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MultiQuas (Multiple reference quasispecies reconstruction protocol) V.4

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In Development



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ABSTRACT

The following protocol summarizes the major steps to run the MultiQuas pipeline to evaluate viral variability and reconstruct the viral quasispecies from NGS data (particularly Miseq reads). It is based on the assumption that 1 o more known references are available. These references could be obtained using other haplotype reconstruction softwares. Nonetheless, it is recommended that only a few trusted references are used.

PROTOCOL CITATION

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52873

MATERIALS TEXT

A



FastQC 0.11.9 © by Simon Andrews

```
Align_to_references.sh
#!/bin/bash
start=`date +%s`
bbduk.sh in1=$2 out1=reads 1.fq in2=$3 out2=reads 2.fq ref=
[path/to/bbmap/instalation]/bbmap/resources/adapters.fa ktrim=r k=23 mink=11
hdist=1 tpe tbo qtrim=rl trimq=20 minlen=50 maq=20
bowtie2-build $1 VFAref
bowtie2 --no-discordant --no-mixed -p $4 -x VFAref -1 reads_1.fq -2 reads_2.fq |
samtools view -@ 4 -bT $1 - > SAMPLE.bam
samtools sort -@ 4 -m 2G SAMPLE.bam > SAMPLE_sorted.bam
samtools view -@ 4 -h -F 4 -b SAMPLE_sorted.bam > SAMPLE_map.bam
samtools index -@ 4 SAMPLE map.bam SAMPLE map.bai
samtools depth -d10000000 SAMPLE_map.bam > coverage.txt
lofreq viterbi -f $1 -o SAMPLE_map_viterbi.bam SAMPLE_map.bam
samtools sort -@ 4 -m 2G SAMPLE_map_viterbi.bam >
SAMPLE_map_viterbi_sorted.bam
samtools index -@ 4 SAMPLE_map_viterbi_sorted.bam
SAMPLE_map_viterbi_sorted.bai
lofreq indelqual --dindel -f $1 -o SAMPLE map viterbi sorted indels.bam
SAMPLE map viterbi sorted.bam
samtools index -@ 4 SAMPLE_map_viterbi_sorted_indels.bam
SAMPLE_map_viterbi_sorted_indels.bai
lofreq call-parallel --pp-threads $4 --call-indels --use-orphan -f $1
SAMPLE_map_viterbi_sorted_indels.bam -o variants.vcf
end=`date +%s`
echo Execution time was 'expr $end - $start' seconds.
```



samtools 1.12 ⇔ source

bcftools 1.12 © source

Bowtie2 2.4.4 © source

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Brief pipeline description

Reads are trimmed and filtered using

bbduk © source by Brian Bushnell

2 Filtered and trimmed reads are aligned to a set of user-defined references (multifasta format) with

Bowtie2 2.4.4 © source

3 Reads are then split into different classes (one for each reference and one for the unmapped reads) using

SAMtools 1.8 ©
Linux
source by Wellcome Trust Sanger Institute

4 For each class, reads are merged using

PEAR - Paired-End reAd mergeR

source by Alexandros Stamatakis

and then haplotypes are reconstructed with

QuRe © source by Mattia C. F. Prosperi

Important note: If the time limit is reached this step is repeated with a subset of the reads in order to reduce computation time and resources required by QuRe. Steps include 0.99, 0.9, 0.8, 0.7, 0.6 and 0.5 proportions of the toal reads for each class in that order.

- 5 Next, the proportions of each haplotype class (predicted by QuRe) are adjusted to reflect the number of reads of the corresponding class.
- 6 All reconstructed haplotypes are aligned to the first reference in the multifasta file using

mafft 7.487 © source by Kazutaka Katoh

7 Additionally, reads are then aligned to the first reference in the multifasta file. Single Nucleotide Variants (SNVs) are called using

Lofreq 2 © source by Andreas Wilm

8 Finally, concordance between the SNVs (expected minor allele frequency) from the predicted quasispecies and the Lofreq variants using an in-house R script. Higher value of R-squared indicates a better quasispecies reconstruction. Important: this step does not meant as a validation of the obtained results, but it allows the user to choose between different haplotypes reconstructions.

Installing docker

- 9 In order to run the pipeline, a wrapper file is available (bash) which automatically perfoms all the above numbered steps. A docker image is available that includes all the necessary dependencies. If you do not yet have docker installed, do so at this time, and ensure that is in your PATH. For more information please visit https://www.docker.com/get-started
- 10 The docker image ("multiquas") is available at Docker hub To pull the image, use the command below:

docker pull

docker pull cacciabue/multiquas:latest

This will download and install the corresponding docker image. Only has to be run the first time (it may take several minutes depending on your internet connection)

- 10.1 Alternatively, If you don't want to use Docker, you can install all dependencies by yourself (only for linux users). The dependencies are:
 - BCFtools v1.8 (or later version) http://www.htslib.org/download/
 - Samtools v1.8 (or later version) http://www.htslib.org/download/
 - Bowtie2 v2.2.4 http://bowtie-bio.sourceforge.net/bowtie2/index.shtml
 - PEAR https://cme.h-its.org/exelixis/web/software/pear/doc.html
 - seqtk <u>https://github.com/lh3/seqtk</u>
 - bbmap https://jgi.doe.gov/data-and-tools/bbtools/bb-tools-user-guide/bbduk-guide/
 - Lofreq v2 https://csb5.github.io/lofreq/
 - mafft https://mafft.cbrc.jp/alignment/software/
 - R v4.1 (or later version) https://www.r-project.org/
 - R package seqinr https://cran.r-project.org/web/packages/seqinr/index.html
 - R package ape https://cran.r-project.org/web/packages/ape/
 - R package VariantAnnotation
 - https://bioconductor.org/packages/release/bioc/html/VariantAnnotation.html
 - R package Biostrings https://bioconductor.org/packages/release/bioc/html/Biostrings.html
 - R package ggplot2 https://ggplot2.tidyverse.org/
 - R package ggrepel https://cran.r-project.org/web/packages/ggrepel/vignettes/ggrepel.html and contact the author for the most up-to-date wrapper scripts.

Preparing analysis

11 Depending on your operating system follow these steps

On windows: open a windows terminal (WIN + R), type "cmd" and press enter. A Windows terminal

- 11.1 should be up and running.
- 11.2 Create a folder to work with and *navigate* to the location for your new *folder*. For example:

mkdir test_dir
cd test_dir
linux

- 11.3 Copy the fastq files and the reference file (1 or more sequences to use as references in multifasta format) to the test_dir folder.
- 12 On linux the steps are similar.
 - 12.1 Open a terminal (ctrl + alt + T), A bash terminal should be up and running.
 - 12.2 Create a folder to work with and navigate to the location for your new folder. For example:

mkdir test_dir cd test_dir linux

12.3 Copy the fastq files and the reference file (1 or more sequences to use as references in multifasta format) to the test_dir folder.

Running MultiQuas workflow

13 The following command will create a docker container, mount the test_dir folder into the container and perform all the necessary step of the MultiQuas workflow.



Backups files of the fastq reads and reference should be made and store in a different location before running the following command in order to prevent loss of data.

13.1 For Windows:

run reconstruction

docker run -it --volume %cd%:/nexus cacciabue/multiquas:latest reconstruction.sh complete -o OUTPUT_FOLDER -1 R1.fq -2 R2.fq -r REFERENCE.fasta

windows

The user should replace each of the following:

- OUTPUT_FOLDER: folder name inside test_dir to automatically save all files.
- R1.fq and R2.fq: corresponding names of the paired-end reads (use the correct extension in case you have .fastq)
- REFERENCE.fasta: name of the fasta file to use.

Optional paramenters can be set:

- l: use to indicate a specific label (default: SAMPLE)
- -p/--proc: Number of threads to use (default=2)
- -m/--mem: ram memmory available to use by the java machine (default=4).
- --min_quality: Phred value cutoff for filtering the reads (default=25)
- --timeout: seconds (per reference) before shutting down QuRe in case it freezes (default=600). Each time, the reads are downsampled and the step is repeated (6 times per reference)

13.2 For linux:

run reconstruction

docker run -it --volume \$(pwd):/nexus cacciabue/multiquas:latest reconstruction.sh complete -o OUTPUT_FOLDER -1 R1.fq -2 R2.fq -r REFERENCE.fasta

ubuntu

The user should replace each of the following:

- OUTPUT_FOLDER: folder name inside test_dir to automatically save all files.
- R1.fq and R2.fq: corresponding names of the paired-end reads.
- REFERENCE.fasta: name of the fasta file to use.

Optional paramenters can be set:

- -l: use to indicate a specific label (default: SAMPLE)
- -p/--proc: Number of threads to use (default=2)
- -m/--mem: ram memmory available to use by the java machine (default=4).
- --min_quality: Phred value cutoff for filtering the reads (default=25)
- --timeout: seconds (per reference) before shutting down QuRe in case it freezes (default=600). Each time, the reads are downsampled and the step is repeated (6 times per reference)

Output files

- 14 Regardless of the operating system a set of folders should be created:
 - Filtering: Filtered and trimmed reads are stored here.
 - Multiple_Aligning: Alignment files are stored here.

- unmapped: reads extracted from the alignment are stored here.
- Additionally, for each reference a corresponding folder will be created.
- lofreq: the alignment to the first reference is store here. Also you will find the variant.vcf file here.

A set of output files will also be saved in test_dir/OUTPUT_FOLDER. Most relevant are:

- SAMPLE_haplotypes.fasta (and SAMPLE_haplotypes_aligned.fasta): the reconstructed haplotypes (first sequence is always the first sequence in the REFERENCE.fasta file and should NOT be considered an haplotype).
- SAMPLE_adjusted_proportions.txt: proportions of each reconstructed haplotype.
- SAMPLE_graphs.png: Concordance graph between lofreq and the reconstructed haplotypes. It should include a linear regression equation (the higher R-squared the better)