**Instruction to apply rabies-tailored ARTIC network bioinformatic pipeline**

Adapted from <https://artic.network/ncov-2019/ncov2019-bioinformatics-sop.html>

Last updated: 20th Feb 2023

**Remember**: these instructions give the generic structure for commands but require editing to make them specific to your data, they are not a straight copy and paste! I have tried to indicate what you will need to edit.

**Make a new directory for analysis**

This instruction organises output files into an ‘analysis’ directory in your home directory, within which each run should have its own subdirectory. If you wish to change the location of the directory, modify the path accordingly (**/path/**analysis/run\_name). The commands below will only create these directories if they do not exist already i.e. will not overwrite an existing output directory for the named run.

run\_name should be replaced with the name of your sequencing run.

mkdir -p analysis/run\_name

cd analysis/run\_name

**Activate the ARTIC environment:**

All steps should be performed in the artic-rabv conda environment:

source activate artic-rabv

**Basecalling with Guppy**

If you did basecalling with MinKNOW, you can skip this step and go to Demultiplexing.

Run Guppy basecaller on the MinION output folder from your sequencing run:

For fast mode basecalling:

guppy\_basecaller -c dna\_r9.4.1\_450bps\_fast.cfg -i /path/to/reads -s run\_name -x auto -r

You need to substitute /path/to/reads to the folder where the FAST5 files from your

run are. The default locations for MinKNOW output folders are:

*Mac*: /Library/MinKNOW/data/run\_name

*Linux*: /var/lib/MinKNOW/data/run\_name

*Windows* c:/data/reads

This will create a folder called run\_name with the basecalled reads (fastq files) in it.

**Demultiplexing**

For the current version of the ARTIC protocol it is essential to demultiplex using strict

parameters to ensure barcodes are present at each end of the fragment (the --require\_barcodes\_both\_ends option)

guppy\_barcoder --require\_barcodes\_both\_ends -i run\_name -s output\_directory --barcode\_kits "EXP-NBD104"

Modify argument –-barcode\_kits according to kit used for run e.g.

- if you used native barcodes 1-12 and 13-24 it should be "EXP-NBD104 EXP-NBD114"

- 96 native barcode kit would be “EXP-NBD196”

Note: the output files from this command (or from MinKNOW) will be labelled according to their barcode id i.e. NB01, or barcode01.

The following steps must be done for every barcode

**Read filtering**

This step is performed for each barcode in the run.

We collect all the fastq files (typically stored in files each containing 4000 reads), filter out reads below the expected amplicon length (less than 350) and concatenate them into a single file.

To collect and filter the reads for barcode03, we would run:

artic guppyplex --skip-quality-check --min-length 350 --directory output\_directory/barcode03 --prefix run\_name

You will now have a files called: run\_name\_barcode03.fastq

**Run the MinION pipeline: medaka**

For each barcode you wish to process (e.g. run this command 12 times for 12 barcodes),

replacing the file name and sample name as appropriate:

artic minion --medaka --medaka-model r941\_min\_fast\_g303 --normalise 200 --threads 4 --scheme-directory ~/Github/artic-rabv\_ea/primer\_schemes --read-file <xx.fastq> <rabv\_ea/V1> <samplename>

Replace the read-file, samplename and primer scheme (the bit after fastq) as appropriate.

E.g. To process a sample sequenced with barcode02 (and therefore output from Minknow labelled as barcode02.fastq), prepared using the east Africa primer scheme (rabv\_ea) but a sample name of exampleRun1\_GBR\_RV921…

artic minion --medaka --medaka-model r941\_min\_fast\_g303 --normalise 200 --threads 4 --scheme-directory ~/Github/artic-rabv\_ea/primer\_schemes --read-file barcode02.fastq rabv\_ea/V1 exampleRun1\_GBR\_RV921

Relevant primer schemes:

rabv\_ea/V1 – East Africa primers

rabvSEasia/V1 – Philippines primers

rabvPeru2/V1 – Peru primers

Double check the filepath to the artic-rabv folder. If this does not match the location on *your* laptop, modify as necessary.

**Output files**

samplename.rg.primertrimmed.bam - BAM file for visualisation after primer-binding site

trimming

samplename.trimmed.bam - BAM file with the primers left on (used in variant calling)

samplename.merged.vcf - all detected variants in VCF format

samplename.pass.vcf - detected variants in VCF format passing quality filter

samplename.fail.vcf - detected variants in VCF format failing quality filter

samplename.primers.vcf - detected variants falling in primer-binding regions

samplename.consensus.fasta - consensus sequence

**Gather consensus files together into one fasta file**

To put all the consensus sequences in one file called my\_consensus\_genomes.fasta , run

cat \*.consensus.fasta > <my\_consensus\_genomes>.fasta

Choose an informative name for the output fasta!

**Sequence alignment**

MAFFT can be used for alignment (it should already be installed on your laptop). To align sequences in your fasta file, run

mafft <my\_consensus\_genomes>.fasta > <my\_consensus\_genomes>.aln.fasta

This runs mafft with default parameters. Try and distinguish between unaligned and aligned fasta files in the filename e.g. I always add a .aln to the output file to annotate it as an alignment.

**To visualise genomes in Tablet**

Open a new Terminal window and type:

tablet

Go to “Open Assembly”

Load the BAM (binary alignment file) as the first file.

Load the reference file (e.g. in artic/artic-rabv/primer\_schemes/rabv\_ea/V1/rabv\_ea\_reference.fasta or whatever is the relevant one for your primer scheme) as the second file.

Select Variants mode in Color Schemes for ease of viewing variants.