

# Quality Control of Reads Using Trimmomatic (Cyverse) Version 2

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#### **Abstract**

A collection of protocols designed to guide the user in processing a viral metagenome from raw sequence data to assembly, and subsequent analysis. The user uses *actual* reads from <u>Ocean Sampling Day (2014)</u> and processes them entirely within Cyverse, a NSF-supported cyberinfrastructure.

Quality trimming of reads is important!

**Citation:** Benjamin Bolduc Quality Control of Reads Using Trimmomatic (Cyverse). **protocols.io** https://www.protocols.io/view/Quality-Control-of-Reads-Using-Trimmomatic-Cyverse-eygbftw

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### **Guidelines**

This is part of a larger protocol *Collection* that involves the end-to-end processing of raw viral metagenomic reads obtained from a sequencing facility to assembly and analysis using Apps (i.e. tools) developed by iVirus and implemented within the Cyverse cyberinfrastructure.

### **Before start**

To run this protocol, users must first <u>register</u> for Cyverse account. All data (both inputs and outputs) are available within Cyverse's data store at /iplant/home/shared/iVirus/ExampleData/

### **Protocol**

Quality control of reads

Step 1.

## **Open Trimmomatic**

Open Trimmomatic 0.35.0 from 'Apps'



#### Quality control of reads

Step 2.

## **Select Inputs**

Select the 'Inputs tab.

### For Reads file (for paired or unpaired):

Navigate to Community Data --> iVirus --> ExampleData --> Trimmomatic --> Inputs --> Reads.
Select OSD46\_R1\_shotgun\_workable.fastq.gz Alternatively, copy-and-paste the location:
/iplant/home/shared/iVirus/ExampleData/Trimmomatic/Inputs/Reads into the navigation bar and select the fastq.gz file.

For **Reads file (ONLY USE FOR PAIRED END)**: Only use this file if it's read pair 2. Don't try to double-up two single-end reads.

Navigate to Community Data --> iVirus --> ExampleData --> Trimmomatic --> Inputs --> Reads.
SelectOSD46\_R2\_shotgun\_workable.fastq.gz Alternatively, copy-and-paste the location:
/iplant/home/shared/iVirus/ExampleData/Trimmomatic/Inputs/Reads into the navigation bar and select the fastq.gz file.

### For **Illumina adapter file**:

Navigate to Community Data --> iVirus --> ExampleData --> Trimmomatic --> Inputs -->
 Adapters. Select TruSeq3-PE.fa Alternatively, copy-and-paste the location:
 /iplant/home/shared/iVirus/ExampleData/Trimmomatic/Inputs/Adapters into the navigation bar
 and select the fasta file.

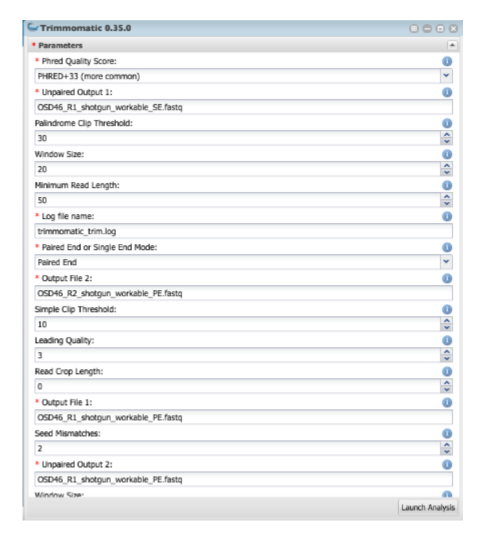


Step 3.

### **Select Parameters**

Select the 'Parameters' tab.

The default options will suffice for this example.



### Quality control of reads

Step 4.

# **Launch Analysis**

Run the job!

Trimmomatic can take minutes to hours to complete.

### Quality control of reads

Step 5.

### **Results**

Trimmomatic will generate 5 output files. 2 paired files where both forward and reverse survived read trimming, 2 unpaired files where their pair/mate did not survive, and a log file.

Expected results can be found from the 'Outputs' directory of Trimmomatic.

