# Illumina metagenomic data analysis

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# Introduction

Welcome to the Illumina metagenomic data analysis manual. This manual contains a step by step guide for performing quality control, filtering host sequences, assembling reads into contigs, annotating the contigs, and then extracing viral contigs and their corresponding reads. In the final chapter of the manual we will show how to automate all of these steps into a single pipeline for speed and convenience.

# 1 Preparation



#### $\triangle$ Important!

In the following sections whenever a "parameter" in brackets {} is shown, the intention is to fill in your own filename or value. Each parameter will be explained in the section in detail.



🕊 Tip

Notice the small "Copy to Clipboard" button on the right hand side of each code chunk, this can be used to copy the code.

## 1.1 Activating the correct conda software environment

Anaconda is a software management tool that can be used for creating specific environments where bioinformatics software can be installed in Linux, as it manages all the dependencies of different softwares for you. We will use Conda to create a dedicated environment for this manual's workflow, ensuring that all required tools are installed and configured correctly.

To create the Conda environment for this manual, you'll need an environment.yml file that specifies the required software and their versions. Here's an example:

```
name: stable_env_1
channels:
  - conda-forge
  - https://repo.anaconda.com/pkgs/main
  - https://repo.anaconda.com/pkgs/r
dependencies:
  - python=3.9
  - bioconda::spades=3.15.4
  - bioconda::samtools=1.3.1
  - conda-forge::diamond=2.0.15
  - bioconda::snakemake=7.32.4
  - conda-forge::biopython=1.85
```

```
- bioconda::cd-hit-auxtools=4.8.1
- bioconda::fastp=0.22.0
- bioconda::bwa=0.7.18
- bioconda::seqkit=2.9.0
```

- 1. Copy the above text and save it to a file named environment.yml in your project directory.
- 2. In your terminal, navigate to your project directory (where you saved the environment.yml file) and create the Conda environment with the following command:

```
conda env create -f environment.yml
```

After the environment has been created, you need to activate it before you can use the software installed within it. To activate the environment, run:

```
conda activate stable_env_1
```

stable\_env\_1 is the name of the Conda environment you created. When the environment is activated, you'll see its name in parentheses at the beginning of your terminal prompt (stable\_env\_1). This indicates that you're now working within the Conda environment.

#### Note

We are now ready to start executing the code to perform quality control of our raw Illumina sequencing data in the next chapter.

# 2 Quality control



#### Important!

In the next steps we are going to copy-paste code, adjust it to our needs, and execute it on the command-line.

Please open a plain text editor to paste the code from the next steps, to keep track of your progress!

For simplicity's sake, most steps will be geared towards an analysis of a single sample. It is recommended to follow a basic file structure like the following below:

```
my_project/
                      # Contains the raw, gzipped FASTQ files
  raw_data/
      sample1_R1_001.fastq.gz
      sample1_R2_001.fastq.gz
      sample2_R1_001.fastq.gz
      sample2_R2_001.fastq.gz
  results/
                      # This is where the output files will be stored
  log/
                      # This is where log files will be stored
```

When running any command that generates output files, it's essential to ensure that the output directory exists before executing the command. While some tools will automatically create the output directory if it's not present, this behavior is not guaranteed. If the output directory doesn't exist and the tool doesn't create it, the command will likely fail with an error message (or, worse, it might fail silently, leading to unexpected results).

To prevent a lot of future frustration, create your output directories beforehand with the mkdir command as such:

```
mkdir -p results
mkdir -p log
# Create a subdirectory
mkdir -p results/assembly
```

# 2.1 Merging and decompressing FASTQ

Any file in linux can be pasted to another file using the cat command. zcat in addition also unzips gzipped files (e.g. .fastq.gz extension). If your files are already unzipped, use cat instead.

#### Modify and run:

```
zcat {input.folder}/{sample}_R1_001.fastq.gz > {output.folder}/{sample}_R1.fastq
zcat {input.folder}/{sample}_R2_001.fastq.gz > {output.folder}/{sample}_R2.fastq
```

- {input.folder} is where your raw .fastq.gz data is stored.
- {sample} is the name of your sample.
- {output.folder} is where your decompressed .fastq files will be stored.

## 2.2 Deduplicate reads

This step removes duplicate reads from the uncompressed FASTQ files. Duplicate reads can arise during PCR amplification or sequencing and can skew downstream analyses. We'll use the cd-hit-dup program to identify and remove these duplicates.

After cd-hit-dup command is finished, we'll remove the .clstr file that cd-hit-dup creates. This file contains information about the clusters of duplicate reads, but it's not needed for downstream analysis, so we can safely remove it to save disk space.

```
cd-hit-dup -u 50 -i {input.R1} -i2 {input.R2} -o {output.R1} -o2 {output.R2} > {logs}/{samplerm {output.dir}/*.clstr}
```

- {input.R1} and {input.R2} are the decompressed R1 and R2 reads from step 2.1.
- {output.R1} and {output.R2} are your deduplicated .fastq reads. Think of where you want to store your results, something like results/dedup/ will be sufficient, so output.R1 turns into result/dedup/SampleName\_R1.fastq, etc.
- {output.dir} This should be pointed to wherever your deduplicated reads are stored.

# 2.3 Running fastp quality control software

The fastp software is a very fast multipurpose quality control software to perform quality and sequence adapter trimming for Illumina short-read and Nanopore long-read data.

#### Run and modify:

```
fastp -i {input.R1} -I {input.R2} \
  -o {output.R1} -O {output.R2} \
  --unpaired1 {output.S} --unpaired2 {output.S} --failed_out {output.failed} \
  --length_required 50 \
  --low_complexity_filter \
  --cut_right \
  --cut_right_window_size 5 \
  --cut_right_mean_quality 25 \
  --thread {threads} \
  -j {output.J}/qc_report.json -h {output.H}/qc_report.html > {log} 2>&1
```

- {input.R1} and {input.R2} are the reads belonging to the deduplicated sample step 2.2.
- {output.R1} and {output.R2} are the the quality controlled .fastq filenames.
- {output.S} –unpaired1 and –unpaired2 tells fastp to write unpaired reads to a .fastq file. In our case, we write unpaired reads (whether they originated from the R1 or R2 file) to the same file, output.S.
- {output.failed} .fastq file that stores reads (either merged or unmerged) which failed the quality filters
- {sample} is the name of your sample.
- {output.J} is the directory for the json report file
- {output.H} is the directory for the html report file
- {log} is the directory for log files.

#### Note

{threads} is a recurring setting for the number of CPUs to use for the processing. On a laptop this will be less (e.g. 8), on an HPC you may be able to use 64 or more CPUs for processing. However, how much performance increase you get depends on the software.

# 2.4 Host filtering

This step removes reads that map to a host genome (e.g., human). This is important if you're studying metagenomes from a host-associated environment (e.g., gut microbiome, skin surface).

To improve computational speed and reduce memory usage, it is required to index your reference sequence before proceeding to the host filtering step.

```
# Index reference genome
bwa index {reference}

# Host filtering
bwa mem -aY -t {threads} {reference} {input.R1} {input.R2} | \
samtools fastq -f 4 -s /dev/null -1 {output.R1} -2 {output.R2} -
bwa mem -aY -t {threads} {reference} {input.S} | \
samtools fastq -f 4 -> {output.S}
```

- {input.R1} and {input.R2} are QC-filtered FASTQ files from step 2.3.
- {output.R1} and {output.R2} are FASTQ files (R1, R2) containing reads that did not map to the host genome.
- {input.S} are singleton reads from the previous step (output.S).
- {reference} is a reference genome sequence.

#### Note

We now have our quality controlled sequence reads which we can use to create an assembly in the next chapter.

# 3 De novo assembly

#### 3.1 metaSPades

We will perform de novo assembly of the host-filtered reads to create contigs (longer, assembled sequences) with SPades.

```
COUNT=$(cat {input.S} | wc -1)
if [[ $COUNT -gt 0 ]]
then
    spades.py -t {threads} \
    --meta \
    -o {output} \
    -1 {input.R1} \
    -2 {input.R2} \
    -s {input.S} > {log} 2 \times 2
else
    spades.py -t {threads} \
    --meta \
    -o {output} \
    -1 {input.R1} \
    -2 \{input.R2\} > \{log\} 2 > \&1
fi
```

The if [[ \$COUNT -gt 0 ]] condition checks if singleton reads are present. If so, we will include them with the -s option in the spades.py command

- {input.R1}, {input.R2} and {input.S} are host-filtered FASTQ files (R1, R2, and S) from step 2.3.
- {output} defines the directory where the assembly results will be stored.
- {log} specifies the log directory.

# 3.2 Renaming contigs:

We will add sample names to the beginning of each contig name. This will make sure that each sample's contig names are unique before we start aggregating contigs. If you are only dealing with a single sample, then this step can be seen as optional.

```
seqkit replace -p "^" -r "{sample}_" {input} > {output}
```

- {sample} is the sample name that will be placed in front of the contig name
- {input} is your contig file from step 3.1
- {output} is fasta file with the renamed contig

# 3.3 Aggregating contigs:

Next we will combine all renamed contigs into a single fasta file so we can perform taxonomic annotation across all samples in one go. Once again, this step can be seen as optional if you have a single sample.

#### cat {input} > {output}

- {input} are your renamed contig files from step 3.2
- {output} a single .fasta file

#### Note

We have created contigs that are ready for taxonomic annotation in the next chapter.

# 4 Taxonomic classification

#### 4.1 Diamond

Now we will annotate the aggregated contigs by assigning taxonomic classifications to them based on sequence similarity to known proteins in a database using diamond blastx.

The -f 6 parameter will ensure the output is in a tabular format. The 6 may be followed by a space-separated list of various keywords, each specifying a field of the output. For a full description of the output, please visit here.

```
diamond blastx \
-q {input} \
-d {db} \
-o {output} \
-f 6 qseqid sseqid pident length mismatch gapopen qstart qend sstart send evalue bitscore st
--taxonmap {tx} \
--threads {threads} \
-b 10 -c 1 > {log} 2>&1
```

- {input} is the conting file created in either step 3.3 or step 3.1, depending on your amount of samples.
- {db} is the protein database to be searched against.
- {output} is a .tsv file containing the annotation results.
- {tx} is a mapping file to convert protein accessions to taxonomic IDs.
- {log} is the log directory.

Multiple different databases and taxomaps can be found in: \\cifs.research.erasmusmc.nl\viro0002\workgr

# 4.2 Split annotation files

We will split the combined annotation file back into individual annotation files for each sample. This step can be seen as optional if you are dealing with a single sample.

Modify and run:

```
mkdir -p tmp_split

sed 's/_NODE/|NODE/' {input} | awk -F'|' '{
    identifier = $1;  # Construct the identifier using the first two fields

    output_file = "tmp_split/" identifier;  # Construct the output filename

if (!seen[identifier]++) {
        close(output_file);  # Close the previous file (if any)
        output = output_file;  # Update the current output file
    }

    print $2 > output;  # Append the line to the appropriate output file
}'

for file in tmp_split/*; do
    mkdir -p {output}/$(basename "$file")/;
    mv "$file" {output}/$(basename "$file")/diamond_output.tsv;
done

rmdir tmp_split
```

- {input} is the combined annotation file from step 4.1.
- {output} is a directory path. This directory will be automatically filled with subdirectories for each sample. In each subdirectory you will find a diamond\_out.tsv file.

## 4.3 Parsing diamond output

Now we will process the DIAMOND output files with a custom Python script called **post\_process\_diamond.py**. This script will further enrich taxonomic information for each contig based on the DIAMOND alignment results. If a contig has multiple matches in the database, it will select the best hit based on a combined score of bitscore and length. Lastly, it separates the contigs into two lists: those that were successfully annotated and unannotated.

This python script utilizes the biopython library.

```
python /mnt/viro0002/workgroups_projects/Bioinformatics/scripts/post_process_diamond.py \
-i {input.annotation} \
-c {input.contigs} \
-o {output.annotated} \
```

# -u {output.unannotated} \ -log {log}

- {input.annotation} is the annotation file step 4.2.
- {input.contigs} are the contigs from the SPAdes step 3.1.
- {output.annotated} is a .tsv file with a set of annotated contigs.
- {output.unannotated} is a .tsv file with a set of unannotated contig IDs.
- {log} is the log directory.

## Note

We can now move on to the final steps where we will create various files needed for downstream analysis.

# 5 Extracting Viral Sequences and Analyzing Mapped Reads

We will conclude the pipeline with various steps which will create valuable data files for further downstream analysis.

#### 5.1 Extract viral annotations

A basic grep command can be used to extract contigs that have been annotated as viral from the annotation file in **step 4.3**.

```
grep "Viruses$" {input.annotated} > {output.viral} || touch {output.viral}
```

- {input.annotated} is the annotation file from step 4.3.
- {output.viral} contains all of your viral contigs.

## 5.2 Mapping reads to contigs

We can map the quality-filtered and host-filtered reads back to the assembled contigs to quantify the abundance of different contigs in each sample. We will create a mapping file for paired reads (R1 and R2) and singletons (S) and then merge these two files together.

```
bwa index {input.contigs}
bwa mem -Y -t {threads} {input.contigs} {input.R1} {input.R2} | samtools sort - > {output.pa
bwa mem -Y -t {threads} {input.contigs} {input.S} | samtools sort - > {output.S}/tmp_singlets
samtools merge {output.merged}/contigs.bam {output.paired}/tmp_paired.bam {output.S}/tmp_singlets.rm {output.paired}/tmp_paired.bam {output.S}/tmp_singlets.bam
```

- {input.contigs} contains the assembled contigs from step 3.1.
- {input.R1}, {input.R2} and {input.S} are the quality controlled and host-filtered reads from step 2.4.
- {output.paired} is the directory for the .bam mapping file based on paired reads.
- {output.S} is the directory for the .bam mapping file based on singleton reads.
- {output.merged} is the directory for the merged .bam file.

## 5.3 Extract mapped reads

We will extract the reads for each annotation file that we've created.

```
#Create temporary BED file to extract the annotated mappings from the BAM of all mappings
cut -f1 {input.annotated} | awk -F'_' '{{print $0 "\t" 0 "\t" $4}}' > {output}/tmp.bed
samtools view -bL {output}/tmp.bed {input.mapped} > {output.annotated}

#Do the same for the unannotated contigs
cut -f1 {input.unannotated} | awk -F'_' '{{print $0 "\t" 0 "\t" $4}}' > {output}/tmp.bed
samtools view -bL {output}/tmp.bed {input.mapped} > {output.unannotated}

#Do the same for the viral contigs
cut -f1 {input.viral} | awk -F'_' '{{print $0 "\t" 0 "\t" $4}}' > {output}/tmp.bed
samtools view -bL {output}/tmp.bed {input.mapped} > {output.viral}
rm {output}/tmp.bed
```

- {input.annotated}, {input.unannotated} and {input.viral} are the .tsv annotation files from step 4.3 and 5.1.
- {input.mapped} is the combined .bam file from step 5.2.
- {output} is a folder where temporary .bed files will be stored.
- {output.annotated}, {output.unannotated} and {output.viral} are .bam files for each annotation input.

## 5.4 Count mapped reads

Next we count the number of reads that mapped to each contig in the annotated, unannotated, and viral BAM files.

```
samtools view -bF2052 {input.annotated} | seqkit bam -Qc - | awk '$2 != 0 {{print}}' > {output.annotated} | seqkit bam -Qc - | awk '$2 != 0 {{print}}' > {output.annotated} | seqkit bam -Qc - | awk '$2 != 0 {{print}}' > {output.annotated} | seqkit bam -Qc - | awk '$2 != 0 {{print}}' > {output.annotated} | seqkit bam -Qc - | awk '$2 != 0 {{print}}' > {output.annotated} | seqkit bam -Qc - | awk '$2 != 0 {{print}}' > {output.annotated} | seqkit bam -Qc - | awk '$2 != 0 {{print}}' > {output.annotated} | seqkit bam -Qc - | awk '$2 != 0 {{print}}' > {output.annotated} | seqkit bam -Qc - | awk '$2 != 0 {{print}}' > {output.annotated} | seqkit bam -Qc - | awk '$2 != 0 {{print}}' > {output.annotated} | seqkit bam -Qc - | awk '$2 != 0 {{print}}' > {output.annotated} | seqkit bam -Qc - | awk '$2 != 0 {{print}}' > {output.annotated} | seqkit bam -Qc - | awk '$2 != 0 {{print}}' > {output.annotated} | seqkit bam -Qc - | awk '$2 != 0 {{print}}' > {output.annotated} | seqkit bam -Qc - | awk '$2 != 0 {{print}}' > {output.annotated} | seqkit bam -Qc - | awk '$2 != 0 {{print}}' > {output.annotated} | seqkit bam -Qc - | awk '$2 != 0 {{print}}' > {output.annotated} | seqkit bam -Qc - | awk '$2 != 0 {{print}}' > {output.annotated} | seqkit bam -Qc - | awk '$2 != 0 {{print}}' > {output.annotated} | seqkit bam -Qc - | awk '$2 != 0 {{print}}' > {output.annotated} | seqkit bam -Qc - | awk '$2 != 0 {{print}}' > {output.annotated} | seqkit bam -Qc - | awk '$2 != 0 {{print}}' > {output.annotated} | seqkit bam -Qc - | awk '$2 != 0 {{print}}' > {output.annotated} | seqkit bam -Qc - | awk '$2 != 0 {{print}}' > {output.annotated} | seqkit bam -Qc - | awk '$2 != 0 {{print}}' > {output.annotated} | seqkit bam -Qc - | awk '$2 != 0 {{print}}' > {output.annotated} | seqkit bam -Qc - | awk '$2 != 0 {{print}}' > {output.annotated} | seqkit bam -Qc - | awk '$2 != 0 {{print}}' > {output.annotated} | seqkit bam -Qc - | awk '$2 != 0 {{print}}' > {output.annotated} | seqkit bam -Qc - | awk '$2 != 0 {{print}}' > {output.annotated} | seqkit bam -Qc - | awk
```

- {input.annotated}, {input.unannotated} and {input.viral} are the .bam output files from step 5.3.
- {output.annotated}, {output.unannotated} and {output.viral} are .tsv files containing the read counts for each .bam file

# 5.5 Extract contigs

Lastly, we will extract contigs for the annotated, unannotated and viral contigs.

```
seqkit grep -f <(cut -f1 {input.annotated}) {input.contigs} > {output.annotated}
seqkit grep -f <(cut -f1 {input.unannotated}) {input.contigs} > {output.unannotated}
seqkit grep -f <(cut -f1 {input.viral}) {input.contigs} > {output.viral}
```

- {input.annotated}, {input.unannotated} and {input.viral} are .tsv annotation files from steps 4.3 and 5.1.
- {input.contigs} is the .fasta file containing all contigs for a sample from step 3.1.
- {output.annotated}, {output.unannotated} and {output.viral} are .fasta files containing the contigs.

## Note

You can now move to the final chapter to automate all of the steps we've previously discussed.

# 6 Automating data analysis

In the previous chapters, you learned how to perform each step of the metagenomic data analysis pipeline manually. While this is a valuable learning experience, it's not practical for analyzing large datasets or for ensuring reproducibility in the long term.

We can make use of a tool called Snakemake to automate the previous steps into a single pipeline. With Snakemake, you can define the steps of your analysis in a Snakefile and then let Snakemake handle the execution, dependency management, and error handling.

### 6.1 Preparing to run the workflow.

To run the automated workflow, you'll need to make sure that your project directory is set up correctly and that you have activated the necessary Conda environment.

To make the project setup process even easier, we've created a command-line tool called prepare\_project.py. This tool automates the creation of the project directory, the sample configuration file (sample.tsv), and the general settings configuration file (config.yaml), guiding you through each step with clear prompts and error checking.

To proceed, make sure to activate the appropriate conda environment and check if the command line has (stable\_env\_1) in front.

```
conda activate stable_env_1 # Replace environment name is needed
```

Run prepare project.py with Python as follows:

```
python prepare_project.py -n {name} -p {project.folder} -r {reads}
```

- {name} is the name of your study, no spaces allowed.
- {project.folder} is your project folder.
- {reads} is the folder that contains your raw .fastq.gz files.

Now, move to your project directory. First check your current directory with the pwd command:

```
pwd
```

Change your current directory using cd if needed:

```
cd /{folder1}/{folder2} # Replace with the actual path to your project directory
```

Next, use the ls command to list the files in the project directory and check if the following files are present: sample.tsv, config.yaml and Snakefile.

- The sample tsv should have 3 columns: sample (sample name), fq1 and fq2 (paths to raw read files).
- The config.yaml contains more general information like the reference, database and taxonmap you supplied as well as the amount of default threads to use.
- The Snakefile is the "recipe" for the workflow, describing all the steps we have done by hand, and it is most commonly placed in the root directory of your project (you can open the Snakefile with a text editor and have a look).

# 6.2 Running the workflow

After setting everything up, we can redo the analysis for all samples in a single step. First we will test out a dry run to see if any errors appear. A dry run will not execute any of the commands but will instead display what would be done. This will help identify any errors in the Snakemake file.

Run inside of your project directory:

```
snakemake \
--snakefile \
Snakefile \
--use-conda \
--cores {threads} \
--dryrun
```

If no errors appear, then remove the **--dryrun** argument and run it again to fully execute the workflow.