mkdir -p mapped_sample

# Build the index of the reference
bwa index phiX174.fa

# Do the mapping twice on the two input FastQ files
bwa mem phiX174.fa A.fastq.gz | samtools view -Sb - > A.bam
bwa mem phiX174.fa B.fastq.gz | samtools view -Sb - > B.bam
```

Issues

Good start. Simple but what about some variables and scalability?

A shell solution



Shell loop and variables

```
#!/bin/sh
REFERENCE="phiX174.fa"
ODIR="mapped_sample"
SAMPLES='ls *.fastq.gz'

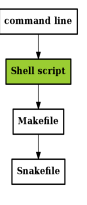
#Create a directory
mkdir -p $ODIR

# Build the index of the reference
bwa index $REFERENCE

# Do the mapping twice on the two input FastQ files
for var in $SAMPLES
do
    TARGET=${var/.fastq.gz/.bam}
    bwa mem $REFERENCE $SAMPLES | samtools view -Sb - > $ODIR/$TARGET
done
```

Introduction Toy example Snakemake tutorial Advanced topics Conclusions and discussions

A shell solution



Shell loop and variables

```
#!/bin/sh
REFERENCE="phiX174.fa"
ODIR="mapped_sample"
SAMPLES='ls *.fastq.gz'

#Create a directory
mkdir -p $ODIR

# Build the index of the reference
bwa index $REFERENCE

# Do the mapping twice on the two input FastQ files
for var in $SAMPLES
do
    TARGET=${var/.fastq.gz/.bam}
    bwa mem $REFERENCE $SAMPLES | samtools view -Sb -> $ODIR/$TARGET
done
```

Issues

Still simple but sequential. What about dependencies between tasks? What if a file exists already? Do we start from scratch?

The Makefile solution: a set of directives (rules)



A Makefile consists of a set of rules in the following form

rule syntax

target: dependencies
 system command(s)

Makefile interests:

- handles the dependencies between rules
- avoids re-rerunning a task if the targets exist already Widely used in C / C++ community for compilation of libraries.

The Makefile solution: a set of directives (rules)



Makefile: a bwa_mapping and bwa_index rule

```
SAMPLES = sample_A sample_B
ODIR = "mapped_sample"
FASTQS = $(patsubst %, %.fastq.gz, $(SAMPLES))
BAMS = $(patsubst %,$(ODIR)/%.bam,$(SAMPLES))
INDEX = phiX174.fa.bwt
REFERENCE = phiX174.fa
# Main rule
all: $(BAMS)
# bwa_mapping
$(ODIR)/%.bam: %.fastq.gz $(INDEX) $(ODIR)
  bwa mem $(REFERENCE) $< | samtools view -Sb - > $@
$(ODTR):
  mkdir -p $(ODIR)
# bwa_index
$(INDEX): $(REFERENCE)
  bwa index $<
```

The Snakefile solution



```
Snakefile
SAMPLES = ["sample_A", "sample_B"]
rule all:
    input: expand("mapped_sample/{sample}.bam", sample=SAMPLES)
rule bwa index:
    input: "phiX174.fa"
    output: "phiX174.fa.bwt"
    shell: "bwa index {input}"
rule bwa_mapping:
    input:
        ref = "phiX174.fa",
        index = "phiX174.fa.bwt",
        fastq = "{sample}.fastq.gz"
    output: "mapped_sample/{sample}.bam"
    shell:
        "bwa mem {input.ref} {input.fastq} | samtools view -Sb - > {c
```

The Snakefile solution

Snakemake logic: Makefile

Snakemake takes the best of Makefile:

- · infers dependencies and execution order
- rules define obtain output files from input files
- structured pipelines

Snakemake syntax: Python

- Own domain specific syntax (rules and keywords)
- Use Python as the glue language
- The snakemake library itself is in Python

Snakemake tutorial

The problem:

- convert a WAV signal into a frequency plot (spectrogram)
- Repeat for N input files
- Effect of the time window parameter W on the resolution
- Add an HTML / server report

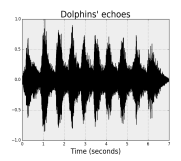


Figure: Input: a time series in WAV or OGG format

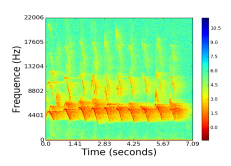


Figure: Output: a PNG image representing the time-frequency content

Rules

```
rule spectrogram:
   input: "DOLPHINS.wav"
   output: "DOLPHINS.png"
   shell: "python spectrogram.py {input}"
```

Save the file into a file called Snakefile

Execution

In a shell, type:

snakemake

Stdout

Wildcards: generalized inputs/outputs

Wildcards can be used to generalize a rule.

```
Wildcards
rule sort:
    input: "{dataset}.wav"
    output: "{dataset}.png"
    shell: "python spectrogram.py {input}"
```

Wildcards: generalized inputs/outputs

Snakemake automatically resolved multiple named wildcards. You still need to set the final targets.

```
Wildcards
rule all:
  input: ["DOLPHINS.png", "WHALES.png"]
rule sort:
    input: "{dataset}.wav"
    output: "{dataset}.png"
    shell: "python spectrogram.py {input}"
```

Wildcards: generalized inputs/outputs

If the rule's output matches a requested file, the substrings matched by the wildcards are propagated to the input files and to the variable wildcards

```
Wildcards + expands + variables
samples = ['DOLPHINS', 'WHALES']
rule all:
  input: expand("{name}.png", name=samples)
rule sort:
    input: "{dataset}.wav"
    output: "{dataset}.png"
    shell: "python spectrogram.py {input}"
```

Several wildcards

```
samples = ["DOLPHINS", "WHALES"]
windows = [512, 1024, 2048, 4096]
rule all:
    input: expand("{name}_{ws}.png",
                           name=samples,
                           ws=windows)
rule spectrogram:
    input: "{dataset}.wav"
    output: "{dataset}_{window}.png"
    shell.
      "python spec.py {input} {wildcards.window}
```

Config file

We can use a configuration file for parameters. Format are either JSON or YAML The community seems to prefer YAML.

```
config.yaml

samples: [DOLPHINS, WHALES]
windows: [512,1024,2048,4096]
```

Snakefile

Add a rule without input/output

We could add a cleanup rule:

```
rule clean:
shell: "rm -f DOL*png WHALE*png"
```

Since the rule does not produce any outputs and does not depend on other rules, it is not part of the workflow: the rule must be called explicitly:

```
snakemake clean
```

dependencies

If not target is specified, snakemake tries to apply the first rule in the workflow Do not put the clean rule at the beginning!

Handle logs

```
configfile: "config.yaml"
rule all:
 input: expand("data/{name}_{ws}.png",
                        name=config['samples'],
                        ws=config['windows'])
rule spectrogram:
 input: "data/{dataset}.wav"
 output: "data/{dataset}_{window}.png"
 log: "logs/{dataset}_{window}.log"
 shell:
   "python spec.py {input} {wildcards.window} > {log}"
```

The **param** and **run** keywords

```
rule all:
   input:
      expected_output_list,
      "server.ready"
rule server:
 output: touch ("server.ready")
 params:
    port=config['port']
 run: # Some python code
   from easydev import browse
    browse("http://127.0.0.1:%s" % params['port'])
rule spectrogram:
```

onsuccess section

If the workflow succeeds, the onsuccess section is ran if provided (same for onerror section).

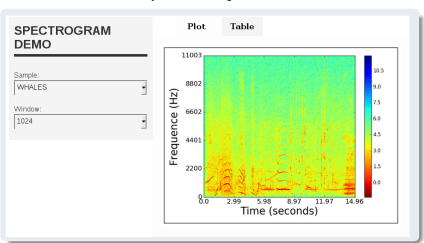
```
onsuccess:
  from myapp import SpecExample
  app = SpecExample(samples, window, "data")
  app.launch(port=config['port'])
```

You also have a **onerror** and **onstart** sections if needed.

demo

Get materials from

https://github.com/sequana/presentations/tree/master/2016/snakemake_IPday_dec/examples/demo



Advanced topics alternative title: cheeries on the cake

Job execution (source: J. Koster talk)

A job is executed if

- output file target does not exist
- output file needed by another executed job and does not exist
- input file newer than output file
- input file will be updated by other job
- execution is enforced

determined via breadth-first-search on DAG of jobs

T.Cokelaer Snakemake overview 27 / 39

Threading

Disjoint paths in the DAG of jobs can be executed in parallel using --cores argument:

```
snakemake --cores 8
```

We can be more specific inside the rules:

```
rule bwa_mapping:
   input: test.fastq
   output: test.bam
   threads: 4
   shell: bwa mem -t {threads} {input} > {output}
```

And use the same command:

```
snakemake --cores 8
```

but here only two jobs are executed at the same time

T.Cokelaer Snakemake overview 28 / 39

Resources (memory)

We can be specific about memory used by a job with the resources keyword:

```
rule bwa_mapping:
    input: test.fastq
    output: test.bam
    threads: 4
    resources: mem_mb=1000
    shell: bwa mem -t {threads} {input} > {output}
```

and use the resources parameter when calling Snakemake:

```
# execute with only 8 cores and 1Gb memory
snakemake --cores 8 --resources mem_mb=1000
```

so here only one job at a time is executed

T.Cokelaer Snakemake overview 29 / 39

Cluster execution

No intrusive code. It just worked on SGE and then on a SLURM cluster without changing a single line of code!

```
# execute the workflow on cluster with qsub submission command
# (and up to 100 parallel jobs)
snakemake --cluster qsub --jobs 100

# tell the cluster system about the used threads
snakemake --cluster "qsub -pe threaded {threads}" --jobs 100

# execute the workflow with DRMAA
snakemake --drmaa --jobs 100

# execute the workflow on cluster with sbatch (SLURM)
snakemake --cluster "sbatch --gos fast" --jobs 100
```

T.Cokelaer Snakemake overview 30 / 39

Errors

If an error occurs after hours of computation, fix the error in your code or missing files, and run snakemake again. Finished jobs won't be re-run.

T.Cokelaer Snakemake overview 31 / 39

Other features

- handles temporary and protected files
- run until a given rule
- run from a given rule
- stats about run time
- benchmark: run several times the rules
- any external scripts can be used (R, python, etc)
- remote files (http, ftp, google could, amazon, dropbox)
- rules may have priorities
- cluster time and memory can be fully customized
- modular: can include rules, or sub workflow

T.Cokelaer Snakemake overview 32 / 39

Conclusions and discussions

Conclusions

Mature

Snakemake is a mature tool ready for production.

batteries included

To cite just one great feature: free parallelization on a cluster.

Nice Syntax

The syntax is in Python, the library is in Python. Nevertheless, only a minimalist knowledge is required to get started since nice functions are already provided (e.g. expand).

Large community

Large snakemake community. See also the conda/bioconda community.

T.Cokelaer Snakemake overview 34 / 39

Discussions

Snakemake is great so what's wrong?

Not much but here are some food for thoughts

- Snakefile uses Python syntax but Snakefile are not Python module
- Errors are sometimes too cryptic and definitely not useful for end-users
- Despite lots of sanity checks, if you are not careful you may end up in an infinite loop or delete the content of a file. So do lots of testing and save your data files before production. And just avoid symbolic links same input/output filenames.
- The rule syntax is great but developpers makes different choices on how they use them. So despite a great idea of sharing tools, you end up with many different pipelines and rules that does the same thing...

T.Cokelaer Snakemake overview 35 / 39

References

- Great documentation for developers on Snakemake web site:
 - https://bitbucket.org/snakemake/snakemake/wiki/Documentation
 - https://bitbucket.org/snakemake/snakemake/wiki/Home
- Useful information:
 - http://slides.com/johanneskoester/deck-1
 - http://snakemake.bitbucket.org/snakemake-tutorial.html
 - http://slowkow.com/notes/snakemake-tutorial/
 - http://watson.nci.nih.gov/~sdavis/blog/flexible_bioinformatics_pipelines_with_snakemake/
- This talk and materials
 All snakefiles and data files to play around and available on

 $https://github.com/sequana/presentations/2016/snakemake_IPday_dec$

Snakemake Citations

Köster J., Rahmann S. Snakemake - a scalable bioinformatics workflow engine.

Bioinformatics application note Vol 28 (19) 2012

T.Cokelaer Snakemake overview 36 / 39

Acknowlegments

Snakemake overview

Acknowlegments

```
HELPERS = [
    "Dimitri Desvillechabrol",
    "Rachel Legendre",
    "Christiane Bouchier",
    "The bioinformatics and biostatistics Hub (IP)"]
rule thank:
    input: expand("{helpers}", helpers=HELPERS)
rule you:
    output: temp(touch("{to}"))
```

T.Cokelaer Snakemake overview 37 / 39

Acknowlegments

```
HELPERS = [
     "Dimitri Desvillechabrol",
    "Rachel Legendre",
     "Christiane Bouchier",
     "The bioinformatics and biostatistics Hub (IP)"1
rule thank:
     input: expand("{helpers}", helpers=HELPERS)
rule you:
    output: temp(touch("{to}"))
 snakemake -s thanks.rules --dag | dot -Tpng > thanks.png
                    you
to: Rachel Legendre
                                                          you
to: The bioinformatics and biostatistics Hub (IP)
 you
to: Christiane Bouchier
                                     you
to: Dimitri Desvillechabrol
                                  thank
```

T.Cokelaer Snakemake overview 37 / 39

Questions

What happens when the snakemake is interrupted

If you stop your snakemake (i.e. ctrl+c):

```
Terminating processes on user request.
Will exit after finishing currently running jobs.
Removing output files of failed job samtools since they might be corrupted:
reference.fa.fai
```

- On the cluster, the current job is not kill
- If you close your shell (Crash simulation):

• We can rerun the snakemake with --rerun-incomplete.

T.Cokelaer Snakemake overview 39 / 39