ProkGenomics

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Contents

1	START HERE	5		
2	Introduction	7		
	2.1 Overview of the pipeline	7		
3	Getting Started	9		
	3.1 Make sure you have Nextflow running	9		
	3.2 Download the nextflow pipeline	9		
	3.3 Set up your working directory	10		
	3.4 How to transfer raw data to cluster/server	10		
4	Run Pipeline	13		
	4.1 Simple run	13		
5	Understand Outputs	17		
	5.1 Additional outputs description	17		
6	Write Methods 21			
7	Troubleshooting	23		
	7.1 git@github.com: Permission denied	23		
8	Resources	25		
9	Command Line Intro	27		
	9.1 General syntax and conventions:	27		
	9.2 Basic commands:	28		
	9.3 Files system	28		
10	Connect to the cluster	31		

4 CONTENTS

START HERE

I know command-line and SSH I don't know where to start

Introduction

This nextflow pipeline allows you to run several programs using a one-line command. It wraps programs to assemble, annotate, taxonomically identify, genotypic characterize and perform simple comparative genomics on prokaryotic sequencing data. This pipeline is self-contained and is NOT meant to be run in modules (not independent parts of it). This with the intention of simplifying the user interaction and the user knowledge about the bioinformatics behind this processes. However, making yourself familiar with the programs behind the scenes helps you to make informed decisions. Please go to resources to read more about what each program is doing.

2.1 Overview of the pipeline

- · Assembly, Annotation and Classification
 - Quality control of reads (short, long, or both)
 - Cleaning reads
 - De-novo Assembly (short, long or hybrid)
 - Quality Control of the assembly and detection of contamination
 - Identification of the species in the sample
 - Identification of plasmids, phages and prophages
 - Genes Annotation
- Comparative genomics (when genome reference provided)
 - Coverage of reference genome
 - Identification of core genome
 - Identification of Single Nucleotide Variants (SNV)
 - Identification of rearrangements and larger deletions
- Characterization of genes of interest (when genes sequences provided)
 - Alignments to genes of interest
 - Summary table to identy presence, absence and truncated genes

– Generation peptides sequences from the denovo assembly

Getting Started

3.1 Make sure you have Nextflow running

Before you get started with the pipeline, make sure you have installed nextflow.

If you are using a **cluster** there are high chances that nextflow is a module already installed. Please call the module. Type the following command and use **tab** after the word **nextflow** to see if you have different versions available. Ideally use the 21.10.6 version (which this pipeline has been successfully tested).

```
# if you are in a cluster
module add nextflow
```

If you can have conda installed in your system (either in your **computer** or your **server**), you could create a conda environment and install nextflow following the next commands

```
# if you can create conda env, this could work for a cluster, server or your own computer.
conda create -n nextflow
conda activate nextflow
conda install -c bioconda nextflow=21.10.6
```

Use conda activate nextflow every time you want to use nextflow

Alternatively, you could install nextflow using the nextflow installation instructions

3.2 Download the nextflow pipeline

Run the following command

when private

```
git clone git@github.com:Grinter-Lab/ProkGenomics.git
# when public
git clone https://github.com/Grinter-Lab/ProkGenomics.git
If you have issues consulte the troubleshoot section for help
A successful download of the repository should look like this:
Cloning into 'ProkGenomics'...
remote: Enumerating objects: 3, done.
remote: Counting objects: 100% (3/3), done.
remote: Compressing objects: 100% (2/2), done.
remote: Total 3 (delta 0), reused 0 (delta 0), pack-reused 0
Receiving objects: 100% (3/3), done.
To be able to run the program from any location without using the complete
path, run the following commands
# move to the program folder
cd ProkGenomics/
# print your working directory
pwd
# This will print your location something like: /path/to/dir/program/
# export the path of the folder into your PATH
export PATH="$PATH:/path/to/dir/program/"
```

3.3 Set up your working directory

Create a working directory for your project

mkdir Project1 cd Project1

Create a folder for your raw data

mkdir rawdata
cd rawdata

3.4 How to transfer raw data to cluster/server

Open a tab in your terminal from your local computer

```
# from tab in your local computer
```

scp <location in the server, notice structue as serve:path> <location in your comput

notice that the wild card allows you to move all files ending in fq.gz. if your file

scp *fq.gz <username>@<cluster_name>:/srv/home/username/folder/

Run Pipeline

Remember to put the program in your \$PATH if you haven't done it. This step has to be done every time you start a new terminal session. If you want to make this change permanent you could modify your bash profile (don't play around with it if you don't feel confident about it)

If you added the program to your \$PATH successfully you should be able to run

#remember to activate the conda env if you are using one conda activate nextflow

#run the pipeline
ProkGenomics

If it starts correctly, you will see something like

```
N E X T F L O W ~ version 21.10.6
Launching `main.nf` [focused_noether] - revision: eb930f0e69
```

If you don't see a version of that go to troubleshoot to look for possible solutions

4.1 Simple run

#remember to activate the conda env if you are using one, and if you haven't activated it yet conda activate nextflow

```
#run the pipeline
ProkGenomics --sample_name '1-77321' -profile conda
```

Parameters you can use:

Command	Description
sample_path ./rawdata/	The default path for the reads is the folder rawdata in the working directory (please follow the instructions for setting up the working folder). if you have your reads somewhere else you should set this parameter to that path.
sample_name 1-77321	The sample name is the prefix of your samples files. it doesn't have a default because I don't know your sample names. Please don't use sample names with spaces in them. Best approach is to use the name of the file as it comes from the sequencing facility
assembly_type short	This parameter can be short long or hybrid. The default is 'short'. if you have short reads you don't have to specify this parameter. If you pick the argument long or hybrid the longreads parameter should be specify. For hybrid make sure to give a path for short and long reads.
longreads	Path to the long reads.
./rawdata/longreads/	
threads 16 outdir 1-77321	Number of threats to use. More threats faster your processing. Make sure you know what is available for you. The results will be in a folder in the working directory with the same sample
reference	name and _results ex. 1-77321_results. If you have a reference genome put the
	tapath here. This will activate all the comparative genomics steps. This file can be formatted as fasta or genbank. If you provide a genbank file your Single
adapter_file TruSeq3-PE.fa	Nucleotide Variant file will be annotated (tell you what gene has the mutations). To trim your short reads you need to specify what adaptors where used when sequencing. Arguments are TruSeq2-SE.fa, TruSeq2-PE.fa, TruSeq3-PE.fa. The default is
genes_interest GenesBD/*fasta	TruSeq3-PE.fa. Path to a folder that contains all genes of interest. The correct formatting is a gene per file in fasta format.

15

Command	Description
assembly genome_assembly.fasta	If you already have an assembly you can set this parameter and the pipeline will skip all the steps of assembly

Understand Outputs

This pipeline produce a folder per program run and several main outputs. The main results are described in the final report named <sample_name>_ProkGenomics_report.html (click hyperlink to see)

5.1 Additional outputs description

```
Folder structure
```

```
sample_name_results
  fastqc
      sample_prefix_1_fastqc.html
      sample_prefix_1_fastqc.zip
      sample_prefix_2_fastqc.html
      sample_prefix_2_fastqc.zip
      sample_prefix.R1.trim_fastqc.html
      sample_prefix.R1.trim_fastqc.zip
      sample_prefix.R1.unpaired.trim_fastqc.html
      sample_prefix.R1.unpaired.trim_fastqc.zip
      sample_prefix.R2.trim_fastqc.html
      sample_prefix.R2.trim_fastqc.zip
      sample_prefix.R2.unpaired.trim_fastqc.html
      sample_prefix.R2.unpaired.trim_fastqc.zip
      software_details.txt
  trimmomatic
      sample_prefix.R1.trim.fastq
      sample_prefix.R1.unpaired.trim.fastq
      sample_prefix.R2.trim.fastq
      sample_prefix.R2.unpaired.trim.fastq
      software_details.txt
```

```
unicycler
    sample_prefix
       001_spades_graph_k027.gfa
       001_spades_graph_k053.gfa
       001_spades_graph_k071.gfa
       001_spades_graph_k087.gfa
       001_spades_graph_k099.gfa
       001_spades_graph_k111.gfa
       001_spades_graph_k119.gfa
       001_spades_graph_k127.gfa
       002 depth filter.gfa
       003_overlaps_removed.gfa
       004_bridges_applied.gfa
       005_final_clean.gfa
       assembly.fasta <<<< Complete de novo assembly
       assembly.gfa
       unicycler.log
    software_details.txt
checkm
   sample_prefix.tsv <<<< QC de novo assembly</pre>
   software_details.txt
checkv
   sample_prefix
       complete_genomes.tsv
       completeness.tsv
       contamination.tsv
       proviruses.fna
       quality_summary.tsv <<<< Phages or Provirus detected
       viruses.fna
   software_details.txt
prokka
   sample_prefix_annotation_output
       sample_prefix.err
       sample_prefix.faa
       sample_prefix.ffn
       sample_prefix.fna
       sample_prefix.fsa
       sample_prefix.gbk <<<< Gene Annotation</pre>
       sample_prefix.gff <<<< Gene Annotation</pre>
       sample_prefix.log
       sample_prefix.sqn
       sample_prefix.tbl
       sample_prefix.tsv
       sample_prefix.txt
   software details.txt
pharokka
```

```
sample_prefix_annotation_output
   sample_prefix.err
   sample_prefix.faa
   sample_prefix.ffn
   sample_prefix.fsa
   sample_prefix.gbk
   sample_prefix.gff <<<< Phages or Provirus annotation
   sample_prefix.log
   sample_prefix.sqn
   sample_prefix.tsu
   sample_prefix.tst</pre>
```

Write Methods

Workflow of pipeline

Troubleshooting

7.1 git@github.com: Permission denied

You may see this error:

Cloning into 'ProkGenomics'...
git@github.com: Permission denied (publickey).
fatal: Could not read from remote repository.

Please make sure you have the correct access rights and the repository exists.

Note that this is a private repository, you may required to log in using your github details. Github now requires for you to setup a token key to access private repositories, please follow the github instructions to set up one

Resources

```
https://www.bioinformatics.babraham.ac.uk/projects/fastqc/http://www.usadellab.org/cms/?page=trimmomatichttps://github.com/rrwick/Unicyclerhttps://singularity-tutorial.github.io/01-installation/Other programshttps://bactopia.github.io/v3.0.0/https://proksee.ca/https://genome.usegalaxy.org.au/
```

Command Line Intro

If you are completely new to working with command line the following short introduction should be useful to get you started. Please read carefully this section, it will help you to understand instructions in later sections.

9.1 General syntax and conventions:

• Code or command are instructions directly given to the computer through a console or terminal window. Code or command lines in this tutorial are written with this style or

in this boxes

• If a string is written between < > it means that you have to type what that means in your case. For example: Login as: <your username> this means you have to type your user name in that space without the < >

For example, the following instructions should look like:

```
cp <file_name> <file_destination>
```

My file name os myfile.txt and my file destination is newfolder

cp myfile.txt newfolder

- When a sterisk * is used it means all of that kind. For example: 1s *.fasta will print a list of all files that have the extension .fasta
- Every line starting with # is a comment. There lines are not interpreted by your computer, there are there only to give you additional information.

9.1.1 Programs:

Command lines for executing programs usually looks like:

program --input <inputfile>
where
program is the program in question
--input is the option or parameter
inputfile is the argument

- Options/parameters for a program are denotated by a dash and a letter as:-f or a double dash and a string as: --file. If an option is not required but optional is often explained using [], for example: [-t 8]
- Arguments are the input to the options/parameters. For example -f myfile.txt. -f is the option to input your file and myfile.txt is the argument for that option, the name of your file. The arguments are often explained using < >. When several arguments are possible for an option pipes are used to show the different possibilities, for example [-f sam|bam]. This means the option -f allows sam or bam formats

9.2 Basic commands:

When you enter your terminal your prompt consists of: HOST_NAME:MACHINE CURRENT_DIRECTORY \$ everything after \$ is your command line. You can use the following basic commands to access information or perform tasks in your computer.

• change directory

cd <name of directory you want to change to>
cd or cd ~ move you to your home directory

• print working directory

pwd

• list your files

ls

• make dir*ectory

mkdir <new folder name>

• copy (needs file to be copied and destination).

cp <path of file to be copy> <destination path>

9.3 Files system

• Please note that directories are structured in a hierarchical system. You have to know where you are standing to ask the computer to move to the correct

```
folder.

Example of folder structure:

| subfolder_1 |
|main_folder |
| subfolder_2 |
| subfolder_2.1 (YOU ARE HERE)

#where am I?

pwd

#shows this path: /main_folder/subfolder_2/subfolder_2.1

# I want to go to the folder conteining this folder

cd ..

# moves to /main_folder/subfolder_2/

# I want to go to the folder conteining this folder and change to a folder that is there

cd ../subfolder_1

#moves to subfolder_1
```

Connect to the cluster

10.0.1 Macs

If you are working on Mac you can directly open the terminal from applications or click the Launchpad icon in the Dock, type Terminal in the search field, then click Terminal. You will see a version of this:

type the following command

ssh <username>@<cluster_name>

where <username> is your authcate and the <cluster_name> is the cluster you are connecting to. Click enter, you will be asked for a password. Enter your password and click enter. Note you will not see the characters as they are typed. You are now in your home directory on the cluster.

10.0.2 Windows

If you are on a windows-based PC, you will need to download PuTTY.

In the hostname (or IP address) box, enter the hostname that you were provided, ie. <username>@<cluster_name>, where <username> where <username> is your authcate and the <cluster_name> is the cluster you are connecting to. Ensure the connection type is SSH. Click open. You will be prompted to enter your username (authcate) and password in the terminal window. Enter your credentials and click enter. Note you will not see the characters as they are typed. You are now in your home directory on the cluster.

Now you are ready to go Let's get started