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## Exploring the archaeal virosphere by metagenomics

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### Abstract

During the past decade, environmental research has demonstrated that archaea are abundant and widespread in nature, and play important ecological roles at a global scale. Currently, however, the majority of archaeal lineages cannot be cultivated under laboratory conditions and are known exclusively or nearly exclusively through metagenomics. A similar trend extends to the archaeal virosphere, where isolated representatives are available for a handful of model archaeal virus-host systems. Viral metagenomics provides an alternative way to circumvent the limitations of culture-based virus discovery and offers insight into the diversity, distribution and environmental impact of uncultured archaeal viruses. Presently, metagenomics approaches have been successfully applied to exploring the viromes associated with various lineages of extremophilic and mesophilic archaea, including Asgard archaea (Asgardarchaeota), ANME-1 archaea (Methanophagales), thaumarchaea (Nitrososphaeria), altiarchaea (Altiarchaeota) and marine group II archaea (Poseidoniales). Here, we provide an overview of methods widely used in archaeal virus metagenomics, covering metavirome preparation, genome annotation, phylogenetic and phylogenomic analyses, and archaeal host assignment. We hope that this summary will contribute to further exploration and characterization of the enigmatic archaeal virome lurking in diverse environments.

**Key words:** archaeal viruses, Archaea, metagenomics, hyperthermophiles, major capsid protein, CRISPR spacers, host prediction, virome

### 1. Introduction

Recent advances in high-throughput genome sequencing and computational approaches have transformed our appreciation of the diversity, ubiquity and importance of archaea in natural environments [1-9]. Similar to bacteriophages [10-15], archaeal viruses represent one of the major factors controlling the diversity and metabolic activity of archaeal populations [16, 17]. Although culture-based approaches are revealing extraordinary morphological and genomic diversity of archaeal viruses isolated from extreme geothermal and hypersaline environments [18-23], only a handful of virus isolates infecting mesophilic archaea have been described thus far [24, 25], limiting our appreciation of their diversity and ecological impacts. Nevertheless, the culture-based virus discovery efforts are increasingly complemented by culture-independent metagenomic approaches. For instance, metagenomics has uncovered a number of family-level groups of viruses and mobile genetic elements associated with Asgard archaea [26-29], a prominent phylum Asgardarchaeota widely considered to represent the ancestors of eukaryotes [1, 2]; ANME-1 clade (order Methanophagales), a group of methane oxidizing archaea implicated in modulation of greenhouse gas emission [30]; methanogenic archaea [31, 32]; ubiquitous marine archaea of the order Poseidoniales [33-35] and ammonia oxidizing thaumarchaea [33, 36-38]; as well as Altiarchaeota, abundant primary producers in subsurface ecosystems [39]. These studies have provided precious insights into the parts of the archaeal virosphere which are currently inaccessible through classical culture-based techniques. In this chapter, we provide an overview of protocols and practices used in archaeal virus metagenomics, covering metavirome preparation, genome annotation, phylogenetic and phylogenomic analyses, abundance profiling and archaeal host assignment.

### 2. Materials

#### 2.1 Bioinformatics tools

Software packages and bioinformatic tools commonly used for the analysis of archaeal virus genomes and proteins are listed in [Table 1](#).

**Table 1.** Software and web servers

Tool	Available at	Function	Reference
Fastp	github.com/OpenGene/fastp	Reads quality control	[40]
Trimmomatic	github.com/usadellab/Trimmomatic	Reads quality control	[41]
Megahit	github.com/voutcn/megahit	Sequence assembly	[42]
metaSPAdes	cab.spbu.ru/software/spades	Sequence assembly	[43]
Blast+	ftp.ncbi.nlm.nih.gov/blast/executables	Sequence homology search	[44]
VirSorter2	github.com/jiarong/VirSorter2	Viral sequence identification	[45]
VIBRANT	github.com/AnantharamanLab/VIBRANT	Viral sequence identification	[46]
DeepVirFinder	github.com/jessieren/DeepVirFinder	Viral sequence identification	[47]
Cenote-Taker 2	github.com/mtisza1/Cenote-Taker2	Viral sequence identification	[48]
Geneious Prime	Biomatters, Inc.	Sequence extension and reads mapping	N/A
ContigExtender	github.com/dengzac/contig-extender	Sequence extension	[49]
CheckV	bitbucket.org/berkeleylab/CheckV	Virus genome completeness assessment	[50]
CRISPRCasFinder	crisprcas.i2bc.paris-saclay.fr/CrisprCasFinder/Index	CRISPR spacer extraction	[51]
CRISPRDetect	github.com/davidchyou/CRISPRDetect_2.4	CRISPR spacer extraction	[52]
CD-HIT	cd-hit.org	Sequence redundancy removal	[53]
tRNAscan-SE	github.com/UCSC-LoweLab/tRNAscan-SE	tRNA gene detection	[54]
Batch CD-Search	www.ncbi.nlm.nih.gov	Protein annotation	[55]
eggNOG-mapper	eggnog-mapper.embl.de	Protein annotation	[56, 57]
DRAM-v	github.com/WrightonLabCSU/DRAM	Viral AMG detection	[58]
HostG	github.com/KennthShang/HostG	Host prediction	[59]
MArVD2	bitbucket.org/MAVERICLab/marvd2	Archaeal virus identification	N/A
WISH	github.com/soedinglab/wish	Host prediction	[60]
PHISDetector	github.com/HIT-ImmunologyLab/PHISDetector	Host prediction	[61]
RaFAH	sourceforge.net/projects/rafah	Host prediction	[62]
iPHoP	bitbucket.org/srouxjgi/iphop	Host prediction	N/A
Pharokka	github.com/gbouras13/pharokka	Genome annotation	[63]
HHsearch	github.com/soedinglab/hh-suite	Genome annotation	[64]
VIRFAM	biodev.cea.fr/virfam	Genome annotation	[65]
T-Coffee	tcoffee.crg.eu	Sequence alignment	[66]
MUSCLE	www.drive5.com/muscle	Sequence alignment	[67]
PROMALS3D	prodata.swmed.edu/promals3d	Sequence alignment	[68]
trimAl	trimal.cgenomics.org/	Alignment trimming	[69]
PhyML	www.atgc-montpellier.fr/phyml	Phylogeny construction	[70]
FastTree	www.microbesonline.org/fasttree	Phylogeny construction	[71]
IQ-TREE	www.iqtree.org	Phylogeny construction	[72]
iTOL	itol.embl.de	Tree visualization	[73]
Evolview	www.evolgenius.info/evolview	Tree visualization	[74]
vConTACT2	bitbucket.org/MAVERICLab/vcontact2	Gene-sharing network construction	[75]
compareM	github.com/dparks1134/CompareM	comparative genomic analyses	N/A
Easyfig	easyfig.sourceforge.net	Genome map construction	[76]
Clinker	github.com/gamcil/clinker	Genome map construction	[77]
Cytoscape	cytoscape.org	Network visualization	[78]
ViPTree	www.genome.jp/viptree/	Comparative genome analyses	[79]
R package	www.r-project.org/	Heatmap construction	N/A
Bowtie2	bowtie-bio.sourceforge.net/bowtie2	Reads mapping	[80]

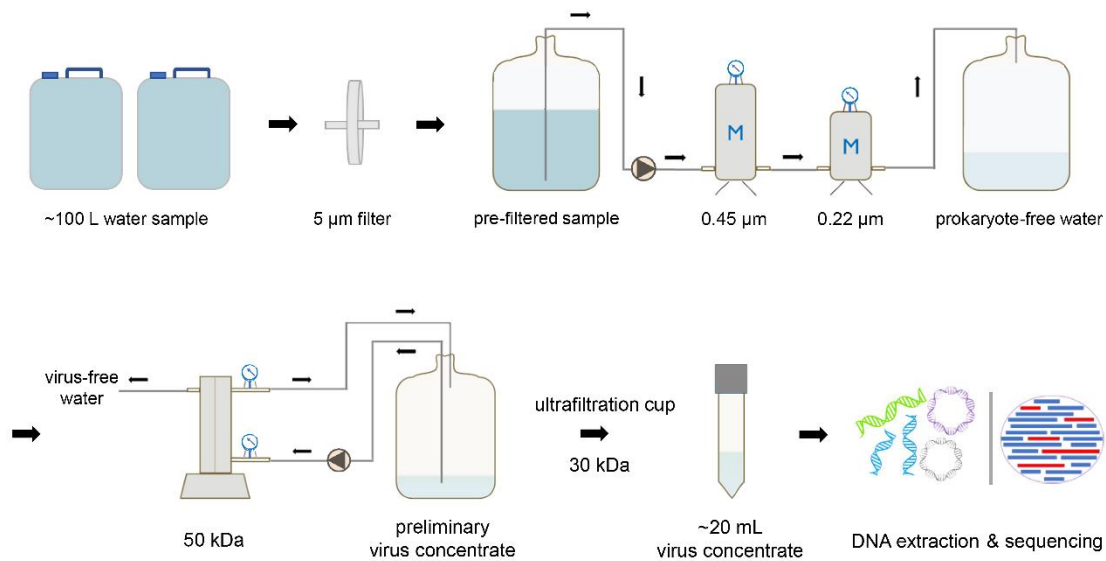
### 3. Methods

Archaea typically represent an abundant or even dominant component of the microbial communities in geothermal (e.g., terrestrial hot spring), hypersaline (e.g., crystallizer pond) and certain marine (e.g., estuary, sediments) environments [81-85]. Not surprisingly, archaeal viruses are also prevalent in such archaea-dominated ecosystems. Thus, to exemplify the protocols for exploration of the archaeal virome, in this chapter, we focus on samples originating from hot springs, hypersaline ponds and marine environments. Generally, preparation of the viral DNA for archaeal virus metagenomics is not different from that used for other prokaryotic DNA viruses and, given that there are many excellent dedicated studies on this topic (e.g., [86-91]), this part will be discussed only briefly. Notably, all currently known archaeal viruses contain DNA genomes [92], thus protocols for RNA virus discovery will not be considered. For environmental samples with low archaeal abundance, whenever possible, an archaeal enrichment step [93] might be considered prior to collecting the viral DNA for the downstream metagenomics studies.

#### 3.1. Concentration of viral particles and DNA purification (adapted from [88-91, 94, 95])

For environmental samples with low cellular (and viral) density, large initial volumes, in the range of tens or hundreds of liters, are typically processed. For instance, in the case of marine samples, ~100 L of the water sample can be filtered through a 5 µm pore size filter to remove large inorganic particles and eukaryotic cells (Figure 1). Next, the prefiltered water sample is further passed through two successive

0.45  $\mu\text{m}$  and 0.22  $\mu\text{m}$  pore size filter-columns to remove the remaining cells, mainly archaea and bacteria. The virus particles can be concentrated by pumping the cell-free water through a 50 kDa cutoff tangential-flow filtration (TFF) cassette until the retentate reaches a volume of  $\sim 500$  mL (see **Note 1** and **Note 2**). Finally, the viral preparation can be further concentrated to  $\sim 20$  mL using 30 kDa cutoff ultrafiltration cups or ultracentrifugation.



**Figure 1.** A workflow for the virus metagenome preparation.

Chemical flocculation is an alternative approach which has been successfully applied for concentration of viruses from seawater samples. This method depends on iron (III) chloride to precipitate viruses which are recovered by filtration onto large-pore size membranes and then resuspended using a buffer containing magnesium and a reductant (ascorbic acid or oxalic acid) at slightly acid pH [87, 96].

Next, total nucleic acids are extracted from the virus concentrate and prepared for metagenomic sequencing (see **Note 3** and **Note 4**). Generally, libraries are subjected to 150-bp, 250-bp, or 300-bp paired-end sequencing on Illumina platforms such as HiSeq X Ten, NovaSeq 6000 and MiSeq, respectively. Furthermore, the long-read sequencing, for example, provided by PacBio (Pacific Biosciences) or Nanopore (Oxford Nanopore Technologies), is becoming increasingly popular and affordable, providing single-molecule long-read sequences which can cover the entire virus genomes or considerably facilitate the assembly of large virus genomes [97].

### 3.2 Quality control of sequencing reads

Sequencing adapters and low-quality reads ( $<Q20$ ) are trimmed off by using quality control tools such as Fastp and Trimmomatic to obtain a clean metagenomic dataset (see **Note 5**).

Example:

```
'fastp -i input_R1.fastq.gz -o clean_R1.fastq.gz -l input_R2.fastq.gz -O clean_R2.fastq.gz'
```

### 3.3 Sequence assembly

In the next step, clean reads are assembled to generate metagenomic contigs. The assembly is one of the most memory-consuming steps. A server with at least 200 GB of RAM is required. It is advisable to use more than one sequence assembler, as they produce slightly different results. Tools such as Megahit and metaSpades are applied to assemble the clean reads into metagenomic contigs. Contigs with sequence length over 1 kb are retained for the subsequent analyses.

Examples:

```
'megahit -1 clean_R1.fastq.gz -2 clean_R2.fastq.gz -o output_assembly -t 48'
```

```
'spades.py --meta -1 clean_R1.fastq.gz -2 clean_R2.fastq.gz -t 48 -m 360 -o output_assembly'
```

### 3.4 Identification of viral contigs

It is not possible to completely eliminate prokaryotic DNA from environmental metaviromes, because cellular DNA can be packaged into virus capsids and be transferred from one host to another by a process known as general transduction [98]. Alternatively, cellular DNA can be protected from DNase digestion within virus-sized extracellular membrane vesicles, which are known to be produced by many archaeal species [99-102]. To assess the extent of contamination with prokaryotic DNA, the 16S ribosomal RNA gene sequences can be downloaded from the SILVA database (latest release, [www.arb-silva.de/](http://www.arb-silva.de/)) and used as queries to search against the assembled metagenomic sequences. Depending on the results, instead of using all sequences derived from the 'virus metagenome', it is recommended to implement a step of viral sequence sorting.

Tools such as VirSorter2, VIBRANT, DeepVirFinder or Cenote-Taker 2 (here and elsewhere, see Table 1 for references) can be used to identify and extract viral sequences from the assembled contigs (see **Note 6**).

Examples:

```
'virsorter run -w vir_outputVS2 -i Assembly_contigs.fasta --include-groups dsDNAphage --high-confidence-only'
```

```
'python run_cenote-taker2.py -c Assembly_contigs.fasta -r vir_outputCT2 -p True -m 256 -t 48 --exact_dtrs True'
```

### 3.5 Sequences extension and sequence quality check

The viral contigs can be extended by recruiting the sequencing reads that overlap with the edges of the de novo assembled contigs [103, 104]. To this end, all identified viral sequences can be pooled together and used as seed sequences to perform the reference assembly. Geneious Prime 'map to reference' function can be used to compare all the metagenomic reads to the seed sequences. If there are significant matches (i.e.,  $\geq 30$  bp overlap and  $\geq 95\%$  overlap identity), the reads will be assembled to the corresponding seed sequence, yielding longer viral contigs. This procedure can be repeated until the seed sequence ceases to extend (see **Note 7**). Alternatively, sequence extension can be performed using ContigExtender. Example:

```
'extender_wrapper.py --m1 clean_R1.fastq --m2 clean_R2.fastq --out VirContigExtension --min-overlap-length 30 --stop-length 55 --threads 48 vir_contigs.fasta'
```

'Repeat Finder' (Geneious Prime plugin) can be used to detect direct terminal repeats (DTR) and inverted terminal repeats (ITR) on the extended viral contigs. If the contig contains either DTR or ITR regions, the sequence can be considered to represent a complete circular or linear virus genome, respectively. Note that for formal classification, the International Committee on Taxonomy of Viruses (ICTV) only considers viruses with complete genome sequences [105] (see **Note 8**).

CheckV can also be used to estimate genome completeness by examining the terminal repeats and similarity of the viral contigs to related viral genomes.

Example:

```
'checkv end_to_end viral_contigs.fasta checkv_output -t 48 -d ~/checkv_database'
```

Note that due to its inherent dependency on viral reference genomes, CheckV shows inadequate performance when assessing the completeness of truly novel viral genomes which have only distant relatives or no relatives in the reference database.

### 3.6 Assignment of archaeal hosts

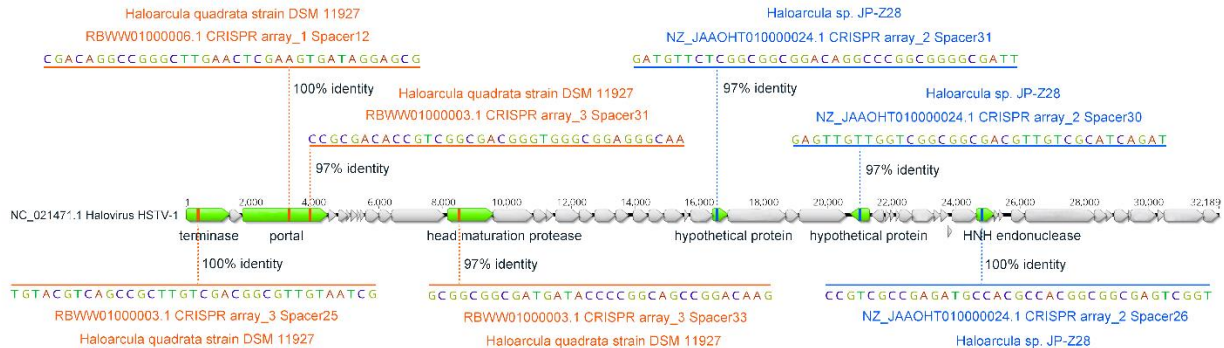
Arguably, the most challenging step in archaeal virus metagenomics is differentiation between viruses infecting bacteria and archaea and accurate assignment of viruses to their archaeal hosts. Below we introduce several host assignment approaches which have been previously applied to identify archaeal viruses in metagenomic datasets.

#### 3.6.1 CRISPR spacer targeting

CRISPR-Cas is an adaptive immune system encoded by most archaeal species [106]. The spacer sequences stored in CRISPR arrays represent the immune memory of past encounters with foreign mobile genetic elements. Therefore, by matching CRISPR spacers to the corresponding protospacer sequences in the viral genomes, it is possible to identify archaeal virus-host pairs in metagenomic dataset. See an



example of archaeal CRISPR spacers matching an archaeal virus genome in [Figure 2](#). This method is currently by far the most reliable among host prediction approaches. However, it should be noted that in the case of archaeal viruses, the accuracy of host prediction is typically limited to the level of family [107] or even order [30]. For instance, in the case of rudiviruses, viruses infecting *Saccharolobus* and *Metallosphaera* species were targeted by spacers from CRISPR arrays of *Metallosphaera* and *Saccharolobus*, respectively [107].



**Figure 2.** Seven CRISPR spacers from two *Haloarcula* species match the genome of *Haloarcula sinaiensis* tailed virus 1 (HSTV-1).

To assemble the database of CRISPR spacers, all archaeal genomic sequences can be downloaded from the Genome Taxonomy Database (GTDB) or any other genome database. The retrieved archaeal genomes can then be analyzed using CRISPRDetect or CRISPRCasFinder to detect CRISPR arrays and extract the archaeal CRISPR spacer sequences.

Example:

```
'CRISPRCasFinder -in archaeal_seqs.fasta -out CRISPR_results -md 10 -t 15 -mr 25 -xr 55 -ms 25 -xs 55 -pm 0.7 -px 2.5 -s 55 -fl 120 -cpuM 48'
```

CD-HIT can be used to remove redundant spacers with 100% sequence identity cutoff. In order to maintain the targeting specificity, it is recommended to discard spacers with sequence length shorter than 25 bp.

Example:

```
'cd-hit -i archaeal_spacer.fasta -o unique_archaeal_spacers.fasta -c 1 -M 36000 -T 12 -d 0'
```

Next, a BLASTn database should be prepared from the extracted viral contigs:

```
'makeblastdb -i viral_contigs.fasta -out nt_db/viral_contigs -dbtype nucl'
```

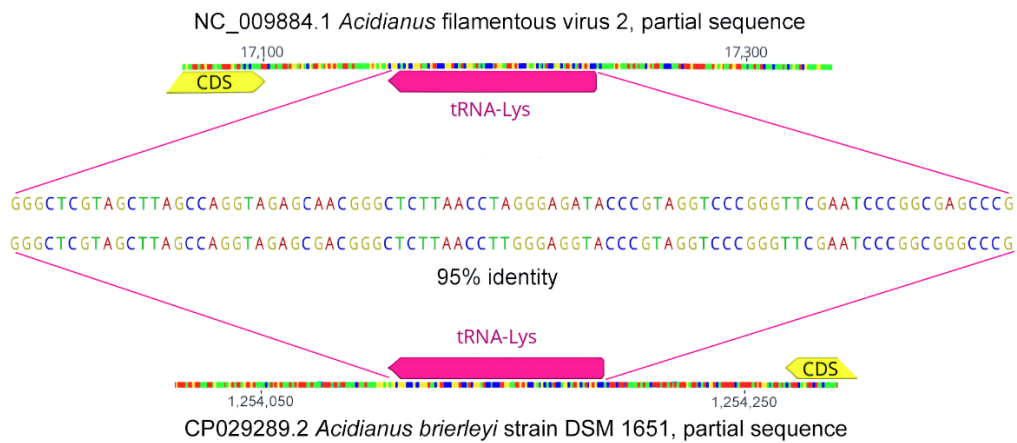
The archaeal spacers can then be used as queries to search against the viral contig database (see **Note 9** and **Note 10**). Hits with at least 95% coverage and 95% identity are considered as valid protospacers and link the protospacer-containing viral contigs to the corresponding archaeal hosts.

```
'blastn -query unique_archaeal_spacers.fasta -db nt_db/viral_contigs -out ArSpacers_vs_VirContigs.xls -outfmt 6 -word_size 7 -dust no -qcov_hsp_perc 95 -perc_identity 95 -num_threads 48'
```

More than one spacer hit increases the reliability of the host prediction. The sequences of the CRISPR-targeted viral contigs should be retrieved for the subsequent analyses.

### 3.6.2 tRNA gene matching

tRNAs are an integral part of the cellular protein translation system. Certain archaeal viruses occasionally also encode host-derived tRNA genes [18, 108]. Therefore, the tRNA gene match between a virus and an archaeal genome is highly suggestive of a virus-host relationship. See an example of tRNA gene match between an archaeal virus and its host in [Figure 3](#).



**Figure 3.** The tRNA-Lys gene encoded by the genome of *Acidianus filamentous virus 2* shares 95% identity with a counterpart from the genome of *Acidianus brierleyi* strain DSM 1651.

The tRNA genes encoded by viral contigs can be predicted using tRNAscan-SE 2.0 with '-A' option (archaeal mode):

```
'tRNAscan-SE viral_contigs.fasta -A -o viral_tRNA.results -a viral_tRNA.fasta'
```

The viral tRNA gene sequences should then be extracted and used as queries in BLASTn search against the GTDB archaeal sequences (or other archaeal sequence database with available taxonomy information).

```
'makeblastdb -in archaeal_seqs.fasta -out nt_db/archaeal_seqs -dbtype nucl'
```

```
'blastn -query viral_tRNA.fasta -db nt_db/archaeal_seqs -out Vir_tRNA_vs_ArSeqs.xls -outfmt 6 -word_size 16 -dust no -qcov_hsp_perc 95 -perc_identity 95 -num_threads 48'
```

Hits with at least 95% coverage and 95% identity can be considered as genuine matches, providing host assignments for the tRNA-encoding viral contigs (see **Note 11**).

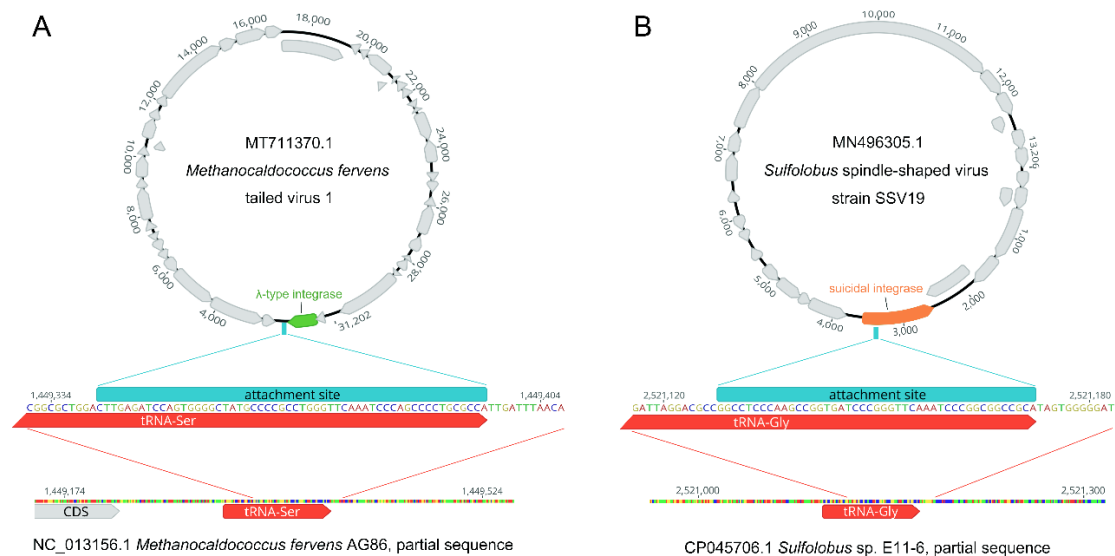
### 3.6.3 Matching of the viral and host attachment site(s)

Many archaeal viruses with circular dsDNA genomes encode integrases of the tyrosine recombinase superfamily and are commonly found as integrated proviruses within archaeal genomes [109-114]. The integration involves integrase-mediated homologous recombination between identical sequences of variable lengths on the viral and cellular genomes, known as the viral and archaeal attachment sites, attV and attA, respectively [115]. In the viral genome, the attV is typically located in the vicinity of or within the integrase gene, whereas the most common attA site occupies the 3'-proximal regions of the tRNA genes, although cases of integration into 5'-distal regions of tRNA genes, intergenic regions as well as protein-coding genes have been also described [112, 113]. The identity between the viral and archaeal attachment sites can be used for the host assignment for temperate viruses which integrate into the genome of their host [116]. See two examples of attachment site sharing between archaeal viruses and their respective hosts in [Figure 4](#).

To identify the attachment sites within the viral genomes, the integrase genes have to be identified first. This can be done using batch protein annotation tools, such as Batch CD-Search or eggNOG-mapper v2. Once identified, the sequences of the viral integrase genes along with the upstream and downstream non-coding regions ( $\leq 500$  bp) can be extracted and used as queries in BLASTn searches against archaeal sequences to detect the potential attachment sites.

```
'blastn -query integrase_region.fasta -db nt_db/archaeal_seqs -out Vir_integrase_vs_ArSeqs.xls -outfmt 6 -word_size 6 -dust no -perc_identity 95 -num_threads 48'
```

Hits with at least 95% coverage and 95% identity can be considered as genuine matches (see **Note 12**).



**Figure 4.** Examples of attachment sites located next to (A) and within (B) the integrase genes. A. The attachment site of *Methanocaldococcus fervens* tailed virus 1 is located next to the gene encoding the  $\lambda$ -type integrase of the tyrosine recombinase superfamily (green) and is identical to the 3'-proximal region of the host's tRNA-Ser gene (red). B. The attachment site of *Sulfolobus* spindle-shaped virus 19 (SSV19) is located within its integrase gene (orange) and is identical to the 3'-proximal region of the host's tRNA-Gly gene (red).

### 3.6.4 MCP homology searches

The ability to form virions distinguishes viruses from other types of mobile genetic elements and cellular organisms [117]. The major capsid proteins (MCP) are highly diverse, with some structurally related MCPs being widespread in viruses infecting hosts from different domains of life and others being specific to particular domains, including archaea [118]. Regardless, at the sequence level, MCPs are typically virus family or order specific. Therefore, homology searches using MCP as a signature protein has become one of the most commonly used approaches for the identification of new prokaryotic viruses in metagenomic dataset [33, 36, 37, 109, 119].

To assemble a database of archaeal virus MCPs, all archaeal virus protein sequences can be downloaded from the NCBI virus database ([www.ncbi.nlm.nih.gov/labs/virus/vssi/#/](http://www.ncbi.nlm.nih.gov/labs/virus/vssi/#/)) and the MCP sequences extracted. Once the MCP database is assembled, open reading frames in the metagenomic viral contigs can be predicted using the software Prodigal with '-p meta' option. The corresponding *in silico* translated protein sequences can then be used as BLASTp queries to search against the archaeal virus reference MCP database.

```
'prodigal -i viral_contigs.fasta -a ORFs_viral_contigs.fasta -p meta'
```

```
'makeblastdb -in Ref_ArVir_MCPs.fasta -out prot_db/ Ref_ArVir_MCPs -dbtype prot'
```

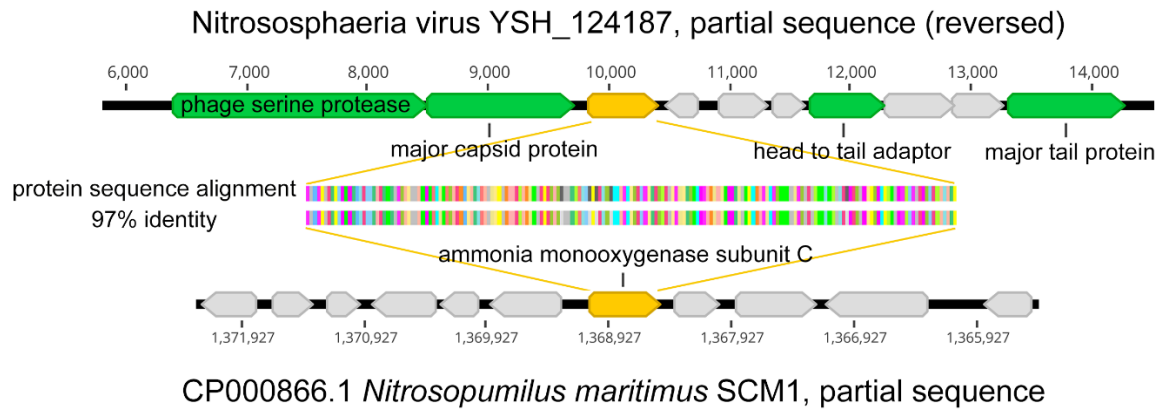
```
'blastp -query ORFs_viral_contigs.fasta -db prot_db/Ref_ArVir_MCPs -out  
viral_contigs_vs_Ref_ArVir_MCPs.blastp -evalue 1e-5 -num_threads 48'
```

BLASTp hits with E-value  $<10^{-5}$ ,  $\geq 30\%$  identity,  $\geq 50\%$  coverage, and  $\geq 100$  bit score can be considered significant. The corresponding viral contigs should be extracted for further analyses. More accurate host assignment can be inferred from phylogenetic analysis of the MCP sequences from the reference viruses and viruses assembled from metagenomes.

### 3.6.5 Host-specific AMGs

It is increasingly recognized that viruses can modulate the metabolic processes within the infected prokaryotes by expressing auxiliary metabolic genes (AMGs) [12]. In order to efficiently tinker with the host metabolism, the virus-encoded versions of the metabolic genes are usually recognizably similar to the host homologs [37, 38]. This property offers a possibility to trace potential virus-host pairs from environmental samples by analyzing the virus-encoded AMGs. Figure 5 shows an example of high sequence similarity between the ammonia monooxygenase subunit C (AmoC) encoded by a virus and an archaeon, *Nitrosopumilus maritimus*.





**Figure 5.** The ammonia monooxygenase subunit C encoded by the *Nitrososphaeria* virus YSH\_124187 [33] shares 97% protein sequence identity with the counterpart encoded by *Nitrosopumilus maritimus* SCM1.

Viral AMGs can be predicted using the software DRAM-v, the mode of DRAM for viral sequences (see **Note 13**).

```
'DRAM-v.py annotate -i viral-combined-for-dramv.fa -v viral-affi-contigs-for-dramv.tab -o dramv_annotate --skip_trnscan --threads 48'
```

```
'DRAM-v.py distill -i dramv_annotate/annotations.tsv -o dramv-distill'
```

Viral AMGs should then be extracted and used as BLASTp queries to search against the archaeal proteome. The archaeal proteome database can be constructed using the following commands:

```
'prodigal -i archaeal_seqs.fasta -a ORFs_archaeal_seqs.fasta -p meta'
```

```
'makeblastdb -in ORFs_archaeal_seqs.fasta -out prot_db/archaeal_proteome'
```

Next, the viral AMGs should be assigned to specific archaeal species using the following command:

```
'blastp -query viral_AMGs.fasta -db prot_db/archaeal_proteome -out Vir_AMGs_vs_archaeal_proteome.blastp -evalue 1e-5 -num_threads 48'
```

Hits with at least 90% identity and 90% coverage are considered particularly indicative of a virus-host relationship, whereas hits with lower sequence identity should be evaluated more cautiously.

### 3.6.6 Host prediction tools (see **Note 14**)

A number of host prediction tools have been recently developed for prokaryotic viruses, e.g., HostG, WiSH, PHISDetector, RaFAH, iPHoP, and others. These tools rely on certain databases and yield results with host prediction with associated confidence scores.

Example (HostG):

```
'python run_Speed_up.py --contigs viral_contigs.fasta --len 1000 --t 0'
```

Example (iPHoP):

```
'iphop predict --fa_file viral_contigs.fasta --db_dir ~/iPHoP_db --out_dir iphop_prediction --num_threads 48'
```

In addition, the tool MARVD2 has been specifically developed for the identification of archaeal viruses from a set of viral contigs.

```
'MARVD2.py -i viral_contigs.fasta -o marvd_prediction --db-pvog ~/AllvogHMMprofiles.hmm --db-nr ~/nr.faa --db-accession2tax ~/prot.accession2taxid --marine-jackhmmmer-db ~/pVOG_prots_ref_marine_pVOG.faa --viral-refseq-txt ~/viruses.txt --pvog-dir ~/pVOGs --cpu-count 48 -load-model ~/rf_model.pkl'
```

### 3.7. Genome annotation

All viral contigs can be automatically annotated using such tools as Batch CD-Search, eggNOG-mapper or Pharokka. However, for comprehensive functional annotation of the complete or near-complete archaeal virus genomes, it is advisable to use sensitive hidden Markov model (HMM) profile–profile comparisons with HHsearch v3.3.0 against the publicly available databases: CDD, Pfam, Protein Data Bank (PDB), uniprot\_sprot\_vir70, and PHROG. In the case of *Caudoviricetes*, viral structural proteins can be also predicted using VIRFAM.

### 3.8 Phylogenetic & phylogenomic analyses

Relationships between related viruses can be assessed using different methods, including single gene phylogenies, network or phylogenomic analyses as briefly detailed below.

#### 3.8.1 Phylogenetic analysis

For the purpose of understanding the relationship between evolutionarily related viruses, a protein conserved in a given group of viruses has to be selected. The commonly used viral hallmark proteins include MCP, portal protein or large subunit of the terminase and other genome packaging ATPases. The protein sequences can be aligned using tools such as MUSCLE, T-COFFEE or PROMALS3D, followed by removal of highly divergent, uninformative positions, e.g., using trimAl. Maximum likelihood phylogenetic trees can be constructed using IQ-tree or PhyML. Both IQ-tree and PhyML can select the amino acid substitution model best fitting the given dataset. In the case of very large datasets, approximate maximum likelihood trees can be calculated using FastTree. The phylogenies can be annotated and visualized using iTOL v5 or Evolview v3.

#### 3.8.2 Gene-sharing networks

The relationships between the identified archaeal viruses and other known prokaryotic viruses can be assessed using network analysis. vConTACT2 can be used to generate the gene-sharing networks with the latest prokaryotic virus database. Given that nodes (viral genomes) are connected only when they share three gene families, ideally, the length of input viral sequences should exceed 10 kb. Alternatively, the relationships between the viral genomes can be explored using bipartite networks, which include two types of nodes, viral genomes (type 1 nodes) connected through shared gene families (type 2 nodes) [120]. The resulting networks can be visualized using Cytoscape.

#### 3.8.3 Viral proteomic tree

A virus proteomic tree is a dendrogram that represents global genomic relationships between viral sequences calculated from comparison of all protein sequences encoded by a given set of viruses. The viral proteomic tree generally corresponds well with the established virus taxonomy [33, 35]. The virus proteomic tree can be calculated for any given dataset using ViPTree, which can be either locally installed or run through the web server (see **Note 15**).

#### 3.8.4 Estimation of the orthologous protein fraction

For classification purposes, it is useful to know the fraction of genes shared with other viruses (i.e., the degree of relatedness between viruses). For example, members of head-tailed archaeal viruses of the same family in the class *Caudoviricetes* generally share ~20–50% of orthologous genes, while viruses from different families share less than 10% [18]. The fraction of orthologous proteins can be estimated using the CompareM software toolkit with the following command:

```
'comparem aai_wf -e 0.0001 -p 30 -a 50 -c 4 ~/seqs output_results'
```

Based on the results (see **Note 16**), taxonomic classification can be tentatively assigned to the sequenced viruses according to the established taxon-specific demarcation criteria. It is important to note that depending on the virus group, different parameters can be adopted, resulting in different estimates of the orthologous fractions.

### 3.9 Abundance and distribution

To gain ecological and evolutionary insights into how archaeal viruses interact with their hosts and environments, it is important to explore the distribution and abundance of these viruses in different ecosystems. This information can be obtained by recruiting sequencing reads from metagenomes to the identified archaeal virus genomes, using read mappers, such as Bowtie2 or Geneious.

Example (Bowtie2):

```
'bowtie2-build ref_virus.fasta ref_virus'
```

```
'bowtie2 -x ref_virus -1 clean_R1.fastq -2 clean_R2.fastq -S results.sam'
```

The relative abundance of viruses in any particular sample or environment can be estimated by mapping the sequence reads from a metavirome to the reference genomes and expressed as **Reads recruited Per Kb** of genome per **Gb** of metagenome (RPKG). This way, the sequencing depth is normalized and is comparable to the distribution and abundance of the viruses in different environments.

#### 4. Notes

1. The sample is passed through the TFF system by using a peristaltic pump and the pressure of the flow within the system should always be kept below 10 p.s.i. (~62 kPa) to avoid disruption of the viral particles [89].
2. One should consider the pH of the water sample to choose the corresponding filters. Ideally, all filtration operations should be conducted in a cold room to minimize the enzymatic degradation of virus particles and/or nucleic acids. Reservoirs, e.g., 500 mL flasks or bottles and 10-, 25- and 100-liter plastic containers, should be sterilized before use.
3. Filter the obtained viral concentrate through a 0.22  $\mu\text{m}$  pore size filter to remove any potentially remaining prokaryotes. The absence of cells in the virus concentrate and the number of purified virus particles can be verified by fluorescence microscopy after staining of an aliquot of the sample with nucleic-acid-staining fluorescent dyes, such SYBR Green or DAPI (4',6-diamidino-2-phenylindole), as described previously [121-123].
4. The majority of "free" DNA should be filtered out during the process of virus concentration. Nevertheless, an additional step of DNase treatment prior to extraction of the viral DNA from the virus particles can be performed to further reduce the contamination of the preparation with cellular DNA [124].
5. The result of reads quality control can be visualized as graphical and statistical reports by FastQC ([www.bioinformatics.babraham.ac.uk/projects/fastqc/](http://www.bioinformatics.babraham.ac.uk/projects/fastqc/)).
6. For tools that are reference-based, it is necessary to apply the most updated databases. For more reliable identification of viral contigs, at least 2 different tools can be used in parallel.
7. It is advisable to manually check the mapping of assembled reads to avoid the possible mis-assembly.
8. For reporting sequences of uncultivated virus genomes, the genome quality is one of the requirements by the Minimum Information about an Uncultivated Virus Genome (MIUViG), with other qualifiers being information about virus origin, assembly tool, virus identification software, genome type, taxonomic classification, biogeographic distribution and *in silico* host prediction [125]. For recommendations on official classification of uncultivated viruses, see [126].
9. Since the default BLASTn parameters are not optimal for short sequences (e.g., 30 bp), it is recommended to use a word size of 7 and dust filtering turned off to identify the targets of CRISPR spacers [127, 128].
10. Although highly reliable, the CRISPR-targeting host assignment approach is dependent on the richness of the CRISPR spacer database available for a particular host organism. To further enrich the spacer database, one may consider extracting spacers from taxonomically unclassified sequences by identifying group-specific CRISPRs (e.g., Asgard-archaea-specific CRISPRs [26]). In parallel or alternatively, additional CRISPR spacers for the archaeal groups of interest can be recovered by amplifying CRISPR arrays with CRISPR-specific PCR primers from the environmental sample from which viral metagenome is being prepared (e.g., [129]).
11. The tRNA genes of unknown viruses (here, the extracted viral contigs) can be also searched against the tRNA genes of viruses for which the hosts have been assigned. The shared tRNA genes could indicate that the two viruses infected the same host.
12. Some attachment sites are as short as 8 bp [112], but such hits can hardly be considered significant without further validation. Thus, to avoid short random matches and to improve the specificity of this host assignment method, we recommend using matches with nucleotide alignment length not shorter than 25 bp.
13. Follow this Standard Operating Procedure [www.protocols.io/view/viral-sequence-identification-sop-with-virsorter2-5qpvoqebg4o/v3](http://www.protocols.io/view/viral-sequence-identification-sop-with-virsorter2-5qpvoqebg4o/v3).
14. The performance of these host-prediction tools depends on the representation of the actual host organism within the initial training and reference dataset. Thus, it is important to make sure that the suspected host species were included in the training dataset. Otherwise, reasonable results are hardly to be expected. We recommend using these predictions only as supporting evidence to complement the host predictions inferred using other methods.
15. ViPTree web server also provides an informative genome map visualization useful for comparative genomics. Genome maps can be also compared using Easyfig or Clinker.
16. The output file of compareM can be converted into a matrix using tidyR (R package) and visualized using pheatmap (R package).

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