

Snakemake Presentation

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Workflow management

- Many methods to implement a workflow.
- They must handle:
 - Parallelization
 - Suspend/Resume

Workflow management

- Many methods to implement a workflow.
- They must handle:
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Snakemake

- Workflow management system based on python.
- Simplify the workflow's conceptions.
- Decompose workflow into rules.

Basic rule

```
rule bwa_mapping:
    input:
        ref = "genome.fa",
        fastq = "sample_A.fastq.gz"
    output:
        "mapped_sample/sample_A.bam"
    shell:
        "bwa mem {input.ref} {input.fastq} | samtools view -Sb - > {output}"
```

Basic rule

Snakefile

```
rule bwa_mapping:
    input:
        ref = "genome.fa",
        fastq = "sample_A.fastq.gz"
    output:
        "mapped_sample/sample_A.bam"
    shell:
        "bwa mem {input.ref} {input.fastq} | samtools view -Sb - > {output}"
```

Shell

```
$ snakemake
Job counts:
count jobs
1 bwa_mapping
1
rule bwa_mapping:
input: genome.fa, sample_A.fastq.gz
output: mapped_sample/sample_A.bam
1 of 1 steps (100%) done
$ snakemake
Nothing to be done.
```

Generalizing the rule

```
SAMPLE = ["sample_A", "sample_B"]

rule all:
    input:
        expand("mapped_sample/{sample}.bam", sample=SAMPLE)

rule bwa_mapping:
    input:
        ref = "genome.fa",
        fastq = "{sample}.fastq"
    output:
        "mapped_sample/{sample}.bam"
    shell:
        "bwa_mem {input.ref} {input.fastq} | samtools_view_-Sb_- > {output}"
```

Adding a rule

```
SAMPLE = ["sample_A", "sample_B"]
rule all:
   input:
        expand("sorted/{sample}.bam", sample=SAMPLE)
rule bwa_mapping:
   input:
        ref = "genome.fa",
        fastq = "{sample}.fastq"
    output:
        "mapped_sample/{sample}.bam"
    shell:
        "bwa mem {input.ref} {input.fastq} | samtools view -Sb - > {output}"
rule sort_mapping:
    input:
        bam = "mapped_sample/{sample}.bam"
    output:
        "sorted/{sample}.bam"
    shell:
        "samtools sort -0 bam {input.bam} > {output}"
```

Config file

We can add a config file (JSON/YAML) in our Snakefile.

```
config.yaml
samples:[sample_A, sample_B]
```

```
Snakefile

configfile: "config.yaml" # Ou .json

rule all:
    input:
        expand("mapped_sample/{sample}.bam", sample=config["samples"])

rule bwa_mapping:
    input:
        ref = "genome.fa",
        fastq = "{sample}.fastq"
    output:
        "mapped_sample/{sample}.bam"
    shell:
        "bwa mem {input.ref} {input.fastq} | samtools view -Sb - > {output}"
```

YAML file

config_vc.yaml

```
reference: reference.fa
sample:
    - ERR036019.bam
markduplicate: true
freebayes:
    --ploidy: 1
vcf_filter:
    QUAL: 10000
FREQ: 0.9
INFO:
    DP: ">10"
```

Optional rule

Snakefile MARKJOB = config["markduplicate"] MARKTAG = "" if MARKJOB: MARKTAG = "_undup" rule markDuplicate: input: bam = "{sample}.bam" output: a = "{sample}%s.bam" % MARKTAG, b = temp("{sample}.metrics") run: if MARKJOB: shell("./MarkDuplicates I={input.bam} O={output.a} M={output.b}") else: shell("touch {output.b}")

• NB: We can't use the ".format()" method.

Optional parameter

```
rule samtools_index:
    input:
        bam = "{sample}%s.bam" % MARKTAG
    output:
        "{sample}%s.bam.bai" % MARKTAG
    shell:
        "samtools index {input.bam}"
rule freebayes:
    input:
        bam = "{sample}%s.bam" % MARKTAG,
        bai = "{sample}%s.bam.bai" % MARKTAG,
        ref = config["reference"]
    output:
        vcf = "{sample}.vcf"
    params:
        " ".join(['%s %s' % (key, value) for (key, value) in \
            config["freebayes"].items()])
    shell:
        "freebayes {params} -f {input.ref} -b {input.bam} -v {output.vcf}"
```

Running python code

```
rule vcf_filter:
    input:
        vcf = "{sample}.vcf"
    output:
        vcf = "{sample}_filter.vcf"
run:
        from sequana import vcf_filter
        vcf_record = vcf_filter.VCF(input["vcf"])
        vcf_record.filter_vcf(config["vcf_filter"], output["vcf"])
```

Running snakemake

If all your files are in the current directory:

snakemake

• We can specify the path of Snakefile or a config file:

```
snakemake -s path/to/Snakefile --configfile path/to/configfile
```

If you want to run on 4 cores:

```
snakemake -cores 4
```

Interesting option

Dry-run is able to test if your Snakefile is valid:

```
snakemake -n
```

Change config variable in command line:

```
snakemake --config markduplicates=false
```

Doesn't work with boolean parameter because it becomes a string.

• Running only one rule, here freebayes rule:

```
snakemake --force freebayes
```

Doesn't work if the rule uses wildcards declared in other rule.

```
RuleException in line 42 of /home/desvillechabrol/Documents/pasteur/variant_calling/Snakefile:
Could not resolve wildcards in rule freebayes:
sample
```

Running snakemake on bic

It is very simple to run a workflow on bic:

```
module load snakemake
/local/gensoft2/exe/snakemake/3.5.4/bin/snakemake -p --cluster \
"gsub -g pf4 -cwd -V -b y" --jobs 5
```

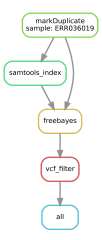
Module load must be added inside rules.

```
rule bwa_mapping:
   input:
       ref = "genome.fa",
       fastq = "sample_A.fastq.gz"
   output:
       "mapped_sample/sample_A.bam"
shell:
       """
       . /etc/profile.d/pasteur_modules.sh
       module load bwa/0.7.5a
       bwa mem {input.ref} {input.fastq} | samtools view -Sb - > {output}
       """
```

Directed Acyclic Graph (DAG)

We can generate the DAG of your workflow:

```
snakemake --dag | dot -Tsvg > dag.svg
```



References

- Snakemake website:
 - 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/
- Advanced example:
 - https://github.com/sequana/sequana/tree/master/pipelines/variants
- Enable syntax in vim for Snakemake
 - http://tinyurl.com/zm4b6c4

Additional keywords

threads Set the number of threads needed log Set a log file for a rule

```
Snakefile
rule bwa_mapping:
    input:
        ref = "genome.fa".
        fastq = "sample_A.fastq.gz"
    output:
        "mapped_sample/sample_A.bam"
    threads: 8
    log:
        "logs/bwa_mapping/sample_A.log"
    shell:
        (bwa mem -t {threads} {input.ref} {input.fastq} | \
        samtools view -Sb - > {output}) 2> {log}
        . . .
```

Cluster special features

- Snakemake generates log files with stdout and stderr of your jobs
- Your current jobs are displaying with qstat

```
# ddesvill@bic ~/Test_fastq_dir > qstat
iob-ID prior
                                         state submit/start at
aueue
                               slots ja-task-ID
3781631 0.63541 snakejob.b ddesvill
                                               03/23/2016 12:17:04
pf4@bic-n119.cluster.pasteur.f
3781632 0.63541 snakejob.b ddesvill
                                               03/23/2016 12:17:04
                                         r
pf4@bic-n119.cluster.pasteur.f
3781633 0.63541 snakejob.b ddesvill
                                               03/23/2016 12:17:04
                                         r
pf4@bic-n106.cluster.pasteur.f
3781634 0.63541 snakejob.b ddesvill
                                               03/23/2016 12:17:04
                                         r
pf4@bic-n422.cluster.pasteur.f
3781635 0.63541 snakejob.b ddesvill
                                               03/23/2016 12:17:04
                                         r
pf4@bic-n422.cluster.pasteur.f
```

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.