# **ONTbarcoder**

A pipeline for MinION based DNA barcoding

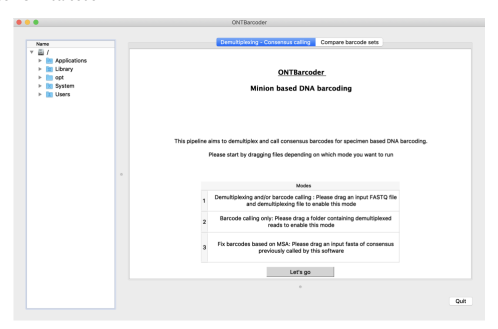
**ONTBarcoder** (available from <a href="https://github.com/asrivathsan/ONTbarcoder">https://github.com/asrivathsan/ONTbarcoder</a>) is installed by unzipping the folder with the version of the program that supports the operating system on your computer. The folders are available from here:

#### ONTBarcoder's three modules.

- 1. <u>Demultiplexing</u>: MinION reads are assigned to specimen-specific bins.
- 2. <u>Barcode calling</u>: The reads in the specimen-specific bins are used to derive the barcodes based on alignment and consensus calling.
- 3. <u>Barcode comparison module</u>: Two or more sets of barcodes can be compared.

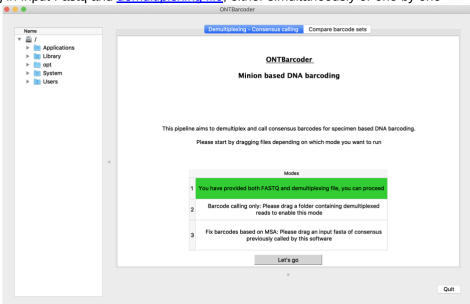
We recommend that you try the pipeline using our DatasetA containing 257 amplicons generated from Flongle. It is available from <a href="here">here</a>. Information to download the various datasets is available <a href="here">here</a>. Prior to running any dataset, ensure you have at least as much space as is required for the input FASTQ file.

1. Open ONTbarcoder



## Demultiplexing and barcode calling using the "Demultiplexing-Consensus calling tab"

2. Drag in Input Fastq and demultiplexing file, either simultaneously or one by one



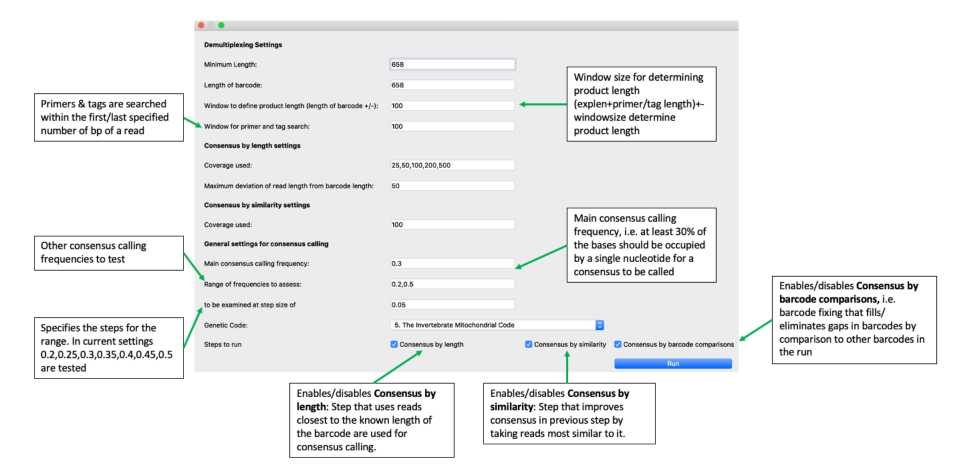
3. Press "Let's go"

4. Configure your run based on the settings described below. DatasetA does not require any changes to the default. The default assumes that you are working on invertebrate barcodes that are 658 bp in length. The remaining settings define which reads are demultiplexed, where the primers are found, how many reads are used in the different steps, and how the consensus barcodes are determined.

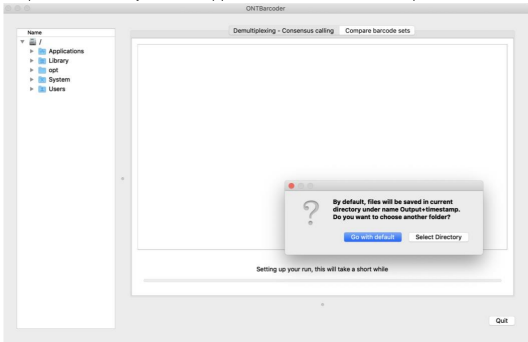
### **BASIC SETTINGS**

#### **Analysis workflow** Expected barcode length, used Reads longer than this are for QC and determining expected accepted for demultiplexing product length 1. Demultiplexing based on primers and tags **Demultiplexing Settings** Minimum Length: 658 In iterative mode, consensus Number of reads used for are called at the lowest Length of barcode 2a. Consensus by Length: consensus by length. coverage. Those barcodes Window to define product length (length of barcode +/-): subset to reads closest to If multiple coverages are not passing the QC criteria of expected barcode length to specified translation, meeting barcode 100 Window for primer and tag search: length and being ambiguity specified coverages and call (delimiter=comma), then Consensus by length settings free are passed for barcode iterative mode is used. consensus using MAFFT calling at higher coverage. 25,50,100,200,500 Maximum deviation of read length from barcode length Number of reads used for Consensus by similarity settings 2b. Consensus by consensus by similarity. 100 Similarity: subset to reads Coverage used most similar to consensus General settings for consensus calling in 2a (similarity >90%) and Genetic code 0.3 Main consensus calling frequency: call consensus using MAFFT 0.2.0.5 Range of frequencies to assess to be examined at step size of 0.05 Genetic Code 5. The Invertebrate Mitochondrial Code 3. Consensus by barcode ✓ Consensus by similarity ✓ Consensus by barcode comparisons comparisons: subset to Steps to run Consensus by length reads most similar to consensus in 2a (similarity >90%) and call consensus using MAFFT Start the run

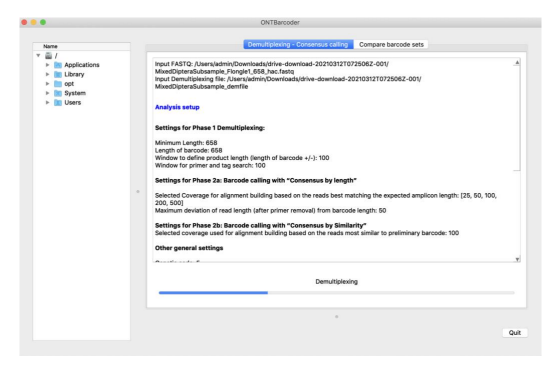
# **ADVANCED SETTINGS**



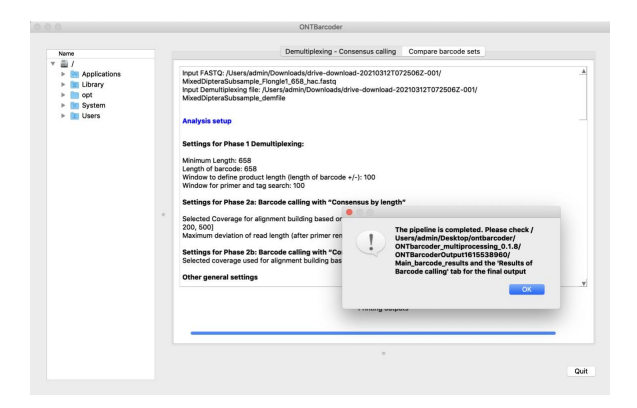
5. Select an output folder. If you select "default", it will create a folder within the directory containing the software with a unique name starting with ONTbarcoderOutput. If you choose "Select Directory" ensure that the directory name has no empty space and is empty. Generally avoid complex characters in your names (spaces, brackets, slashes, colons)



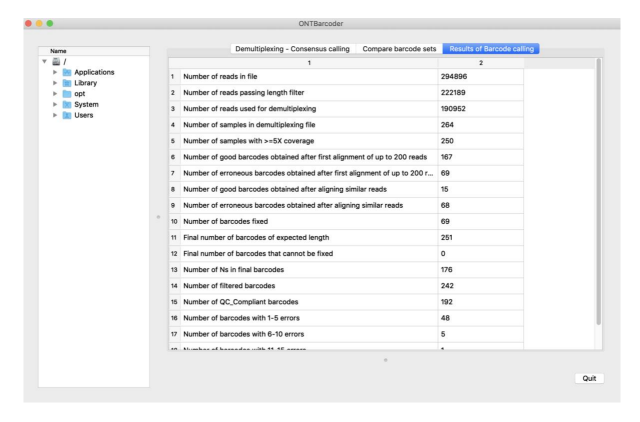
6. The analysis starts, a progress bar appears, and results are generated in respective folders as the analysis advances.



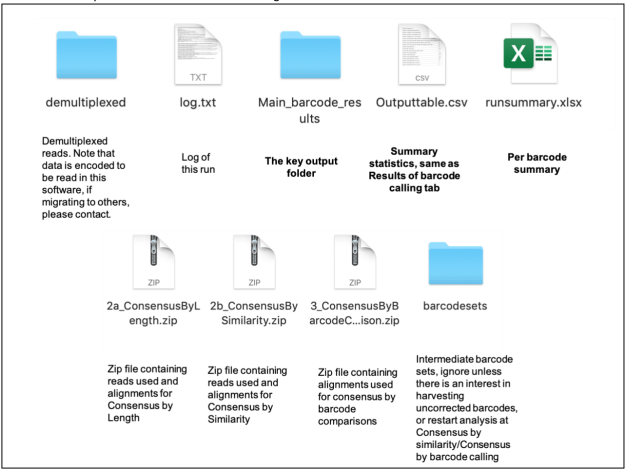
7. The analysis is complete and all output files have been created.



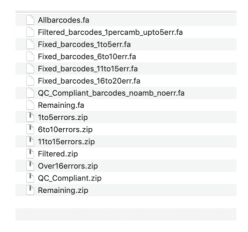
8. Results Table: The table reports that 242 "Filtered" barcodes were obtained of which 192 are QC-Compliant. This means they have no ambiguities, are translatable, have expected barcode length and are not having any gaps in an internal Multiple Sequence Alignment (MSA) check. The filtered sets include addition 50 barcodes that have <1% ambiguous bases, are translatable, have expected length, and may have been corrected for up to 5 indels. The table can be copied but it also stored in the "Outputtable.csv" file in the output folder.



9. The output folder contains the following



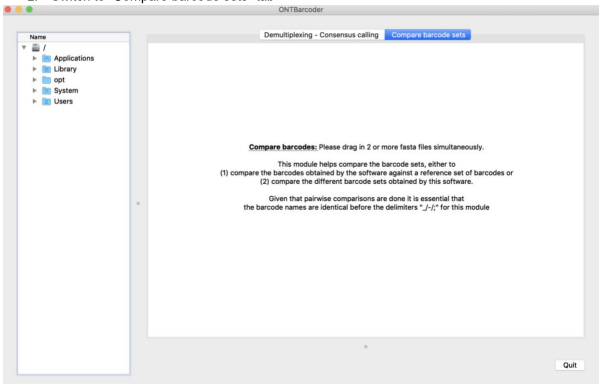
10. The Main Barcode Results folder contains the main barcode output file "Filtered\_barcodes" fasta file. The barcodes without ambiguous bases are in the QC\_Compliant\_barcodes file. The zip folders contain demultiplexed reads per dataset divided into each of the categories. All barcodes should be checked via BLAST for contamination and/or verification that the barcode obtained belongs to the expected taxon.



### 2. Compare barcodes in the "Compare barcodes sets tab"

This can be used in different modes (4 modes: single barcode file against single reference, multiple barcode files against single reference, pairwise comparisons of two barcode files, and all-vs-all comparisons of multiple barcode files), which depend on how many input files are dragged in. Here we show how to compare the output for Dataset A with reference sequences generated with Sanger sequencing.

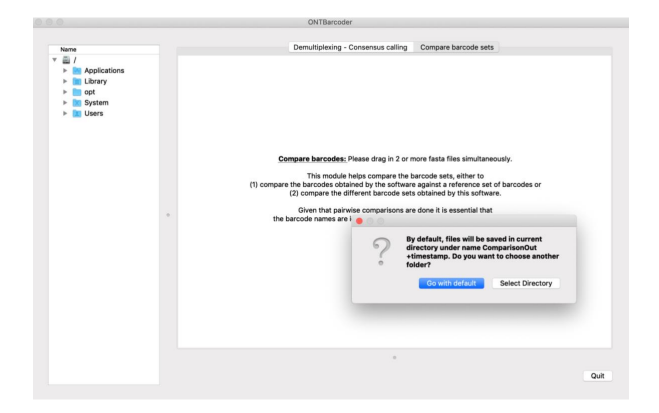
1. Switch to "Compare barcode sets" tab



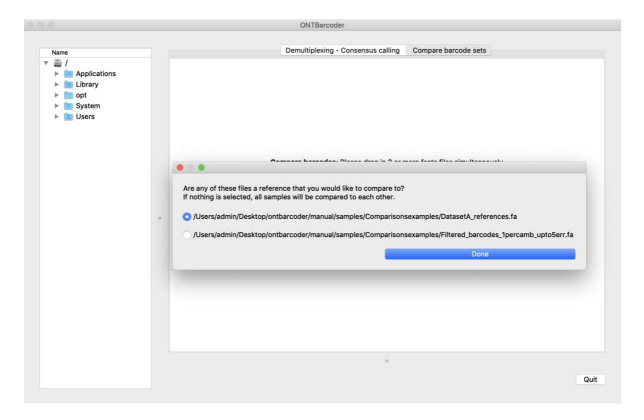
Drag at least two fasta files into the window. For the example, go to the "Main\_barcode\_results" folder and use the following two files:

DatasetA_references.fa	
Filtered_barcodes_1percamb_upto5err.fa	

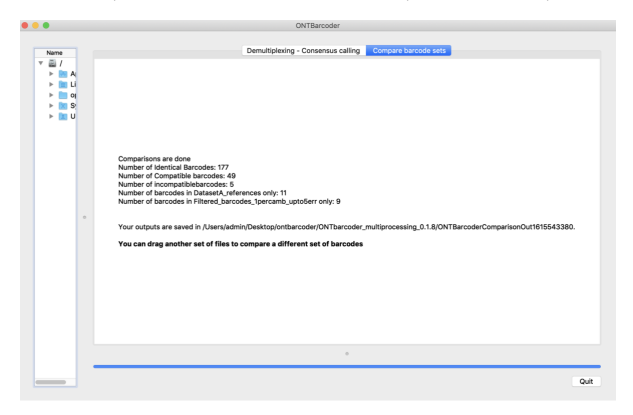
Select an output mode, if you select default, it will create a folder within the directory containing
the software starting with ONTBarcoderComparisonOut. If you choose "Select Directory" ensure
that the directory name has no empty space and is empty. Generally avoid complex characters in
your names (space, brackets, slashes, colons)



4. You can now select a set of reference barcodes if you want all other files to be compared to the barcodes in the reference set. Alternatively, you can just press "Done" to do all pairwise comparisons between the fasta files. Here, since we want the reference mode, we select the reference Sanger file and click "Done"



5. The output will be as shown below and the text can be copied. It is also in the output folder



The barcodes in the sets are compared and classified into three categories: "identical" where sequences are a 100% match and lack ambiguities, "compatible" where the sequences only differ by ambiguities, and "incompatible" where the sequences differ by at least one base pair.

6. The output folder contains the following:



- "statsfile.csv" is the key summary folder having summary of the comparisons and the various incompatible barcodes with the edit distances
- Identicalbarcodes\* and compatible\* barcode files gives one sequence per dataset, in case the user wants to dereplicate barcodes
- The \*onlybarcodes.fa files contain barcodes that cant be compared
- "badbarcodes" folder contains incompatible barcodes and the value before "\_" represents edit distance.



Note that if more than 2 input files were provided, each pairwise comparison is given in a separate folder and a summary of all comparisons is then given the outermost folder

# Summary

#### **Expected input and output**

Demultiplexing module	Input required	Results
Demultiplexing	<ul> <li>FASTQ file obtained after base-calling</li> <li>Demultiplexing file</li> </ul>	<ul> <li>Demultiplexed reads in "demultiplexed" folder</li> <li>Overall summary in "Outputtable.csv"</li> <li>Per barcode summary in "runsummary.xlsx</li> </ul>
2. Barcode calling module	Input required	Results
Barcodes are derived from reads in specimen-specific bins	Folder containing     Demultiplexed data	<ul> <li>Barcode sets in "Main barcode results" folder</li> <li>Overall summary in "Outputtable.csv"</li> <li>Per barcode summary in "runsummary.xlsx"</li> </ul>
Improving barcodes using "Consensus by similarity" and "Consensus by barcode comparisons"	Sequences in "barcodesets" folder	<ul> <li>Barcode sets available from         "Main barcode results" folder</li> <li>Overall summary in "Outputtable.csv"</li> <li>Per barcode summary in         "runsummary.xlsx"</li> </ul>
3. Barcode comparison module	Input required	Results
Comparison of barcodes to references	<ul> <li>One or more barcode fasta file(s)</li> <li>Reference fasta file See format specifications</li> </ul>	"statsfile.csv" that describes the overall summary and erroneous barcodes
Comparison of barcodes to each other	Two or more barcode fasta file(s)     See format specifications	"statsfile.csv" that describes the overall summary and erroneous barcodes

# **Format Specifications**

File			
FASTQ file	Standard fastq, generated after basecalling with ONT software		
Demultiplexing File	A 5-column csv file with the following headers: SpecimenID, TagFsequence, TagRsequence, PrimerF, PrimerR		
	You can only demultiplex one one primer pair at a time. FASTQ files with data for multiple pairs, have to processed sequentially.		
	Please avoid unusual characters in Specimen ID (e.g. characters like "(){}[]V.,;*\$" will lead to crashes)		
Files for comparison module	Specimen barcodes in different files should have identical name before the following delimiter characters ("_/-/;")		
	2. The barcodes for at least some of the specimens should be present in multiple input files		
Input file for improvement	ONTbarcoder's pipeline can be started at different points.		
(Step 4)	Demultiplexing files can be used to only carry out barcode calling.		
	Consensus by barcode can be started with files that have the format specified in the hyperlink		