${\bf D}$ I F F R N ${\bf A}$: A RNA-Seq analysis pipeline

Izem Mouhoubi Théo Roncalli Gustavo Magaña López

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Prerequisites

1.1 Hardware and Operating System

The pipeline was developed and tested on Ubuntu 20.04.3 LTS on top of the (GNU/Linux 5.4.0-88-generic x86_64) kernel. The output of the commands uname and neofetch are provided to further detail our configuration.

```
$ uname -a
Linux machine329a7396-059f-41aa-94c7-4c41b4ec8290 5.4.0-88-generic #99-Ubuntu SMP Thu Sep 23 17:2
$ neofetch
            .-/+oossssoo+/-.
                                          ubuntu@machine3f9ae3dc-6a4d-46b7-8131-04f00a1be146
        `:+sssssssssssssss+:`
      -+ssssssssssssssyyssss+-
                                           OS: Ubuntu 20.04.3 LTS x86_64
    . \verb"ossssssssssssssdMMMNysssso".
                                          Host: OpenStack Compute 18.2.1-1.el7
                                          Kernel: 5.4.0-88-generic
   /sssssssssshdmmNNmmyNMMMhssssss/
  +sssssssshmydMMMMMMMNddddysssssss+
                                          Uptime: 13 hours, 14 mins
 /ssssssshNMMMyhhyyyyhmNMMNhsssssss/
                                          Packages: 719 (dpkg), 4 (snap)
. \verb"ssssssdMMMNhsssssssshNMMMdsssssss."
                                          Shell: bash 5.0.17
+sssshhhyNMMNysssssssssssyNMMMysssssss+
                                          Theme: Adwaita [GTK3]
ossyNMMNyMMhsssssssssssshmmmhssssssso
                                          Icons: Adwaita [GTK3]
\verb"ossyNMMNyMMhsssssssssssshmmmhssssssso"
                                          Terminal: /dev/pts/0
+sssshhhyNMMNysssssssssssyNMMMysssssss+
                                          CPU: Intel (Haswell, no TSX, IBRS) (16) @ 2.294GHz
.sssssssdMMMNhssssssssshNMMMdsssssss.
                                           GPU: 00:02.0 Cirrus Logic GD 5446
                                          Memory: 635MiB / 64323MiB
 /ssssssshNMMMyhhyyyyhdNMMNhsssssss/
  +ssssssssdmydMMMMMMMddddysssssss+
   /sssssssssshdmNNNnmyNMMMhssssss/
    .ossssssssssssssdMMMNysssso.
      -+ssssssssssssssyyyssss+-
        `:+ssssssssssssssss+:`
```

```
.-/+oossssoo+/-.
```

This configuration was actually a virtual machine hosted on Biosphere's RAIN-Bio a cloud service maintained by the French Institue of Bioinformatics (*Institut Français de Bioinformatique*).

1.2 BioPipes: a Biosphere-commons app

The instance of the virtual machine we used is called *BioPipes*. It provides the most notable bioinformatics pipeline tools:

- nextflow
- snakemake
- cwltool

1.3 Main Tools

Their versions are specified to maximise reproducibility:

```
$ conda --version
conda 4.11.0
$ nextflow -v
nextflow version 21.10.0.5640
$ docker --version
Docker version 20.10.11, build dea9396
```

Detailed information about our development docker installation :

```
$ docker info
Client:
Context:
             default
 Debug Mode: false
 Plugins:
  app: Docker App (Docker Inc., v0.9.1-beta3)
 buildx: Build with BuildKit (Docker Inc., v0.6.3-docker)
  scan: Docker Scan (Docker Inc., v0.9.0)
Server:
 Containers: 0
 Running: 0
 Paused: 0
 Stopped: 0
 Images: 4
 Server Version: 20.10.11
 Storage Driver: overlay2
 Backing Filesystem: xfs
```

Supports d_type: true Native Overlay Diff: true userxattr: false Logging Driver: json-file Cgroup Driver: cgroupfs Cgroup Version: 1 Plugins: Volume: local Network: bridge host ipvlan macvlan null overlay Log: awslogs fluentd gcplogs gelf journald json-file local logentries splunk syslog Swarm: inactive Runtimes: io.containerd.runc.v2 io.containerd.runtime.v1.linux runc Default Runtime: runc Init Binary: docker-init containerd version: 7b11cfaabd73bb80907dd23182b9347b4245eb5d runc version: v1.0.2-0-g52b36a2 init version: de40ad0 Security Options: apparmor seccomp Profile: default Kernel Version: 5.4.0-88-generic Operating System: Ubuntu 20.04.3 LTS OSType: linux Architecture: x86_64 CPUs: 16 Total Memory: 62.82GiB Name: machine3f9ae3dc-6a4d-46b7-8131-04f00a1be146 ID: XT4Y:2HUL:HXEA:CDXV:ERC7:Z7JZ:YYRU:WZBT:ERCU:6GGA:OBZ6:QLXE Docker Root Dir: /mnt/docker-data Debug Mode: false Registry: https://index.docker.io/v1/ Labels: Experimental: false Insecure Registries: 127.0.0.0/8 Live Restore Enabled: false

Dependencies

The pipeline runs on nextflow a domain-specific language created to automate data-analysis pipelines whilst maximising reproducibility. Nextflow enables scientists to focus on their analyses, isolating different parts of the pipeline into processes whose dependencies can be dealt with using containers and virtual environments with technologies such as Docker, Singularity, and Anaconda.

The recommended way to install nextflow is via conda, using the environment file.

```
conda env create -f nextflow_conda_env.yml # will create an env called "nextflow" conda activate nextflow # You can edit the file at your choice, specially if the environment name conflicts # with a preexisting conda env on your system
```

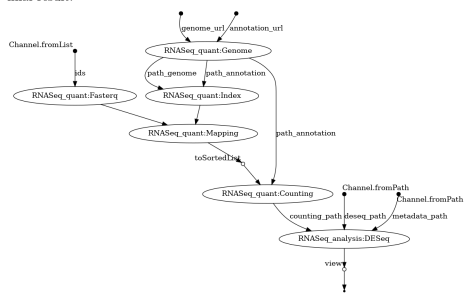
Docker should be installed as well:

```
sudo apt install docker
```

Once nextflow is installed, it will automatically retrieve the docker images used within the pipeline.

Workflow

Nextflow workflows should form a DAG (i.e. directed acyclic graph), which represents the flow of data through the different steps required to produce the final result.



This pipeline will generate a set of figures, representing differential gene expression analysis of RNA-Seq data.

Execution

1. Clone the repo to your machine

git clone https://github.com/bio-TAGI/Hackathon.git
cd Hackathon

2. Create and activate the virtual environment

conda env create -f nextflow_conda_env.yml
conda activate nextflow

3. Run the wokflow with default parameters.

```
cd Nextflow nextflow run main.nf
```

4. If you had to stop the workflow run, or if some error occurred, you can always resume the execution as follows:

```
nextflow run main.nf -resume
```

5. Specifying parameters from the command line

```
nextflow run main.nf --param1 value1\
--param2 value2\
--paramn valuen # these are generic names, not actual parameters for the pipeline
```

Parameters

- index_cpus (number of cpus reserved for the genome indexation process. default=14)
- mapping_cpus (idem. for the mapping process, used to create BAM files. default=14)
- counting_cpus (idem. for the counting process. default=7)
- mapping_memory (RAM reserved for mapping. default=50GB)

If you already possess some of the files needed to execute the pipeline, you can specify them as follows:

- reads (path pointing to a directory containing the fasterq files)
- genome (path pointing to a directory containing the genome FASTA file)
- index (Répertoire contenant les fichiers d'index)
- mapping (Répertoire contenant les fichiers BAM)
- counting (Chemin d'accès entier au fichier de comptage comprend le fichier lui-même)
- metadata (Chemin d'accès entier au fichier de métadonnées comprend le fichier lui-même)

If unspecified, the pipeline will be executed using default values from the config file: nextflow.config. These too, can be tweaked and overriden:

- ids List of SRR accession number to fetch paired-end fastq files.
 - default=['SRR628582', 'SRR628583', 'SRR628584', 'SRR628585', 'SRR628586', 'SRR628587', 'SRR628588', 'SRR628589']
- genome_url URL to download the reference genome.
 - $-\ default\ \mathtt{ftp://ftp.ensembl.org/pub/release-101/fasta/homo_sapiens/dna/Homo_sapiens.GRCh38.com/dna/Homo_sapi$
- annotation_url URL to donwload the reference genome's annotation.
 - default ftp://ftp.ensembl.org/pub/release-101/gtf/homo_sapiens/Homo_sapiens.GRCh38.101.ch
- sjdb0verhang (a STAR-specific parameter. default=99)

- For further information about this parameter, see this tutorial, or the STAR manual.

Caveats

- A good internet connection is required. Retrieving fastq can be really slow and is thus a bottleneck.
- fasterq-dump will randomly segfault. At first we thought this was caused by connection problems, but running ping ruled this out. Apparently, the segfault is a known issue.
- The workflow will inevitably fail if you try building the genome's index on a machine with less than $\sim \! \! 30$ GB of RAM available.
 - As a general rule, tweak all parameters to reasonable values that fit your setup and needs. We don't know your hardware, you do;)