

# Scatter/gather-operations in Snakemake

# What does scatter/gather mean?

- **Scatter**: turn input into several pieces of output
- **Gather**: bring together (aggregate) results from the different pieces

Snakemake now has built-in support for scatter/gather processes via the `scattergather` directive. Described further in the documentation: [Defining scatter-gather processes](#). Currently not very flexible though.

# When are scatter-gather processes handy?

- demultiplexing sequencing runs
  - multiple samples per plate
  - split plates into separate files per sample
- extract reads from bam files
  - reads mapped to several genomes
  - split sequences per genome
- parallelize analyses
  - e.g. multiple sequences per sample
  - split input into smaller chunks and run analyses in parallel

etc...

Between scattering and gathering there's some type of analyses performed.

# The basics

```
DATASETS = ["a", "b", "c"]

rule scatter:
    output:
        expand('{dataset}.txt', dataset=DATASETS)
    input:
        data = 'data.tar.gz'
    shell:
        """
        tar xvf {input}
        """

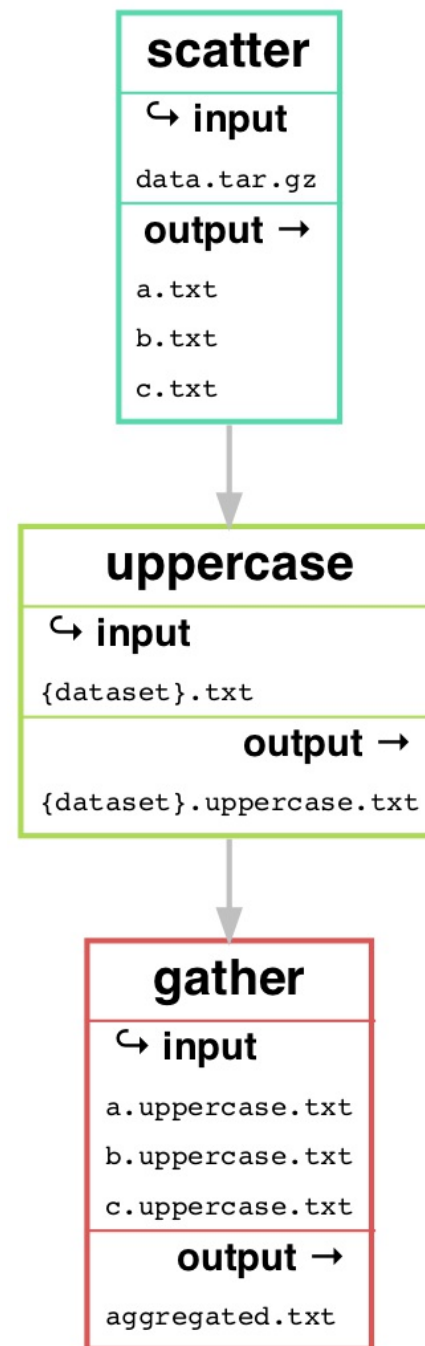
rule uppercase:
    input:
        "{dataset}.txt"
    output:
        "{dataset}.uppercase.txt"
    shell:
        """
        tr [a-z] [A-Z] < {input} > {output}
        """

rule gather:
    output:
        "aggregated.txt"
    input:
        expand("{dataset}.uppercase.txt", dataset=DATASETS)
    shell:
        """
        cat {input} > {output}
        """
```

```
snakemake -c 1
```

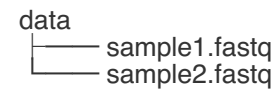
```
Job stats:
job      count  min threads  max threads
-----
gather      1         1         1
scatter     1         1         1
uppercase   3         1         1
total       5         1         1
```

# The basics

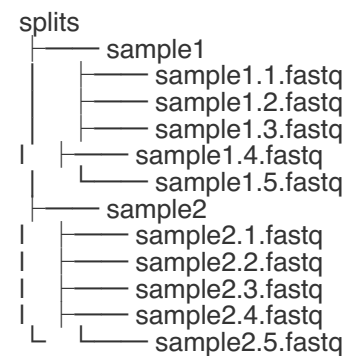


# Example: split files for parallelization

- one fastq file per sample

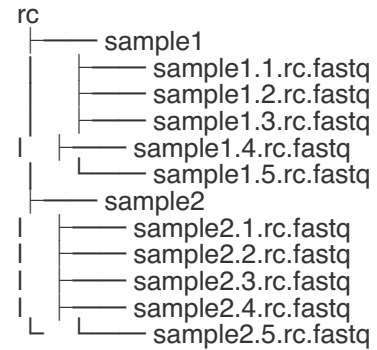


- split into several files (scatter)



# Example: split files for parallelization

- process individual files (parallelization)



- aggregate results (gather)

sample1.rc.fastq  
sample2.rc.fastq

# Example: split files for parallelization

We start with defining the number of splits

```
splits = 5  
scatteritems = range(1, splits+1]
```

Then define a rule to scatter each sample fastq

```
rule scatter:  
    output:  
        expand("splits/{{sample}}/{{sample}}.{{scatteritem}}.fastq", scatteritem = scatteritems)  
    input:  
        "data/{{sample}}.fastq"  
    log:  
        "logs/{{sample}}.scatter.log"  
    conda:  
        "envs/seqkit.yml"  
    params:  
        parts = splits,  
        outdir = lambda wildcards, output: os.path.dirname(output[0])  
    shell:  
        """  
        seqkit split -p {params.parts} -O {params.outdir} {input} > {log} 2>&1  
        rename 's/part_0*/' {params.outdir}/{wildcards.sample}.*.fastq  
        """
```

Here `scatteritem` is not a wildcard because it is expanded using the `scatteritems` list.



# Example: split files for parallelization

Next, a rule to do something with the split files per sample

```
rule reversecomplement:
  output:
    "rc/{sample}/{sample}.{scatteritem}.rc.fastq"
  input:
    "splits/{sample}/{sample}.{scatteritem}.fastq"
  conda:
    "envs/seqkit.yml"
  shell:
    """
    seqkit seq --reverse --complement {input} > {output}
    """
```

Here both `scatteritem` and `sample` are wildcards. The rule is generalized to work on any value for these wildcards.

# Example: split files for parallelization

Then a rule to gather the results per sample

```
rule gather:
  output:
    "{sample}.rc.fastq"
  input:
    expand("rc/{{sample}}/{{sample}}.{{scatteritem}}.rc.fastq", scatteritem = scatteritems)
  shell:
    "cat {input} > {output}"
```

Here `scatteritem` is not a wildcard, but `sample` is. The rule can gather split files for any sample.

# Example: split files for parallelization

Finally we put everything together, and define a pseudo rule 'all' that takes as input the gathered results for all samples.

```
samples = ["sample1", "sample2"]

splits = 5
scatteritems = range(1, splits+1)

rule all:
    input:
        expand("{sample}.rc.fastq", sample = samples)

rule scatter:
    output:
        expand("splits/{sample}/{sample}.{scatteritem}.fastq", scatteritem = scatteritems)
    input:
        "data/{sample}.fastq"

rule reversecomplement:
    output:
        "rc/{sample}/{sample}.{scatteritem}.rc.fastq"
    input:
        "splits/{sample}/{sample}.{scatteritem}.fastq"

rule gather:
    output:
        "{sample}.rc.fastq"
    input:
        expand("rc/{sample}/{sample}.{scatteritem}.rc.fastq", scatteritem = scatteritems)
```

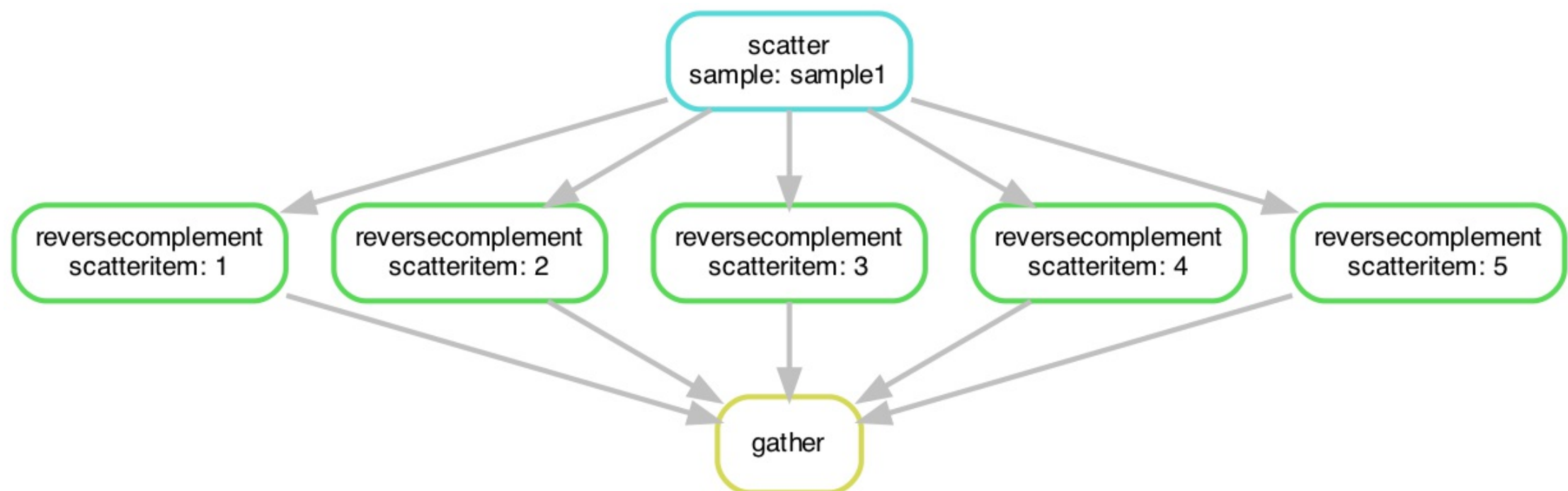
# Example: split files for parallelization

```
snakemake -c 1 --use-conda
```

Building DAG of jobs...

Job stats:

job	count	min threads	max threads
all	1	1	1
gather	2	1	1
reversecomplement	10	1	1
scatter	2	1	1
total	15	1	1



This example workflow is available at the course GitHub repository: [workshop-snakemake-byoc/tree/master/lectures/scatter-gather/](https://github.com/workshop-snakemake-byoc/tree/master/lectures/scatter-gather/)

Questions?