Nextflow: a tutorial through examples

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Introduction to Nextflow

Introduction to Nextflow

Introduction

Resources

 https://github.com/fpsom/CODATA-RDA-Advanced-Bioinformatics-2019/blob/master/ 4.Day4.md

Workflow Languages

Many scientific applications require

- Multiple data files
- Multiple applications
- Perhaps different parameters

General purpose languages not well suited

- Too low a level of abstraction
- Does not separate workflow from application
- Not reproducible

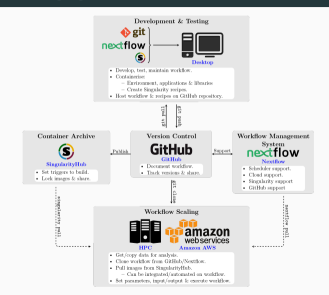
Workflow Languages

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Nextflow

Groovy-based language

- Expressing workflows
- Portable
 - works on most Unix-like systems
- Very easy to install
 - o NB: requires Java 7, 8
- Scalable
- Supports Docker/Singularity
- Supports a range of scheduling systems

Nextflow

Groovy-based language

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Key concepts of Nextflow

- · Processes:
 - actual work being done (usually simple).call program that does the analysis.
- Channels
 - o for communication between processes.
 - handles inputs and outputs.
- When all inputs ready, process is executed.
- Each process runs in its own directory (files are staged).
- Supports resumption of previous partial runs.

Introduction to Nextflow

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Nextflow Script

Simple Example: Using BASH

Input is a file

- With 6 columns
- Column 2 is an index column
- Identify rows with identical field 2
- Remove identical rows

```
11 11:189256 0 189256 A G
11 11:193788 0 193788 T C
11 11:194062 0 194062 T C
11 11:194788 0 194228 A G
11 11:193788 0 193788 A C
```

Using BASH:

```
cut -f 2 data/11.bim | sort | uniq -d > dups
grep -v -f dups data/11.bim > 11.clean
```

Simple Example: Using nextflow

Nextflow and Docker

H3AVarCall

```
#!/usr/bin/env nexflow
     input_ch = Channel.fromPath("data/11.bim")
     process getIDs {
       input:
       file input from input ch
       output:
       file "ids" into id ch
11
       file "11.bim" into orig_ch
13
       script:
14
       "cut -f 2 $input | sort > ids"
15
16
     process getDups {
18
         input:
19
         file input from id ch
20
21
         output:
         file "dups" into dups_ch
24
         script:
26
         unia -d $input > dups
27
         touch ignore
28
29
```

Introduction to Nextflow

Simple Example: Using nextflow

```
#!/usr/hin/env nexflow
     input ch = Channel.fromPath("data/11.bim")
     process getIDs {
       input:
       file input from input_ch
       output:
       file "ids" into id ch
11
       file "11.bim" into orig ch
13
       script:
14
       "cut -f 2 $input | sort > ids"
15
16
     process getDups {
18
         input:
19
         file input from id ch
20
21
         output:
         file "dups" into dups ch
24
         script:
          . . .
26
         unia -d $input > dups
27
         touch ignore
28
29
```

```
30
     process removeDups {
31
         input:
32
         file badids from dups ch
33
         file orig from orig ch
34
35
         output:
36
         file "clean.bim" into output
37
38
         script:
39
          "grep -v -f $badids $orig > clean.bim "
40
41
42
     output.subscribe { print "Done!" }
```

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37 38

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40

41

42

Simple Example: Using nextflow

```
#!/usr/hin/env nexflow
     input ch = Channel fromPath("data/11.bim")
     process getIDs {
       input:
       file input from input ch
       output:
       file "ids" into id ch
       file "11.bim" into orig ch
13
       script:
14
       "cut -f 2 $input | sort > ids"
15
16
     process getDups {
18
         input:
19
         file input from id ch
20
         output:
         file "dups" into dups ch
24
         script:
          . . .
26
         unia -d $input > dups
27
         touch ignore
28
29
```

```
process removeDups {
    input:
    file badids from dups_ch
    file orig from orig_ch

    output:
    file "clean.bim" into output

    script:
    "grep -v -f $badids $orig > clean.bim "
}

output.subscribe { print "Done!" }
```

```
$ nextflow run cleandups.nf
N F X T F I O W ~ version 19.04.1
Launching `cleandups.nf` [soggv jennings] - revision: 795e2aa39d
[warm_up] executor > local
executor > local (3)
[84/7e1ad1] process > getIDs
                               [100%] 1 of 1
[19/cc8bf9] process > getDups [100%] 1 of 1
[f9/ed086d] process > removeDups [100%] 1 of 1
Completed at: 31-Jul-2019 09:00:50
Duration
           · 1 5c
CPU hours
           : (a few seconds)
           . 3
Succeeded
```

Simple Example: Using nextflow

The work directory

```
|--work
| |--90
| | |--cebf3649d883f88381e32b4912b560
| | | |--ids -> /Users/phele/day4/work/b3/aa0380f2a1bca447259b7ffd390083/ids
| | | |--ignore
| |--9c
| | |--e0cb7d8d26682d7d4a1c44392f2bb3
| | | |--11.bim -> /Users/phele/day4/data/11.bim
| | | |--tlan.bim
| | | |--dups -> /Users/phele/day4/work/90/cebf3649d883f88381e32b4912b560/dups
| |--b3
| | |--aa0380f2a1bca447259b7ffd390083
| | | |--11.bim -> /Users/phele/day4/data/11.bim
| | | |--11.bim -> /Users/phele/day4/data/11.bim
```

Introduction to Nextflow

Partial Execution

Partial Execution

If execution of workflow is only partial

- Because of error
- Only need to resume from process that failed

nextflow run cleandups.nf -resume

Introduction to Nextflow

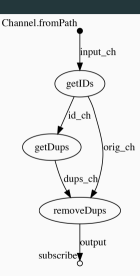
Visualising the Workflow

Visualising the Workflow

Nextflow supports several visualisation tools:

-with-dag

nextflow run cleandups.nf -with-dag <file-name>



Visualising the Workflow

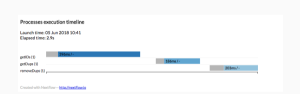
Nextflow supports several visualisation tools:

-with-dag

nextflow run cleandups.nf -with-dag <file-name>

-with-timeline

nextflow run cleandups.nf -with-timeline <file-name>

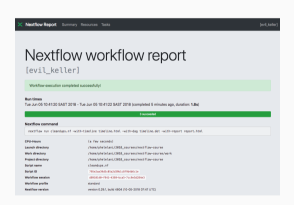


Visualising the Workflow

Nextflow supports several visualisation tools:

-with-dag nextflow run cleandups.nf -with-dag <file-name> -with-timeline nextflow run cleandups.nf -with-timeline <file-name> -with-report

nextflow run cleandups.nf -with-report <filename>



Groovy

Groovy

Nextflow is a DSL built with Groovy

- Can inter-mix Nextflow, Groovy and Java code.
- Very powerful, flexible.
- Don't need to know much (any?) Groovy but a little knowledge is a powerful thing

Groovy

Groovy Closures

Groovy: Closures

Closures are anonymous functions

- Similar to lambdas in Python
- Don't want the overhead of naming a function we only use once
- Typically use with higher-order functions
 - o Functions that take other functions as arguments
- Very powerful and useful

Syntax for a closure that takes one argument:

```
{ parm -> expression }
```

Groovy: Closures

Closures are anonymous functions

- Similar to lambdas in Python
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- Typically use with higher-order functionsFunctions that take other functions as arguments
- Very powerful and useful

Syntax for a closure that takes one argument:

```
{ parm -> expression }
```

```
\{a \to a*a \} (3)
     \{a \rightarrow a*a*7*a - 2\} (3)
     for (n in 1..5) print( {it*it} (n)):
     \{x, y \rightarrow Math.sqrt(x*x + y*y)\} (3,4)
      int doX(f, nums) {
       sum=0:
       for ( n in nums ) {
12
          sum = sum + f(n):
14
        return sum
15
16
     print doX ( {a->a}. [4.5.16] ):
     print doX ( \{a->a*a\}, [4,5,16] );
20
     print doX ( { it*it }, [4,5,16]);
22
23
     m=10
24
     print doX({a->m*a+2}, [1.2.3])
```



Extending the Example

- Parameterise the input
- Want output to go to convenient place
- Workflow takes in multiple input files processes are executed on each in turn.
- Complication : may need to carry the base name of the input to the final output;
- Can repeat some steps for different parameters.

Generalising and Extending

Parameters

Parameters

In Nextflow file:

input_ch = Channel.fromPath(params.data_dir)

And run it like this

nextflow run phylo1.nf --data_dir data/polyseqs.fa

Generalising and Extending

Channels

Data Types in Channels

Channels support different types:

- file
- val
- set

Creating Channels

```
Channel.create()
Channel.empty
Channel.from("blast","plink")
Channel.fromPath("data/*.fa")
Channel.fromFilePairs("data/{YRI,CEU,BEB}.*")
Channel.watchPath("*fa")
```

Many, many operations you can do on channels and their contents

bind	buffer	close
filter	map/reduce	group
join, merge	mix	сору
split	spread	fork
count	min/max/sum	print/view

Generalising and Extending

Generalising Our Example

Generalising and Extending

Workflow: Multiple Inputs

Nextflow and Docker

H3AVarCall

```
params.data dir = "data"
     input ch = Channel.fromPath("${params.data dir}/*.bim")
     process getIDs {
         input:
         file input from input ch
         output:
         file "${input.baseName}.ids" into id ch
         file "$input" into orig ch
         script:
13
         "cut -f 2 $input | sort > ${input.baseName}.ids"
14
15
16
     process getDups {
17
         input:
18
         file input from id ch
19
20
         output:
21
         file "${input.baseName}.dups" into dups_ch
23
         script:
24
         out = "${input.baseName}.dups"
         . . .
26
         unia -d $input > $out
27
         touch ignore
28
29
```

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Workflow: Multiple Inputs

```
params.data dir = "data"
     input ch = Channel.fromPath("${params.data dir}/*.bim")
     process getIDs {
         input:
         file input from input ch
         output:
         file "${input.baseName}.ids" into id ch
         file "$input" into orig ch
         script:
13
         "cut -f 2 $input | sort > ${input.baseName}.ids"
14
15
16
     process getDups {
         input:
18
         file input from id ch
19
20
         output:
         file "${input.baseName}.dups" into dups ch
23
         script:
24
         out = "${input.baseName}.dups"
         . . .
26
         unia -d $input > $out
27
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```

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Workflow: Multiple Inputs

```
params.data dir = "data"
     input ch = Channel.fromPath("${params.data dir}/*.bim")
     process getIDs {
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         file input from input ch
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13
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14
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         input:
18
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19
20
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23
         script:
24
         out = "${input.baseName}.dups"
         . . .
26
         unia -d $input > $out
27
         touch ignore
28
29
```

Generalising and Extending

```
process removeDups {
    publishDir "output", pattern: "${badids.baseName} clean.bim"
      → . overwrite:true. mode:'copy'
    input:
    file badids from dups ch
    file orig from orig ch
    output:
    file "${badids.baseName} clean.bim" into cleaned ch
    script:
    "grep -v -f $badids $orig > ${badids.baseName} clean.bim "
```

H3AVarCall

```
$ nextflow run cleandups.nf
Launching `cleandups.nf` [distracted hodgkin] - revision: 29

← fdh384a6

[warm_up] executor > local
executor > local (9)
[1a/431eb7] process > getIDs
                                [100%] 3 of 3
[cc/fc0aaa] process > getDups
                               [100%] 3 of 3
[03/c31154] process > removeDups [100%] 3 of 3
Completed at: 31-Jul-2019 10:26:23
Duration
           : 25
           · (a few seconds)
CPII hours
Succeeded
           . 9
```

Workflow: Multiple Parameters

Now try splitting the file but use different split values

```
split -l 400 data.txt dataX
```

will produce files dataXaa, dataXab, dataXac and so on ...

Try:

```
splits = [400,500,600]

process splitIDs {
    input:
    file bim from cleaned_ch
    each split from splits

output:
    file ("*-$split-*") into output_ch;

script:
    "split -l $split $bim ${bim.baseName}-$split-"
}
```

Generalising and Extending

Managing Grouped Files

Grouped Files

Use PLINK as an example.

```
## Short version of the command
plink --bfile /path/YRI --freq --out /tmp/YRI

## Long version of the command
plink --bed YRI.bed \
    --bim YRI.bim \
    --fam YRI.fam \
    --freq \
    --out /tmp/YRI
```

Problem:

- Pass the files on another channel(s) to be staged
- Pass the base name as value/or work it out

Pros/Cons

- Simple
- Need extra channel/some gymnastics

Grouped Files

Use **PLINK** as an example.

```
## Short version of the command
plink --bfile /path/YRI --freq --out /tmp/YRI

## Long version of the command
plink --bed YRI.bed \
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    --fam YRI.fam \
    --freq \
    --out /tmp/YRI
```

Problem:

- Pass the files on another channel(s) to be staged
- Pass the base name as value/or work it out

Pros/Cons

- Simple
- Need extra channel/some gymnastics

RECAP CLOSURES

Simply, a *closure* is an anonymous function

- Code wrapped in braces {, }
- Default argument called it

```
[1,2,3].each { print it * it }
[1,2,3].each { num -> print num * num }
```

Grouped Files - Version 1: map

```
#!/usr/bin/env nextflow
     params.dir = "data/pops/"
     dir = params.dir
     params.pops = ["YRI"."CEU"."BEB"]
     Channel
         .from(params.pops)
         .map { pop ->
             [ file("$dir/${pop}.bed").
10
               file("$dir/${pop}.bim"),
11
               file("$dir/${pop}.fam")]
13
         .set { plink data }
14
15
     plink_data.subscribe { println "$it" }
```

Grouped Files - Version 1: map

```
#!/usr/bin/env nextflow
params.dir = "data/pops/"
dir = params.dir
params.pops = ["YRI","CEU","BEB"]

Channel
    .from(params.pops)
    .map { pop ->
        [ file("$dir/${pop}.bed"),
            file("$dir/${pop}.bim"),
            file("$dir/${pop}.fam")]
}
.set { plink_data }

plink_data.subscribe { println "$it" }
```

11

13

14 15

```
[data/pops/YRI.bed, data/pops/YRI.bim, data/pops/YRI.fam]
[data/pops/CEU.bed, data/pops/CEU.bim, data/pops/CEU.fam]
[data/pops/BEB.bed, data/pops/BEB.bim, data/pops/BEB.fam]
```

Grouped Files - Version 1: map

```
#!/usr/bin/env nextflow
                                                                           16
                                                                                process getFreq {
                                                                           17
     params.dir = "data/pops/"
                                                                                  input:
     dir = params.dir
                                                                           18
                                                                                    set file(bed), file(bim), file(fam) from plink data
     params.pops = ["YRI"."CEU"."BEB"]
                                                                           19
                                                                                  output:
                                                                          20
                                                                                    file "${bed.baseName}.frg" into result
     Channel
                                                                           21
                                                                          22
                                                                                   ...
         .from(params.pops)
                                                                          23
         .man { non ->
                                                                                  nlink --bed $bed \
             [ file("$dir/${pop}.bed").
                                                                           24
                                                                                    --bim $bim \
               file("$dir/${pop}.bim"),
                                                                          25
                                                                                    --fam $fam \
11
               file("$dir/${pop}.fam")]
                                                                          26
                                                                                    --frea \
                                                                          27
                                                                                    --out ${bed.baseName}"
                                                                          28
13
         .set { plink data }
14
                                                                          29
15
     plink data.subscribe { println "$it" }
```

```
[data/pops/YRI.bed, data/pops/YRI.fam]
[data/pops/CEU.bed, data/pops/CEU.bim, data/pops/CEU.fam]
[data/pops/BEB.bed, data/pops/BEB.bim, data/pops/BEB.fam]
```

Use fromFilePairs.

• Takes a closure used to gather files together with the same key

```
x_ch = Channel.fromFilePairs( files ) { closure }
```

- Specify the files as a glob
- Closure associates each file with a key
- fromPairs puts all files with same key together
- Returns a list of pairs (key, list)

Use fromFilePairs.

 Takes a closure used to gather files together with the same key

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x_ch = Channel.fromFilePairs( files ) { closure }
```

- Specify the files as a glob
- Closure associates each file with a key
- fromPairs puts all files with same key together
- Returns a list of pairs (key, list)

A more complex example – default closure

```
Channel

fromFilePairs

("${params.dir}/*.{bed,fam,bim}",size:3, flat : true)

ifEmpty { error "No matching plink files" }

set { plink_data }

plink_data.subscribe { println "$it" }
```

Use fromFilePairs.

 Takes a closure used to gather files together with the same key

```
x_ch = Channel.fromFilePairs( files ) { closure }
```

- Specify the files as a glob
- Closure associates each file with a key
- fromPairs puts all files with same key together
- Returns a list of pairs (key, list)

A more complex example – default closure

```
Channel
Channel
fromFilePairs
("${params.dir}/*.{bed,fam,bim}",size:3, flat : true)
flow.ifEmpty { error "No matching plink files" }
flow.set { plink_data }

plink_data.subscribe { println "$it" }
```

```
[CEU, [data/pops/CEU.bed, data/pops/CEU.bim, data/pops/CEU.fam]]
[YRI, [data/pops/YRI.bed, data/pops/YRI.bim, data/pops/YRI.fam]]
[BEB, [data/pops/BEB.bed, data/pops/BEB.bim, data/pops/BEB.fam]]
```

```
process checkData {
    input:
    set pop, file(pl_files) from plink_data

    output:
    file "${pl_files[0]}.frq" into result

    script:
    base = pl_files[0].baseName
    "plink --bfile $base --freq --out ${base}"
}
```

```
process checkData {
                                                                               process checkData {
         input:
                                                                                   input:
         set pop. file(pl files) from plink data
                                                                                   set pop, file(pl_files) from plink_data
        output:
                                                                                   output:
         file "${pl files[0]}.frg" into result
                                                                                   file "${pop}.frg" into result
         script:
                                                                          8
                                                                                   script:
        base = pl_files[0].baseName
                                                                          9
                                                                                   "plink --bfile $pop --freq --out $pop"
10
         "plink --bfile $base --freq --out ${base}"
                                                                         10
11
```

Grouped Files - Final Version

```
#!/usr/hin/env nextflow
     params.dir = "data/pops/"
     dir = params.dir
     params.pops = ["YRI","CEU","BEB"]
 6
     Channel
         .fromFilePairs("${params.dir}/{YRI.BEB.CEU}.{bed.bim.fam}".size:3) {
             file -> file.baseName
10
         .filter { key, files -> key in params.pops }
12
         .set { plink data }
14
     process checkData {
15
         input:
16
         set pop. file(pl files) from plink data
17
18
         output:
19
         file "${pop}.frg" into result
20
21
         script:
22
         "plink --bfile $pop --freq --out $pop"
23
```

Generalising and Extending

On absolute paths

Absolute paths

```
input = Channel.fromPath("/data/batch1/myfile.fa")

process show {
    input:
    file data from input

    output:
    file 'see.out'

script:
    cp $data /home/scott/answer
    ...
```

Nextflow and Docker

Nextflow and Docker

Docker & Singularity Containers

Docker & Singularity Containers

Light-weight virtualisation abstraction layer

- Currently runs on Unix like systems
 - Linux
 - macOS
- Windows support coming

Can create images locally or get from repositories

```
## Docker
docker pull ubuntu
docker pull quay.io/banshee1221/h3agwas-plink
## Singularity
singularity pull docker://ubuntu
singularity pull docker://quay.io/banshee1221/h3agwas-plink
```

Running images

```
## Docker
docker run <some-image-name>
## Singularity
singularity exec <some-image-name>
```

- Docker/Singularity often run images in background
- Can also run interactively

```
## Running Docker interactively
sudo docker run -t -i quay.io/banshee1221/h3agwas-plink
## Running Singularity interactively
singularity shell docker://quay.io/banshee1221/h3agwas-plink
```

Nextflow supports Docker & Singularity

- Well designed script should be highly portable
- Each process gets run as a separate image call
 - o Under the hood, a docker run or a singularity exec is called
- Can use the same or different images for each process
 - o Parameterisable

Assuming all processes use the same image:

```
## For Docker
nextflow run plink2.nf -with-docker quay.io/banshee1221/h3agwas-plink
## For Singularity
nextflow run plink.nf -with-singularity docker://quay.io/banshee1221/h3agwas-plink
```

Nextflow and Docker

Directory & File Access

Directory & File access

Nextflow Docker/Singularity support highly transparent – but pay attention to good practice

- For each process Docker/Singularity mounts the work directory for **that** process on the Docker/Singularity image.
- Files can be staged in and out using Nextflow mechanisms.
- Other files available: directories mounted through Docker/Singularity run time options or on the Docker image
- No other files on the host machine including the current directory
- Process executes in the Docker/Singularity environment

Directory & File access

```
data = Channel.fromPath("data/pops/YRI.bim")
     process see {
         echo true
         publishDir params.publish, overwrite:true, mode:'move'
         input:
         file bim from data
         output:
         file count
         . . .
         hostname
         echo "Path is \$( pwd )\n "
         echo "Parent directory has \$( ls .. )\n"
         echo "My home directory has \$( ls /home/scott )\n"
         wc -l $bim > count
19
         ls
         ...
21
```

```
N E X T F L O W ~ version 0.21.2
Launching show_env.nf
[warm up] executor > local
[94/597f09] Submitted process > see (1)
89ad448ae0b2
Path is /home/scott/witsGWAS/dockerized/work/94/597f09ca6cc01c7be
Parent directory has 597f09ca6cc01c7be
My home directory has witsGWAS

YRI.bim
count
```

Directory & File access

Note that although the script's pwd shows:

/home/scott/witsGWAS/dockerized/work/94/597f09ca6cc01c7be

- Only these specific directories are mounted
- Only the files in the innermost directory are available

Any absolute paths (other than those used in staging) will result in error.

Profiles

In nextflow.config

```
profiles {
    ...
    docker {
    process.container = 'quay.io/banshee1221/h3agwas-plink:latest'
    docker.enabled = true
}

}
```

Now can run as:

```
nextflow run gwas.nf -profile docker
```

This can be extended in many ways

- Different processes can use different containers
- · Can mount other host directories
- Can pass arbitrary Docker parameters



Executors

A Nextflow *executor* is the mechanism which Nexflow runs the code in each of the processes

• Default is local: process is run as a script

Many others

- PBS/Torque
- SLURM
- Amazon (AWS Batch)
- SGE (Sun Grid Engine)

Selecting an executor Annotating each process

- executor directive, e.g. executor 'pbs'
- resource constraints

Or, nextflow.config file

either global or per-process

Nextflow on a cluster (HPC)

Running Nextflow on a cluster (HPC)

Script runs on the *head* node

- Nextflow uses the **executor** information to decide how the job should run
- Each process can be handled differently
- Nextflow submits each process to the job scheduler on your behalf (e.g, if using PBS/Torque, qsub is done)

Example

Scheduler + Docker

Scheduler + Docker

```
process.container = 'quay.io/banshee1221/h3agwas-plink:latest'
docker.enabled = false

process {
    executor = 'pbs'
    queue = 'batch'
    scratch = true
    cpus = 5
    memory = '2GB'
}
```

Amazon EC2

Amazon EC2

Netflow has native support for EC2

- You need an account on EC2
- Image (AMI) with the appropriate support

Launch your code:

nextflow cloud create GenomeCloud -c 5

If successful, Nextflow will give you the name of the headnode of your cluster

- ssh into into it
- run Nextflow on it.

Afterwards shut down:

nextflow shutdown GenomeCloud

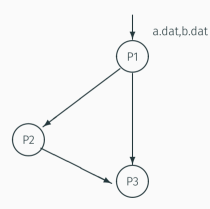
Channel Operations

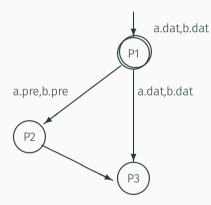
Channel operations

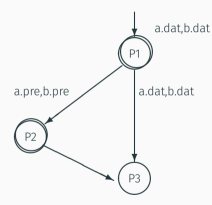
Nextflow tries to maximise concurrency

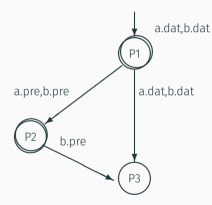
- processes are by default synchronised by channels
- when data arrives on all input channels, process executes

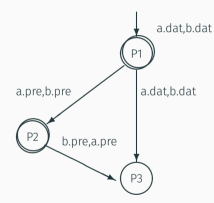
Channel operations











```
Channel.fromPath("data/*.dat").set { data }
     process P1 {
         input:
         file(data)
         output:
         file "${fbase}.pre" into channelA
         file data
                               into channelB
10
11
         script:
12
         fbase=data.baseName
13
         "echo dummy > ${fbase}.pre"
14
15
16
     process P2 {
17
         input:
18
         file pre from channelA
19
20
         output:
21
         file pre into channelC
23
         script:
24
         if (pre.baseName == "a")
25
           "sleep 4"
26
         else
27
           "sleep 1"
28
```

```
Channel.fromPath("data/*.dat").set { data }
     process P1 {
         input:
         file(data)
         output:
         file "${fbase}.pre" into channelA
         file data
                               into channelB
11
         script:
         fbase=data.baseName
13
         "echo dummy > ${fbase}.pre"
14
15
16
     process P2 {
17
         input:
18
         file pre from channelA
19
20
         output:
21
         file pre into channelC
23
         script:
24
         if (pre.baseName == "a")
25
           "sleep 4"
26
         else
27
           "sleep 1"
28
```

Try

```
29
     process P3 {
30
          echo true
31
32
          input:
33
          file(data) from channelB
34
          file(pre) from channelC
35
36
          script:
37
          . . .
38
          echo "${data} - $pre"
39
40
```

Solution: join/merge channels

- x.merge(y)
 Items emmitted by the channels x and y are combined into a new channel.
- x.join(y)
 Items emmited by the channels x and y are joined together into one channel based on existing matching key. Default: first element in each item.

Using join

Using join

```
1 ch1 = Channel.from( "a","b","c" )
2 ch2 = Channel.from( "a","d","e","a","c","b" )
3 ch1.join(ch2).subscribe { println it }
```

```
a b c
```

Using join

```
1    ch1 = Channel.from( "a","b","c" )
2    ch2 = Channel.from( "a","d","e","a","c","b" )
3    ch1.join(ch2).subscribe { println it }
```

```
a
b
c
```

Tuples:

```
1  ch1 = Channel.from( ["a",1], ["b",4], ["c",5] )
2  ch2 = Channel.from( ["a",10], ["d",8], ["e",7], ["a",9], ["c",1], ["b",10] )
3  ch1.merge(ch2).subscribe { println it }
```

```
[a, 1, 10]
[b, 4, 10]
[c, 5, 1]
```

Using merge

Using merge

```
1 ch1 = Channel.from( "a","b","c" )
2 ch2 = Channel.from( "a","d","e","a","c","b" )
3 ch1.merge(ch2).subscribe { println it }
```

```
[a, a]
[b, d]
[c, e]
```

Using merge

```
1 ch1 = Channel.from( "a","b","c" )
2 ch2 = Channel.from( "a","d","e","a","c","b" )
3 ch1.merge(ch2).subscribe { println it }
```

```
[a, a]
[b, d]
[c, e]
```

Tuples:

```
1  ch1 = Channel.from( ["a",1], ["b",4], ["c",5] )
2  ch2 = Channel.from( ["a",10], ["d",8], ["e",7], ["a",9], ["c",1], ["b",10] )
3  ch1.merge(ch2).subscribe { println it }
```

```
[a, 1, a, 10]
[b, 4, d, 8]
[c, 5, e, 7]
```

join vs merge

join vs merge

join

- If values are singletons, then the values must be the same
- If value is tuple if the, then the first element of the tuple must be the same

merge

• Merges everything into a channel, no matching.

Working version of the example

Working version of the example

```
Channel.fromPath("data/*.dat").set { data }
     process P1 {
         echo true
         input:
         file(data)
         output:
         set val(data.baseName), file("${fbase}.pre") into channelA
         set val(data.baseName), file(data) into channelB
13
         script:
14
         fbase=data.baseName
15
         "echo dummy > ${fbase}.pre"
16
     process P2 {
         echo true
19
20
         input:
         set name, file(pre) from channelA
23
         output:
24
         set name, file(pre) into channelC
```

```
25
         script:
26
         if (pre.baseName = /.*TMP.*/)
27
            "sleen 4"
28
         else
29
            "sleep 1"
30
31
32
     process P3 {
33
         echo true
34
35
         input:
36
         set name. file(data). file(pre) from channelB.join(channelC)
37
38
         script:
39
40
         echo "${data} - ${pre}"
41
42
```

Copying channels

Copying channels

You often need to copy a channel

Copying channels

You often need to copy a channel

Alternatively



H3AVarCall: Hands-on Variant Calling Practical

Prepare your workspace for the variant calling workflow!

```
## Change directory to your day4 working folder:
cd ~/Documents/day4

## Clone the H3AVarCall repository from GitHub:
git clone https://github.com/h3abionet/h3avarcall.git

## Change directory to the repository:
cd h3avarcall

## Create symbolic links to the Singularity images with applications:
ln -s /home/nfs3/h3avarcall/containers/* containers/

## Make a temorary folder in the 'scratch directory' for your 'work' folder:
mkdir -p /scratch/<USERNAME>/work
```

Lets look at some important files:

- main.nf
- main.config
- nextflow.config

DONE!! Now we are ready to start with the variant calling analysis!

Quality Checks - FastQC

H3AVarCall

Quality Checks - FastQC

Time allocated for this step: 10 minutes Run:

```
nextflow run main.nf -profile local -w /scratch/<USERNAME>/work --mode do.QC
```

Results:

H3AVarCall

Read Trimming - Trimmomatic

Read Trimming - Trimmomatic

Time allocated for this step: 10 minutes

Run:

```
nextflow run main.nf -profile local -w /scratch/<USERNAME>/work --mode do.ReadTrimming
```

Results:

```
h3avarcall
|--variant_calling_results
| |--2_Read_Trimming
| | |--workflow_report
| | | |--h3avarcall_report.html
| | | |--h3avarcall_timeline.html
| | | |--h3avarcall_timeline.ont
| | | |--h3avarcall_timeline.ont
| | | |--h3avarcall_timeline.ont
| | | |--sample_1>.1P.fastq.gz .. <sample_N>.1P.fastq.gz
| | |--<sample_1>.2P.fastq.gz .. <sample_N>.2P.fastq.gz
```

H3AVarCall

Read Alignment - BWA, GATK and Samtools

Read Alignment - BWA, GATK and Samtools

Time allocated for this step: 10 minutes

Run:

```
nextflow run main.nf -profile local -w /scratch/<USERNAME>/work --mode do.ReadAlignment
```

Results:

```
h3avarcall
|--variant_calling_results
| |--3_Read_Alignment
| | |--workflow_report
| | | |--h3avarcall_report.html
| | | |--h3avarcall_timeline.html
| | | |--h3avarcall_timeline.ont
| | | |--h3avarcall_timeline.ont
| | | |--h3avarcall_timeline.ont
| | | |--sawarcall_timeline.ont
| | | |--sawarcall_timeline.ont
| | | |--sawarcall_time.ont
| |--sawarcall
```

H3AVarCall

Variant Calling - GATK

Variant Calling - GATK

Time allocated for this step: 60 minutes

Run:

```
nextflow run main.nf -profile local -w /scratch/<USERNAME>/work --mode do.VariantCalling
```

Results:

```
h3avarcall
|--variant_calling_results
| |--4_Variant_Calling
| | |--workflow_report
| | | |--h3avarcall_report.html
| | | |--h3avarcall_timeline.html
| | | |--h3avarcall_timeline.otml
| | | |--h3avarcall_timeline.otml
| | | |--h3avarcall_timeline.otml
```

H3AVarCall

Variant Filtering - GATK

Variant Filtering - GATK

Time allocated for this step: 10 minutes

Run:

```
nextflow run main.nf -profile local -w /scratch/<USERNAME>/work --mode do.VariantFiltering
```

Results:

```
h3avarcall
|--variant_calling_results
| |--5_Variant_Filtering
| | |--workflow_report
| | | |--h3avarcall_report.html
| | | |--h3avarcall_timeline.html
| | | |--h3avarcall_timeline.otml
| | | |--h3avarcall_trace.txt
| | |--savarcall_trace.txt
| | |--senome.SNP-recal.vcf.gz
| | |--genome.SNP-recal.vcf.gz.tbi
```

H3AVarCall

Quality Checks - MultiQC

Quality Checks - MultiQC

Time allocated for this step: 10 minutes

Run:

```
nextflow run main.nf -profile local -w /scratch/<USERNAME>/work --mode do.MultiQC
```

Results:

```
h3avarcall
|--variant_calling_results
| |--MultiQC
| | |-multiqc_data
| | |-multiqc_report.html
```

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