AusARG Phylogenomics Workflow

Keogh Lab: March, 2022

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Introduction

This documents outlines the process of going from raw reads provided from the BioPlatforms Australia data portal through to generating gene trees. There are many ways to do each individual step in this workflow, and probably many ways it could be improved, so take notes on things that don't work as you go. Most of this is adapted from the SqCL Pipeline developed by Sonal Singhal, who has been a **huge** help throughout this process.

The working components of this workflow rely on a series of bioinformatics and phylogenetics software packages that have been *containerized* using Singularity. This means the packages are free standing (including dependencies), so as long as you have $Singularity \ v.3+$ installed on your machine you can run them without any prior installation. The available packages are included in the $SqCL_Pipeline$ directory that this tutorial is a part of. Have a look:

```
/SqCL_Pipeline

|-- bbmap_38.90--he522d1c_3.sif

|-- bcftools_1.12--h3f113a9_0.sif

|-- ...

|-- samtools_1.3.1--h1b8c3c0_8.sif

|-- Scripts

| |-- align_reads1.py

| |-- align_reads2.py

| |-- ...

|-- trimmomatic_0.39--hdfd78af_2.sif

|-- ...
```

Parts of this workflow will take place on your local machine (downloading raw data, sample file construction), but most of it will be executed on a separate computer/server. If you're at ANU this is most likely the machine \mathbf{nick} in Craig Moritz's group (nick.rsb.anu.edu.au). To use this server you'll need to make sure you have a login to access remotely via a terminal and are comfortable moving files back and forth with scp. All of the scripts you need to process your data are included in the directory $SqCL_Pipeline/Scripts$, or we will generate from some basic R functions in the file $AusARG_Tools$. Because you likely have a lot of samples to get through, we'll speed up the computing process by handling things in parallel using GNU Parallel to take advantage of the available resources.

Data Pre-Processing

Let's jump in to get our data sorted and ready for processing. Let's start by opening up a new R file and sourcing the functions in $AusARG_Tools.R$

```
source("Path_to/SqCL_Pipeline/AusArg_tools.R")
```

We want to generate a sample info file that holds the basic info about each sample so we can refer to that later. The sample info file will look roughly like this:

```
## Warning in read.table(file = file, header = header, sep = sep, quote = quote, :
## incomplete final line found by readTableHeader on 'Scripts/sample samples.csv'
##
         sample
## 1 Pnig 350789
## 2 Pgra_350790
## 3 Dpax_350791
##
                                                               read1
## 1 /home/ian/SqCL_Pipeline/Optimization/Pnig_350789_R1_concat.fastq.gz
## 2 /home/ian/SqCL_Pipeline/Optimization/Pgra_350790_R1_concat.fastq.gz
## 3 /home/ian/SqCL_Pipeline/Optimization/Dpax_350791_R1_concat.fastq.gz
##
## 1 /home/ian/SqCL_Pipeline/Optimization/Pnig_350789_R2_concat.fastq.gz
## 2 /home/ian/SqCL_Pipeline/Optimization/Pgra_350790_R2_concat.fastq.gz
## 3 /home/ian/SqCL_Pipeline/Optimization/Dpax_350791_R2_concat.fastq.gz
##
                                                    adaptor1
## 1 GATCGGAAGAGCACACGTCTGAACTCCAGTCAC*ATCTCGTATGCCGTCTTCTGCTTG
## 2 GATCGGAAGACCACGTCTGAACTCCAGTCAC*ATCTCGTATGCCGTCTTCTGCTTG
## 3 GATCGGAAGACCACGTCTGAACTCCAGTCAC*ATCTCGTATGCCGTCTTCTGCTTG
##
                                                                    barcode1
                                                         adaptor2
## 1 AATGATACGCCGACCACCGAGATCTACAC*ACACTCTTTCCCTACACGACGCTCTTCCGATCT
                                                                  TACGCCAAGT
## 2 AATGATACGGCGACCACCGAGATCTACAC*ACACTCTTTCCCTACACGACGCTCTTCCGATCT
                                                                  TCCATGACGG
barcode2
                         lineage
## 1 TTCGGCTCCG Pygopus nigriceps
## 2 AACGTCTCCG Pletholax_gracilis
## 3 CATAAACGTC
                       Delma_pax
```

The file can have more columns, you can add additional information, we just won't use it in any of our scripts. But, it absolutely must ultimately have information for: sample, read1, read2, adaptor1, adaptor2, barcode1, barcode2, and lineage.

Step 0: concatenate collate

What we need as input is our library metadata file (the one you submitted with your samples for sequencing), in a .csv format and a directory where we're storing all our fastq.gz read files. Once we know those locations we can run the function $concatenate_collate$. What this does is (1) identifies all the raw read files, (2) extracts and combines information from the file names and the metadata file, (3) concatenates all forward (R1) and reverse (R2) read files for each sample separately.

```
adaptor1 = "AGTCAC*ATCTC",
adaptor2 = "CTACAC*ACACT")
```

- sample.dir: full path to the folder holding your 'fastq.gz' read files
- metadata.file: file name for the csv metadata sequencing file (assumes it's in the 'sample.dir')
- outfile: name the output file
- out.path: this is the output directory path on the computer you will analyze the data on (not your local machine)
- adaptor1 & adaptor2: sequence info for the adaptors, with barcodes replaced by '*'. The default value for this function will provide the appropriate adaptor sequences since they are the same for all 1536 UDIs we purchased from Perkin Elmer (adaptor+barcodes).

After executing this function, you should have successfully created a sample info file that we will use for the rest of the pipeline. One thing to consider though is how you have the *lineages* column designated. If you have multiple samples of a given species, and you want to assemble them separately, make sure to provide unique *lineage* names for each, e.g. "Homo_sapiens_1", "Homo_sapiens_2". If instead you want to assemble samples separately but collate them into a single representative of the species, you would label both the same, e.g. "Homo_sapiens".

Troubleshooting: This function was designed using data from our optimization experiment—and so like the rest of these scripts—has not been tested extensively on different data types. One way I can conceive of this function breaking is around concatenating the forward (R1) and reverse (R2) read files. The NextSeq500 we ran our initial sequencing on has four physical lanes per flowcell. When the BRF makes fastq files from the Illumina BCL files, it uses bcl2fastq which uses a lane-splitting argument that outputs four forward (L001_R1-L004_R1) and four reverse (L001_R2-L004-R2) read files, instead of the normal two (one R1, one R2). If future sequencing efforts result in a different number of output files per sample, adjust lines 59–68 in the $AusARG_Tools.R$ file.

Read Assembly

Almost all of the assembly and alignment scripts in this workflow can be run individually or across multiple samples in parallel. If running individually, you can check each script's commands with the -help flag.

```
$ python Scripts/dedupe_clean_filter_reads.py --help
```

The -help flag will also provide information on commands for the parallel instances too.

```
$ python Scripts/generate_shell_scripts.py --help
```

Instructions for both instances are shown below, follow along by color (individual | in parallel). To take advantage of the available computer resources, we can also generate shell scripts and run these in parallel using the single python script generate_shell_scripts.py. All this does is write the full python command for each sample in your sample info file to a file, then read it in and simultaneously process multiple commands using GNU Parallel. Most people will want to run this pipeline for many samples simultaneously.

Below, there are explanations of each script and their associated commands, as well as Suggested Usage to optimize speed and sample throughput, and Output to tell you what files/directories to expect.

Step 1: dedupe_clean_filter_reads

The <code>dedupe_clean_filter_reads.py</code> script first removes identical duplicate sequences from your raw reads using <code>BBMap</code> and <code>dedupe.sh</code>. It quickly reformats the deduplicated reads back into two read files, then removes lingering adaptor and barcode sequences using <code>trimmomatic</code> and pairs up the raw reads using <code>pear</code>. Finally, it maps the reads against a reference file of phylogenetically diverse target sequences using <code>BBMap</code> and <code>bbmap.sh</code>, and removes any reads which do not match any targets with at least a user specified minimum identity threshold (<code>-minid</code>).

Suggested Usage: this step can be quite memory intensive, but you should be able to run a number of samples in parallel. Consider *-mem 10 -CPU 4 -parallel 10*. At maximum usage this will use 100Gb memory and 40 CPU.

Output: a /trim reads directory with * adapters.fa, duplicates.fq.gz, and dd.fastq.gz* files.

Run individually:

```
~/SqCL_Pipeline$ python Scripts/dedupe_clean_filter_reads.py \
---dir [...] --file [...] --mem [...] --CPU [...] \
--sample [...] --minid [...] --ref [...]
```

Run in parallel:

- script: the python script you'd like to run, here—dedupe_clean_filter
- dir: the directory path (within SqCL_Pipeline) that holds the fastq.gz files and the sample info file
- file: the sample info file
- parallel: how many instances should run simultaneously? Remember: total memory usage will be $-parallel \times -mem$, and total CPU usage will be $-parallel \times -CPU$
- project: the project name
- mem: memory use per instance
- *CPU*: memory use per instance
- ref: path to the reference fasta file of phylogenetically diverse target sequences
- minid: minimum identity score for bbmap.sh
- analyze: if not included this will just output the shell script, if -analyze yes will automatically run the shell script

Step 2: trinity_filtered_assembly

The *trinity_filtered_assembly.py* script takes as input our deduplicated/cleaned/filtered reads and attempts to assemble them into contigs.

Suggested Usage: different steps in the Trinity assembler have different requirements. The first step (clustering k-mers) is highly memory intensive. The final phase (butterfly) requires high CPU capacity. Best bet is to run just a couple samples in parallel but with sufficient memory and core resources (e.g. -mem 40 - CPU 40 - parallel 2).

Output: *_bb.final.fq.gz, _final.fz.gz, and _unpaired.final.fq.gz* files.

Run individually:

```
~/SqCL_Pipeline$ python Scripts/trinity_filtered_assembly.py \
--dir [...] --file [...] --mem [...] --CPU [...] --sample [...]
```

Run in parallel:

- script: the python script you'd like to run, here—trinity filtered assembly
- dir: the directory path (within SqCL Pipeline) that holds the fastq.gz files and the sample info file
- file: the sample info file
- parallel: how many instances should run simultaneously? Remember: total memory usage will be $-parallel \times -mem$, and total CPU usage will be $-parallel \times -CPU$
- project: the project name
- mem: memory use per instance
- *CPU*: memory use per instance
- analyze: if not included this will just output the shell script, if -analyze yes will automatically run the shell script

Step 3: match_contigs_to_probes

The $match_contigs_to_probes.py$ script uses BLAST to match assembled contigs to our target sequences. Based on the user provided e-value (how likely the contig is to actually be the target), we can categorize each contig/target as:

- easy_recip_match: 1-to-1 unique match between contig and targeted locus
- \bullet complicated_recip_match: 1-to-1 non-unique match, in which one targeted locus matches to multiple contigs
- ditched_no_recip_match: a case in which the contig matches to the targeted locus, but it isn't the best match
- ditched_no_match: a case in which the contig matches to the targeted locus, but the locus doesn't match to the contig
- ditched_too_many_matches: a case in which one contig has multiple good matches to multiple targeted loci

From this, we can further choose which matches we want to keep and which to discard, but that will be in the next step.

Suggested Usage: these steps can be processed quickly for a lot of samples, consider running many samples in parallel.

Output: /matches directory with a blat_results subdirectory and a * matches.csv* file per sample.

Run individually:

```
~/SqCL_Pipeline$ python Scripts/match_contigs_to_probes.py \
--dir [...] --sample [...] --evalue [...] --db [...]
```

Run in parallel:

- script: the python script you'd like to run, here— $match_contigs$
- dir: the directory path (within SqCL Pipeline) that holds the fastq.gz files and the sample info file
- *file*: the sample info file
- parallel: how many instances should run simultaneously? Remember: total memory usage will be $-parallel \times -mem$, and total CPU usage will be $-parallel \times -CPU$
- project: the project name
- evalue: strictness of the contig-to-target match. More info see: What is the Expect value?
- db: the fasta file which holds our target sequences (imagine it as a target database). For most purposes this is Scripts/squamate_AHE_UCE_genes_unique.flipped.fasta
- analyze: if not included this will just output the shell script, if -analyze yes will automatically run the shell script

Step 4: make_PRG

The <code>make_PRG.py</code> script allows us to choose which contig-to-probe matches we would like to keep (one of two options <code>easy_recip_match</code>, or <code>complicated_recip_match</code>), and then generates a Pseudo-Reference Genome for each sample or lineage. If you'd like it to generate each sample individually make sure the <code>lineage</code> field in your sample info file is unique for each sample. This will result in all sampled individuals being represented in the final alignments. If you'd like to generate a PRG for each "lineage" or "species", then make the <code>lineage</code> field reflect that accordingly. This will result in only one individual per "lineage" or "species" being represented in the final alignments.

Output: a /PRG directory holding a .fasta file per sample.

Run individually:

```
~/SqCL_Pipeline$ python Scripts/make_PRG.py \
--dir [...] --file [...] --lineage [...] --keep [...]
```

Run in parallel:

Requirements:

- script: the python script you'd like to run, here—make PRG
- dir: the directory path (within SqCL Pipeline) that holds the fastq.gz files and the sample info file
- *file*: the sample info file
- parallel: how many instances should run simultaneously? Remember: total memory usage will be $-parallel \times -mem$, and total CPU usage will be $-parallel \times -CPU$
- project: the project name
- keep: which matches to keep? Probably stick with easy_recip_match
- analyze: if not included this will just output the shell script, if -analyze yes will automatically run the shell script

Step 5: quality_2_assembly

The quality_2_assembly.py script is a quality control step that aims to give us information about the number of targets recovered per sample or lineage/species. There are two instances of it, so make sure to choose the appropriate one when identifying the -script command.

Output: an output directory named with the -outdir flag, which holds an .assemblyquality.csv file per sample.

Run individually:

```
~/SqCL_Pipeline$ python Scripts/make_PRG.py \
--dir [...] --file [...] --ind [...]
```

Run in parallel:

```
~/SqCL_Pipeline$ python Scripts/generate_shell_scripts.py \
--script quality_2_sample \
```

```
--dir [...] \
--file [...] \
--project [...] \
--outdir [...] \
--analyze yes
```

- script: the python script you'd like to run, here—quality_2_sample or quality_2_lineage. Not sure the lineage implementation works right now.
- dir: the directory path (within SqCL_Pipeline) that holds the fastq.gz files and the sample info file
- file: the sample info file
- project: the project name
- outdir: where should we save the quality score files?
- analyze: if not included this will just output the shell script, if -analyze yes will automatically run the shell script

Alignment and Gene Trees

Step 6: phylogeny_make_alignments

The phylogeny_make_alignments script will run across all available samples and pull contigs into target alignments. We don't need to parallelize this, so don't worry about it. One thing to consider is the -minsamp flag, which determines what is the minimum number of samples required to build an alignment. Most shortcut coalescent methods like ASTRAL require quartets to determine the bipartitions, so having -minsamp < 4 is not useful.

Output: a /phylogeny directory which holds a locus_data.csv summary file and a /alignments subdirectory holding all the rough locus alignment files.

Run individually (same as in parallel):

```
~/SqCL_Pipeline$ python Scripts/phylogeny_make_alignments.py \
--dir [...] --file [...] --minsamp [...]
```

Requirements:

- dir: the directory path (within SqCL_Pipeline) that holds the fastq.gz files and the sample info file
- file: the sample info file
- minsamp: minimum sample number required to build an alignment (default is 4)

Step 7: phylogeny_align_genetrees

The phylogeny_align_genetrees_parallel script has only a parallel implementation. It begins by finding the rough alignments, correcting the direction of contigs using MAFFT to make sure they're all consistent. It then aligns the sequences using MAFFT, and optionally trims the edges and cleans up the alignment with Gblocks. Once the alignments are clean, it builds gene trees for each alignment (if specified with -tree_method). If your computing resources are low, you can always run the alignment and gene tree building iteratively by setting -CPU 1, but expect this to proceed slowly.

Run in parallel:

```
~/SqCL_Pipeline$ python Scripts/phylogeny_align_genetrees_parallel.py \
--dir [...] --trim [...] --CPU 80 --tree_method iqtree
```

- dir: the directory path (within SqCL_Pipeline) that holds the fastq.gz files and the sample info file
- trim: no commands required, but if this flag is included it will automatically clean and trim the reads. There are a number of additional parameters you can identify (-b1, -b2, -b3, -b4), but I suggest you keep them at the default settings as this will be a very light clean/trim.
- CPU: number of available cores, doubles as the number of parallel instances to run

Crumbs

Some additional resources which might be useful.

Moving Files

It can be a nuisance to move files between your local computer and another machine. Use *scp* to copy files using your terminal.

```
local machine$ scp [path on local] [path on server]
```

For example, if we wanted to move the file *geckos.fasta* from our local machine to our home directory on the server **nick**:

```
local_machine$ scp path_to/geckos.fasta ian@nick.rsb.anu.edu.au:/home/ian
```

Alternatively when we want to move an output file (updated_geckos.fasta) from the server back to our local machine:

machine:
local_machine\$ scp ian@nick.rsb.anu.edu.au:/home/ian/path_to/updated_geckos.fasta [local_path_destinati

Sometimes we need to move a folder or a series of files with similar extensions.

To move an entire directory:

```
local_machine$ scp -r [path_to_directory] [path_to_destination]
```

To move a series of fasta files:

```
local_machine$ scp path_to/*.fasta [path_to_destination]
```

Checking Usage Stats

Before running any analyses it's important to make sure there are sufficient available computing resources (e.g. someone else isn't hogging the server).

Quickly check availability with htop.

server\$ htop

This will show the number of available CPUs, amount of memory being used, who is using it, et al. If we're only interested in the process that our account is running:

```
server$ htop -u [username]
```

We can also quickly check available disk space with free.

```
server$ free -h
```

This will show the amount of hard disk space installed, available, and in use.

Downloading Containers

The included containerized packages have the file extension .sif (Singularity Image File) and have been downloaded from *quay.io using Singularity. E.g.

```
$ singularity pull docker://quay.io/biocontainers/mafft:7.480--h779adbc_0
```

There are tons of other available packages/programs, so have a browse.

Mounting Drives

Standard user accounts on nick write to disk /Data which does not have enough room for storing the raw read files and the outputs. To get around this we'll mount our account to a larger drive. Do this **before** starting any of the data processing. This will require superuser privileges, so seek out either Ian or Emily to help.

```
# from in your home user directory (/home/[username]/)
server$ mkdir SqCL_Pipeline
# now bind your SqCL_Pipeline to the SqCL directory on /Data2
server$ sudo mount --bind /Data2/SqCL /home/[username]/SqCL_Pipeline
```