

# Supplementary Methods & Tables - nf-core/viralmetagene: A Novel Pipeline for Untargeted Viral Genome Reconstruction

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## Data Availability

The nf-core/viralmetagene pipeline is freely available at <https://github.com/nf-core/viralmetagene>. The raw data and analysis code is available on <https://github.com/Joon-Klaps/nf-core-viralmetagene-manuscript/>.

## Supplementary Methods

### S1. Influence of Scaffold Reference on Consensus Genome

We simulated mixed HIV-1 samples containing paired-end read data based on four genomes (MN090277.1, MN090188.1, MN090240.1, and MZ766668.1) spanning genetic diversity from 80–99% average nucleotide identity (ANI). Thirteen sample configurations were created with varying abundance ratios to model both single-strain infections (100% abundance) and coinfections with different abundance ratios (75:25, 50:50, 25:75; Supplementary table 1). Reads were generated using InSilicoSeq v2.0.1 (Gourlé et al. 2019) using ‘kde’ mode and ‘lognormal’ abundance for a ‘MiSeq’ model.

We investigated how reference genome selection affects consensus accuracy on the same sample mixture dataset as described above. While similar references ( $\geq 96\%$ ) minimally impact consensus quality, divergent references introduce up to 187 nucleotide mismatches (Supplementary Figure 1). Nf-core/viralmetagene addresses this challenge by automatically selecting the most appropriate reference genome based on sequence similarity; when no suitable genome is found in the supplied database, the contigs are scaffolded to a *de novo* contig (specifically, the cluster representative), ensuring accurate final consensus genomes.

Sample mixtures were run with nf-core/viralmetagene using default settings unless specified otherwise. Reference sequences used to compare the influence of the reference during scaffolding are the same sequences used for read generation, in addition to the databases, Virosaurus (Gleizes et al. 2020) and the Reference Viral Database (Goodacre et al. 2018). Visualisation was done in R (R Core Team 2021) using tidyverse (Wickham et al. 2019).

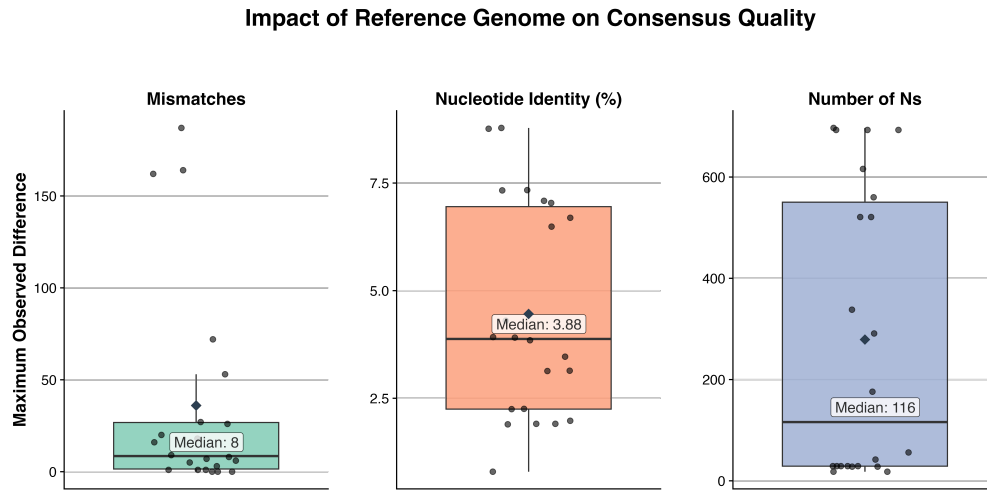


Figure 1: Boxplot of maximum observed differences between consensus sequences generated with different reference genomes during scaffolding. Mean highlighted by a diamond.

## S2. Evaluation of nf-core/viralmetagenome on public data

A public metagenomic dataset was constructed by selecting a randomized subset of 2,000 records from the NCBI Virus database, targeting the following viruses: '*Orthonairovirus haemorrhagiae*', '*Mammarenavirus lassaense*', '*Zika virus*', '*West Nile virus*', '*Monkeypox virus*', '*Influenza A virus*', '*Severe acute respiratory syndrome coronavirus 2*', and '*Human respiratory syncytial virus A*'. Records were filtered to retain only those with associated SRA data, sequenced on the Illumina platform, and with a LibraryStrategy other than 'Amplicon'. From the resulting pool, 28 samples were randomly selected and downloaded using nf-core/fetchngs. Samples were then run with nf-core/viralmetagenome using `-cluster_method 'mmseqs-cluster'` due to sequence size limitations of cdhit, `-keep_unclassified false` and `-skip_read_classification` to speed up the analysis.

For plant pathogens, two samples were randomly selected from the SRA after querying for the following viruses: '*Tobacco mosaic virus*', '*Tomato spotted wilt virus*', '*Cucumber mosaic virus*', and '*Potato virus Y*'. These were also downloaded using nf-core/fetchngs and processed with viral-metagenome using the same parameters, with the addition of the Virosaurus plant virus database as annotation database and reference database.

48 **Supplementary Tables**

Table 1: Overview of computational tools and methods used in the nf-core/viralmetagenome pipeline. The pipeline integrates multiple bioinformatics tools across different stages of viral metagenomic analysis, from preprocessing to consensus calling. Each tool was selected for its specific capabilities and performance characteristics relevant to viral sequence analysis.

Stage	Step	Tool	Explanation	Reference
Pre-processing	Read trimming	fastp	A high-performance preprocessing solution built in C++ that consolidates multiple quality control operations into a single workflow. Fastp delivers enhanced processing speed compared to traditional alternatives, like Trimmomatic, while handling adapter removal, quality assessment, base correction, read and filtering. Fastp also features automated adapter detection capabilities for various Illumina sequencing protocols.	(Chen et al. 2018)
		Trimmomatic	A comprehensive preprocessing solution tailored for Illumina sequencing platforms with specialised paired-end read handling capabilities. Provides multiple pre-processing options, including adapter removal, quality-based trimming using sliding window approaches, and length-based filtering.	(Bolger et al. 2014)
	Complexity filtering	bbduk	A high-performance, multi-threaded tool designed to combine various data-quality-related operations, including trimming, filtering, and masking, into a single pass. It is particularly effective for removing host and other contaminant DNA using k-mer matching strategies, which is crucial for accurate metagenomic analysis.	(Bushnell 2022)
Metagenomic diversity	Taxonomy classification	PRINSEQ++	A C++ implementation that significantly improves upon its predecessor, prinseq-lite.pl, in terms of computational efficiency. It offers extensive quality control features, including filtering by length, GC content, quality scores, N content, and sequence complexity (entropy/DUST score).	(Cantu et al. 2019)
		Kraken2	An advanced taxonomic classification system with an improved k-mer-based approach through optimised memory usage and enhanced processing speed. Incorporates protein-level search capabilities for improved detection of divergent viral sequences, with a library of pre-built RefSeq indexes.	(Wood et al. 2019)

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Stage	Step	Tool	Explanation	Reference
		Kaiju	A protein-level taxonomic classifier that employs exact matching algorithms on translated sequences using the Burrows-Wheeler transform. It can optionally allow amino acid substitutions and tends to have higher sensitivity and precision compared to k-mer-based classifiers (Kraken2). Kaiju is particularly effective for organisms with limited reference representation.	(Menzel et al. 2016)
Assembly & polishing	Assembly	SPAdes	A de Bruijn graph-based assembler designed to overcome challenges in complex sequencing data, such as non-uniform coverage (using multisized de Bruijn graphs) and variable insert sizes. The Spades suite contains multiple specialised modes rnaSPAdes, coronaSPAdes, metaSPAdes for extra flexibility.	(Meleshko et al. 2021)
		MEGAHIT	A succinct de Bruijn graph-based assembler specifically designed for large and complex metagenomic datasets. MEGAHIT has demonstrated the ability to generate larger assemblies with longer contig N50 and average contig length. Its relatively low memory requirements make it well-suited for handling large datasets.	(Li et al. 2016)
		Trinity	A powerful modular transcriptome reconstruction tool, capable of fully reconstructing a large fraction of transcripts, including alternative splice isoforms and transcripts from recently duplicated genes, even in the absence of a reference genome. It partitions sequence data into individual de Bruijn graphs, processing each independently to extract full-length isoforms and resolve paralogous genes.	(Grabherr et al. 2011)
	Read alignment	BWA-MEM2	An optimized version of the BWA-MEM algorithm enhanced with Intel-specific acceleration feature, providing approximately a twofold speedup in alignment throughput over BWA-MEM while ensuring identical SAM output. It is highly efficient for aligning short reads to large reference genomes.	(Vasimuddin et al. 2019)

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Stage	Step	Tool	Explanation	Reference
		BWA	A foundational and widely used Burrows-Wheeler Aligner (BWA-MEM variant) that efficiently maps short reads and long sequences to large reference genomes. It supports long gaps and dynamically adjusts error rates based on sequence length, making it robust to sequencing errors.	(Li 2013)
		bowtie2	A fast and memory-efficient alignment tool for aligning sequencing reads to long reference sequences, such as mammalian genomes. Bowtie2 supports gapped, local, and paired-end alignment modes, combining high speed, sensitivity, and accuracy.	(Langmead et al. 2019)
	Clustering	CD-HIT-EST	A highly efficient and widely used program for clustering large sets of protein or nucleotide sequences. Implementation constraints limit processing to genomes under 10Mbp with minimum 0.8. The program employs a greedy incremental algorithm.	(Li and Godzik 2006)
		MMSeqs	A powerful software suite for fast and sensitive deep clustering and searching of large protein sequence sets. It is significantly faster than other tools like BLASTclust and can cluster large databases down to low sequence identity thresholds. MMseqs offers a variety of clustering modes for a lot of flexibility.	(Steinegger and Söding 2017)
		VSEARCH	An open-source, versatile alternative for USEARCH. VSEARCH supports processing very large datasets, limited primarily by available memory.	(Rognes et al. 2016)
		Mash	A non-alignment based clustering technique that applies MinHash algorithms for rapid genome and metagenome distance estimation. It compresses large sequences into small, representative sketches, enabling ultra-fast estimations of global mutation distances. Mash creates compact sequence representations enabling ultra-fast similarity assessments and large-scale clustering operations, though accuracy decreases below 95% genome similarity.	(Ondov et al. 2019)

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Stage	Step	Tool	Explanation	Reference
		vRhyme	A machine learning-based binning solution specifically developed for viral genome recovery from metagenomic datasets. VRhyme addresses the unique challenges of viral sequences, such as the lack of universal marker genes. Addresses unique viral sequence characteristics through supervised learning approaches combined with coverage-based analysis across multiple samples. Generates high-quality viral bins with minimal computational overhead, though it may produce empty results when insufficient viral evidence is detected.	(Kieft et al. 2022)
Variant & consensus calling	Variant & consensus calling	iVar	A computational package specifically designed for viral amplicon-based sequencing, integrating functions like consensus and variant calling (including iSNVs and insertions/deletions). It is a key component of best-practice pipelines for reconstructing consensus genomes from viral sequencing data.	(Grubaugh et al. 2019)
		BCFtools	A robust suite of utilities for manipulating variant calls in VCF and BCF formats. Provides extensive functionality for variant calling using multiallelic models, data filtering, file merging, and consensus sequence generation through variant application.	(Danecek et al. 2021)

Table 2: Overview of simulated HIV-1 sample mixtures with the used reference genome and meta-data.

Sample name	Reference	Added genome	Similarity to reference (%)	Reads (M)	Read % MN090277	Read % added genome
sample_1	MN090277.1	MN090188.1	98.7	4	75	25
sample_2	MN090277.1	MN090188.1	98.7	4	50	50
sample_3	MN090277.1	MN090188.1	98.7	4	25	75
sample_4	MN090277.1	MN090240.1	96.7	4	75	25
sample_5	MN090277.1	MN090240.1	96.7	4	50	50
sample_6	MN090277.1	MN090240.1	96.7	4	25	75
sample_7	MN090277.1	MZ766668.1	83.5	4	75	25
sample_8	MN090277.1	MZ766668.1	83.5	4	50	50
sample_9	MN090277.1	MZ766668.1	83.5	4	25	75
sample_10	MN090277.1	MN090277.1	100.0	1	100	0
sample_11	MN090277.1	MN090188.1	98.7	1	0	100
sample_12	MN090277.1	MN090240.1	96.7	1	0	100
sample_13	MN090277.1	MZ766668.1	83.5	1	0	100

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