



Taxonomy of prokaryotic viruses: 2018-2019 update from the ICTV Bacterial and Archaeal Viruses Subcommittee

Evelien M. Adriaenssens¹ · Matthew B. Sullivan² · Petar Knezevic³ · Leonardo J. van Zyl⁴ · B. L. Sarkar⁵ · Bas E. Dutilh^{6,7} · Poliane Alfenas-Zerbini⁸ · Małgorzata Łobocka⁹ · Yigang Tong¹⁰ · James Rodney Brister¹¹ · Andrea I. Moreno Switt¹² · Jochen Klumpp¹³ · Ramy Karam Aziz¹⁴ · Jakub Barylski¹⁵ · Jumpei Uchiyama¹⁶ · Rob A. Edwards^{17,18} · Andrew M. Kropinski^{19,20} · Nicola K. Petty²¹ · Martha R. J. Clokie²² · Alla I. Kushkina²³ · Vera V. Morozova²⁴ · Siobain Duffy²⁵ · Annika Gillis²⁶ · Janis Rumnieks²⁷ · İpek Kurtböke²⁸ · Nina Chanishvili²⁹ · Lawrence Goodridge¹⁹ · Johannes Wittmann³⁰ · Rob Lavigne³¹ · Ho Bin Jang³² · David Prangishvili^{33,34} · Francois Enault³⁵ · Dann Turner³⁶ · Minna M. Poranen³⁷ · Hanna M. Oksanen³⁷ · Mart Krupovic³³

Published online: 11 March 2020
© Springer-Verlag GmbH Austria, part of Springer Nature 2020

Abstract

This article is a summary of the activities of the ICTV's Bacterial and Archaeal Viruses Subcommittee for the years 2018 and 2019. Highlights include the creation of a new order, 10 families, 22 subfamilies, 424 genera and 964 species. Some of our concerns about the ICTV's ability to adjust to and incorporate new DNA- and protein-based taxonomic tools are discussed.

Introduction

The prokaryotic virus community is represented in the International Committee on Taxonomy of Viruses (ICTV) by the Bacterial and Archaeal Viruses Subcommittee. Since our last report [1], the committee composition has changed, and over 200 taxonomic proposals (TaxoProps) were submitted to the ICTV Executive Committee (EC) for evaluation and approval. Below, we summarize these new developments.

Changes in subcommittee membership

In an effort to increase the geographical diversity of members, we appointed representatives from Australia (İ. Kurtböke), Canada (L. Goodridge), Georgia (N. Chanishvili), India (B.L. Sarkar), Russia (V.V. Morozova) and Ukraine (A. Kushkina). Lamentably, Nicola Petty has had to withdraw from our committee.

Taxonomic updates

Over the past two years, our subcommittee has proposed the creation of a new order (*Tubulavirales*), ten new families (*Autographiviridae*, *Chaseviridae*, *Demereciviridae*, *Drexleriviridae*, *Finnlakeviridae*, *Halspiviridae*, *Herelleviridae*, *Ovaliviridae*, *Plectroviridae*, *Thaspiviridae*), 22 new subfamilies, 424 new genera and 964 new species. In previous versions of this update, we listed each of the new taxa, but this is not practical with the introduction of almost 500 new genera. Instead, we briefly introduce the new order and each of the new families below.

Tubulavirales

The new bacteriophage order *Tubulavirales* comprises the rearranged family *Inoviridae* (five existing genera and 18 new genera) and the new family *Plectroviridae* (two existing genera and one new genus). The members of the order *Tubulavirales* possess (+)ssDNA genomes and have a unique morphology, visible as flexible filaments or rigid rods, due to helical symmetry of the capsid. This combination of particle morphology and genome type is unique among viruses, with the exception of representatives of archaeal viruses from the family *Spiraviridae*, which, however, are substantially different in virion organization and gene content [2]. The

Handling Editor: Sead Sabanadzovic.

Extended author information available on the last page of the article

genomes of the new order members contain from 4 to 15 ORFs and are similarly organized, with a modular organization. These phages infect Gram-positive, Gram-negative or cell-wall-less bacteria. A prominent characteristic of this order is that its member viruses enter neither typical lytic nor lysogenic cycles. Instead, virions are released from cells by extrusion, causing a chronic infection without killing the host. The phages belonging to the rearranged family *Inoviridae* infect Gram-negative and Gram-positive bacteria, and virions appear as long and flexible filaments. The members of the new family *Plectroviridae*, previously classified as members of the family *Inoviridae*, infect cell-wall-less bacteria and exhibit the morphology of rigid rods.

Autographiviridae

In 2008, Lavigne et al. [3] re-examined the taxonomy of the family *Podoviridae* and, based on shared protein homologs, defined three genera of T7-like phages within a single subfamily, the *Autographivirinae*. The defining characteristic of the subfamily was the presence of a virion-encoded RNA polymerase (RNAP) from which the subfamily derived its name; “auto” and “graphein” derived from the Greek, meaning “self-writing” or “self-transcribing”. The defining morphological characteristics of all these viruses is that they possess a small (ca. 60 nm in diameter) isometric head attached to a short tail. Their genomes are composed of a linear terminally redundant dsDNA of approximately 41 kb, and all encode a large (> 100 kDa) single-subunit RNA polymerase, which is responsible for middle and late transcription. Further common characteristics of these phages include conservation of gene arrangement and apparently genus-specific lysis cassettes and RNAP specificity loops. A new analysis of these viruses reveals nine subfamilies and 132 genera whose members infect bacteria of the classes Betaproteobacteria and Gammaproteobacteria, and the phylum Cyanobacteria. Many of the genera are undersampled and further analyses are required to demonstrate that the new family is not polyphyletic. It is anticipated that further revisions will be necessary in the future.

Chaseviridae

This taxon is named in honour of Martha Cowles Chase (1927–2003), who, together with Alfred Hershey, experimentally demonstrated that DNA rather than protein is the genetic material. The first isolated phage of this type is ϕ EcoM-GJ1 [4, 5]. Members of this family are myoviruses with isometric heads ~62 nm in diameter and contractile tails ~120 nm in length. The hosts belong to the genera *Escherichia*, *Erwinia*, *Pectobacterium*, *Shewanella* and *Aeromonas*, all members of the class Gammaproteobacteria. The genomes possess ~2600-bp direct terminal repeats, are

on average 54.2 kbp (46.5 mol% G + C) in length and encode about 77 proteins and, in some cases, a tRNA. Except where we have identified genera with two species, this group as a whole displays very little overall DNA sequence relatedness, presumably because the GC-content varies from 43.6 to 52.8 mol%. At the protein level, *Erwinia* phage vB_EamM-Y2 and *Shewanella* phage SppYZU05 share 34.8% homologs. Common proteins include the RNA polymerase, DNA polymerase, primase and exonuclease. vConTACT 2.0 [6, 7] clusters *Aeromonas* phage pAh6-C, *Escherichia* phage ϕ iEcoM-GJ1, *Erwinia* phage vB_EamM-Y2, *Pectobacterium* phage PM1, and *Shewanella* phage Spp001 into one cluster (VC 244_0). Our analysis suggests the presence of two subfamilies: one including the genera *Loessnervirus*, *Suwonvirus*, *Carltongylesvirus* and *Faunusvirus*, and the other including the genera *Yushanvirus* and *Pahsxtavirus*.

Demerecviridae

This new family contains three new subfamilies (*Markadamsvirinae* [including the genera *Tequinavirus*, *Epsteptimavirus*, and *Haartmanvirus*]; *Mccorquodalevirinae* [including the genera *Myunavirus* and *Hongcheonvirus*]; and *Ermolyevavirinae* [including the genera *Cetovirus*, *Vipunavirus*, and *Suttonboningtonvirus*]. The hosts for these phages are *Aeromonas*, *Escherichia*, *Klebsiella*, *Pectobacterium*, *Proteus*, *Providencia*, *Salmonella*, *Shigella*, *Vibrio*, and *Yersinia* species. The genome of coliphage T5 (AY543070.1) is 121.75 kbp in length, with a mol%GC content of 39.3 and encodes 162 proteins and 24 tRNAs. It is characterized by long (10,219-bp) direct terminal repeats. The average length of the genomes ranges from 106.2 to 122.8 kbp, but in most cases no evidence has been presented for terminal redundancy. The average mol%GC content varies from 39.1 to 45.2. This new family is named in honour of Milislav Demerec (1895–1966), the Croatian-American geneticist who pioneered the work on bacteriophages, and who, together with the Italian-American physicist Ugo Fano (1912–2001), first isolated *E. coli* phage T5 in 1945 from a mixture provided to them by Tony L. Rakieta (Long Island College of Medicine, USA).

Drexlerviridae

When originally proposed in 1996, the “T1-like phages” included a single species, *Enterobacteria phage T1*. Since that time, numerous taxonomic changes have been introduced, with the genus name changing to “*T1likevirus*” to “*Tunalikevirus*” to “*T1virus*” and ultimately to “*Tunavirus*”. In 2015, the subfamily *Tunavirinae* was introduced, consisting of five genera: *Kp36virus*, *Rogue1virus*, *Rtpvirus*, *T1virus* and *Tlsvirus* [8]. In 2018, three additional genera were included: *Eclunavirus*, *Hanrivervirus* and *Sertoctavirus*,

and the genera *Rogue1virus*, *T1virus* and *Kp36virus* were renamed *Rogunavirus*, *Tunavirus* and *Webervirus*, respectively. In total, 31 species have been classified. GenBank now contains the genome sequences of 84 T1 related phages, calling for re-examination of the relationships within the subfamily. The hosts belong predominantly to the genus *Escherichia*, but *Cronobacter*-, *Enterobacter*-, *Klebsiella*-, *Pantoea*- and *Shigella*-specific isolates have been isolated and sequenced as well. These hosts are all members of the class Gammaproteobacteria. The phage genomes are on average 48.9 kbp long (46.0 mol% G + C) and encode about 79 proteins and 0–2 tRNAs. This resulted in the addition of 12 new genera and three new subfamilies (*Tempevirinae* [including the genera *Tlsvirus*, *Hanrivervirus*, and *Warwickvirus*], *Rogunavirinae* [including the genera *Rogunavirus*, *Eastlansingvirus*, *Wilsonroadvirus*, and *Lindendrivevirus*], and *Braunvirinae* [including the genera *Rtpvirus*, *Shandongvirus*, *Loudonvirus*, and *Guelphvirus*]). This family is named in honour of Henry Drexler, who pioneered research on phage T1.

Finnlakeviridae

This new taxon was created for the new “Finnish lake” virus species *Flavobacterium virus FLiP* (in the new genus *Finnlakevirus*). FLiP is the sole member of this family and is the only known icosahedral single-stranded (ss) DNA bacteriophage with an internal membrane. FLiP and its host bacterium *Flavobacterium* sp. strain B330 were isolated from a boreal lake, Lake Jyväsjärvi, Jyväskylä, Finland [9]. The FLiP genome is a circular ssDNA molecule of 9,174 nucleotides. Nine of the 16 open reading frames have no significant sequence identity to other sequences in the databases. The diameter of the virion is 59 nm, and the vertices have elongated spike structures. The capsid (T=21) is formed by trimeric major capsid proteins with a double β -barrel fold. Network analysis of the major capsid protein sequences has shown that FLiP forms its own group among the bacterial and archaeal icosahedral viruses and related proviruses [10].

Halspiviridae

The spindle-shaped virus His1 has a linear double-stranded DNA genome of 14,464 bp with terminal inverted repeats and encodes a protein-primed family B DNA polymerase [11]. His1 infects the halophilic archaeon *Haloarcula hispanica* (phylum Euryarchaeota) [12] and was originally classified as a member of the unassigned genus *Salterprovirus* [11]. The major capsid protein of His1 shows sequence similarity to those of members of the family *Fuselloviridae*,

which infect hyperthermophilic acidophiles of the order Sulfolobales (phylum Crenarchaeota) [13, 14]. However, unlike His1, fuselloviruses have circular dsDNA genomes and do not encode their own DNA polymerases. Because His1 is sufficiently distinct from all other known cultured viruses, the genus *Salterprovirus* has been assigned to a separate new family, the *Halspiviridae* (hal- for halophilic, spi- for spindle-shaped). Furthermore, the species *His 1 virus* has been renamed “*Salterprovirus His1*”.

Herelleviridae

The family *Herelleviridae* was named after Félix d’Hérelle, celebrating the 100th anniversary of his discovery of bacteriophages [15]. Members of this family are large myoviruses with a virulent lifestyle that infect bacteria of the phylum Firmicutes. The family was extended from the subfamily *Spounavirinae*, with *Bacillus* phage SPO1 being the type member [16], and includes four other subfamilies: *Jasinkavirinae*, *Brockvirinae*, *Bastillevirinae* and *Twortvirinae*. All members of the family *Herelleviridae* share a set of core genes, including the terminase large subunit, major capsid protein, tail tube, tail sheath, and tail tape measure protein. They form a monophyletic clade within the tailed phages when using single and concatenated marker gene phylogenies [15], as well as when using GRAViTy [17], vCONTACT2 [6], and the phage proteomic tree approach [18]. Members of the same genus in this family share at least 50% nucleotide sequence identity over the entire genome, whereas members of the same subfamily share at least 25% translated genome sequence identity as determined with tBLASTx.

Ovaliviridae

The new family *Ovaliviridae* (from the Latin *ovalis*, for oval) is represented by *Sulfolobus ellipsoid virus 1* (SEV1) [19], which harbors a linear double-stranded DNA genome of 23,219 bp and encodes 38 predicted proteins, most of which have no known function. SEV1 has been classified as the first representative of a new family because of its unique morphology, unusual architecture of the virion, and lack of gene content similarity to other characterized viruses. The virion of SEV1 contains a protein capsid with 16 regularly spaced striations and an 11-nm-thick envelope [19]. The virus acquires its envelope intracellularly and exits the host cell through hexagonal pyramidal portals that perforate the host cell envelope.

Sulfolobus ellipsoid virus 1 is designated as the type species of the genus *Alphaovalivirus*.

Thaspiviridae

The virions of Nitrosopumilus spindle-shaped virus 1 (NSV1) measure 64 ± 3 nm in diameter and 112 ± 6 nm in length, with a short tail at one pole [20]. This morphology is similar to that of fuselloviruses and the halspivirus His1 [13]. The genome of NSV1 is a linear dsDNA molecule of 27,548 bp, terminating with 176-bp-long inverted repeats. NSV1 encodes 48 predicted proteins, only one of which (protein-primed family B DNA polymerase) displays significant sequence similarity to the proteins of other known archaeal or bacterial viruses. NSV1 is the first and, thus far, the only virus isolated on ammonia-oxidising archaea of the phylum Thaumarchaeota [20]. NSV1 has been classified into the species *Nitmarvirus NSV1* within the genus *Nitmarvirus* (for *Nitrosopumilus maritimus* virus) and the new family *Thaspiviridae* (*Tha-* for *th*aumarchaeal, *spi-* for *sp*indle-shaped viruses).

Online (10th) Report of the ICTV

Virus Taxonomy: The Classification and Nomenclature of Viruses - The Online (10th) Report of the ICTV is freely accessible at <http://ictv.global/report>, and summaries of the chapters on each virus family are published in the Journal of General Virology. We would like to acknowledge the hard work of David Prangishvili and Mart Krupovic for contributing chapters on archaeal (*Ampullaviridae* [21], *Bicaudaviridae* [22], *Globuloviridae* [23], *Guttaviridae* [24], *Spiraviridae* [25], *Tristromaviridae* [26], and bacterial (*Plasmaviridae* [27]) viruses.

Final thoughts from the Chair, Professor Andrew Kropinski

I would hazard a guess that the Bacterial and Archaeal Viruses Subcommittee (BAVS) has changed the taxonomy of the viruses that are their responsibility over the past five years more than any other group within the ICTV at any point in history. This has resulted in a taxonomy which is much closer aligned to that of other viruses than ever before. To indicate how phage taxonomy has changed: In 2014, within the order *Caudovirales* there were three families, six subfamilies, 72 genera and 484 species. There are now 10 families, 35 subfamilies, 672 genera and 1976 species. The pace and volume of taxonomic changes over the last years may have introduced a period of confusion in the scientific literature. This transition period has been unavoidable but will create a more sustainable clarity in the long term. We

must do our best to guide the community through this transition period towards ‘stable ground’, which will in turn enable textbooks, education tools, databases etc. to follow through.

I would like to thank specifically four individuals: Hans Ackermann (aka Dr. Phage Electron Microscopy/ Dr. Comma), who in spite of misgivings about the direction that we were taking, was always supportive. Whenever I look at an electron micrograph of a phage, I ask myself “What would Hans think of this?” He is greatly missed. Jens Kuhn, who taught me what a taxon is, and who was always ready to support us and write long explanatory emails on the finer aspects of taxonomy, in spite of an extremely busy schedule. Igor Tolstoy from NCBI, who appears to operate without sleep and who sends me queries on deposited phage genomes which have enhanced our taxonomy. Evelien Adriaenssens, who became involved in phage taxonomy with me in 2012, and without whom we would have been far less productive. I know that our subcommittee will be in excellent hands when she takes over its leadership this year.

Future directions

In 2019, it was proposed to the ICTV Executive Committee that the Bacterial and Archaeal Viruses Subcommittee be split. This would result in the formation of the Archaeal Viruses Subcommittee and the Bacterial Viruses Subcommittee.

There is a movement within the ICTV towards a binomial system of viral nomenclature [28]. We therefore request that the phage community move away from giving their isolates names such as “*Escherichia* phage XYZ2020” and instead use a format similar to “*Escherichia* phage Fuddleduddle.” Using words (sensical or not) instead of an alphanumeric will facilitate the expected move to a Linnaean-like system for species names (i.e., if “*Escherichia* phage Fuddleduddle” is a member of the genus “*Twaddlevirus*”, then the new species for “*Escherichia* phage Fuddleduddle” could be called “*Twaddlevirus fuddleduddle*” (with apologies to Pierre Elliot Trudeau). Such a species name, I (AMK) believe, would be more acceptable to the bacteriophage community than the latinized “*Twaddlevirus fuddleduddlensis*”. This naming scheme also has the added benefit that it retains name recognition and better describes the isolate. In GenBank there are currently 426 complete genome sequences for “*Escherichia* phages”, which belong to numerous different families and genera; therefore, the prefix “*Escherichia virus*” has little additional utility in viral taxonomy. Furthermore, in the case of most virome-derived sequences, we do not know the host bacterium; therefore, the current-species naming convention (*host name + virus + species name*) will not work.

We have reached a stage where we need to see fundamental changes in the way Taxonomic Proposals are prepared and assessed. Every year, 400–600 complete phage genome sequences are deposited in the GenBank database, and it has become impossible to rapidly assess potentially new taxa, particularly if they contain numerous representatives. In simple terms, our current way to doing things is too complex and time-consuming. We need to take advantage of the bioinformatically competent ICTV members to develop and implement automated procedures for identification of virus isolates and their easy classification into species, genera, etc. and to generate appropriate documentation. We have recently seen the development of tools such as Pairwise Sequence Comparison (PASC [29]), Sequence Demarcation Tool (SDT [30]), Machine Learning with Digital Signal Processing (ML-DSP [31]), VICTOR [32], ClassiPhage [33], GRAViTy [17], ViPTree [34], vConTACT [6, 7], and the concatenated protein tree method of Low et al. [35], as well as the novel NCBI ORF finder-BLASTp-SymBets; [36] and BLASTN algorithms developed by Igor Tolstoy and Mathew Lueder, respectively. Until such systems are accepted by the ICTV and robust criteria are established to delineate families and subfamilies, phage taxonomy will always lag significantly behind submissions to NCBI, and our committees will not be able to taxonomically address the classification of prophages and metagenomically derived sequences. Environmental virome studies have indicated the existence of huge numbers of diverse and currently unclassified phages, whether it be in oceans, soils or the human gut [37–39]. The case is well illustrated by the families *Microviridae* and *Leviviridae*, which have received relatively little attention from the ICTV (i.e., two subfamilies, six genera and 21 species versus two genera and four species, respectively). Environmental studies have indicated that the family *Microviridae* contains at least five unclassified subfamilies [40–47]. Recently, Callanan et al. [48] predicted 331 species and 247 genera among the single-stranded RNA phages, potentially leading to a significant expansion of the family *Leviviridae*.

Until such a system is in place, we propose to work closely with NCBI to identify new phage species, genera, and subfamilies and to immediately provide names to NCBI. At the annual EC meeting, complete TaxoProps for the newly identified taxa would be presented to the ICTV.

Acknowledgements The committee would like to thank Dr. Graham Hatfull (University of Pittsburgh) for permitting us to use Actinobacteriophage Database electron micrographs in taxonomy proposals. We would also like to thank Professor Colin Hill (University College Cork, Ireland) for letting us see the preprint of their paper on the diversity of ssRNA phages ahead of its publication.

Funding The views and conclusions contained in this document are those of the authors and should not be interpreted as necessarily representing the official policies of their organizations. E.M.A gratefully acknowledges the support of the UK Biotechnology and Biological Sciences Research Council (BBSRC). This research was funded by the BBSRC Institute Strategic Programme Gut Microbes and Health BB/R012490/1 and its constituent project(s) BBS/E/F/000PR10353. B.E.D. was supported by the Netherlands Organization for Scientific Research (NWO), Vidi Grant 864.14.004. R.A.E was supported by Grant MCB-1330800 from the National Science Foundation. J.R.B. was supported by the Intramural Research Program of the National Institutes of Health, National Library of Medicine. R.L. is a member of the PhageBiotics Research Community, supported by FWO Vlaanderen. M.M.P. was supported by the Academy of Finland (272507). H.M.O. was supported by University of Helsinki funding for Instruct-F1 research infrastructure.

Compliance with ethical standards

Conflict of interest The authors declare that they have no conflict of interest.

Ethical approval The authors did not perform any studies with human participants or animals for this article.

References

1. Adriaenssens EM, Wittmann J, Kuhn JH, Turner D, Sullivan MB, Dutilh BE, Jang HB, van Zyl LJ, Klumpp J, Lobočka M, Moreno Switt AI, Rumnieks J, Edwards RA, Uchiyama J, Alfenas-Zerbini P, Petty NK, Kropinski AM, Barylski J, Gillis A, Clokie MRC, Prangishvili D, Lavigne R, Aziz RK, Duffy S, Krupovic M, Poranen MM, Knezevic P, Enault F, Tong Y, Oksanen HM, Rodney Brister J (2018) Taxonomy of prokaryotic viruses: 2017 update from the ICTV Bacterial and Archaeal Viruses Subcommittee. *Arch Virol* 163:1125–1129
2. Mochizuki T, Krupovic M, Pehau-Arnauudet G, Sako Y, Forterre P, Prangishvili D (2012) Archaeal virus with exceptional virion architecture and the largest single-stranded DNA genome. *Proc Natl Acad Sci USA* 109:13386–13391
3. Lavigne R, Seto D, Mahadevan P, Ackermann H-W, Kropinski AM (2008) Unifying classical and molecular taxonomic classification: analysis of the *Podoviridae* using BLASTP-based tools. *Res Microbiol* 159:406–414
4. Jamalludeen N, Johnson RP, Friendship R, Kropinski AM, Lingohr EJ, Gyles CL (2007) Isolation and characterization of nine bacteriophages that lyse O149 enterotoxigenic *Escherichia coli*. *Vet Microbiol* 124:47–57
5. Jamalludeen N, Kropinski AM, Johnson RP, Lingohr E, Harel J, Gyles CL (2008) Complete genomic sequence of bacteriophage phiEcoM-GJ1, a novel phage that has myovirus morphology and a podovirus-like RNA polymerase. *Appl Environ Microbiol* 74:516–525
6. Jang HB, Bolduc B, Zablocki O, Kuhn JH, Roux S, Adriaenssens EM, Brister JR, Kropinski AM, Krupovic M, Lavigne R, Turner D, Sullivan MB (2019) Taxonomic assignment of uncultivated prokaryotic virus genomes is enabled by gene-sharing networks. *Nat Biotechnol* 37:632–639
7. Bolduc B, Jang HB, Doucier G, You ZQ, Roux S, Sullivan MB (2017) vConTACT: an iVirus tool to classify double-stranded DNA viruses that infect Archaea and Bacteria. *PeerJ* 5:e3243

8. Krupovic M, Dutilh BE, Adriaenssens EM, Wittmann J, Vogensen FK, Sullivan MB, Rumnieks J, Prangishvili D, Lavigne R, Kropinski AM, Klumpp J, Gillis A, Enault F, Edwards RA, Duffy S, Clokie MR, Barylski J, Ackermann HW, Kuhn JH (2016) Taxonomy of prokaryotic viruses: update from the ICTV bacterial and archaeal viruses subcommittee. *Arch Virol* 161:1095–1099
9. Laanto E, Mantynen S, De Colibus L, Marjakangas J, Gillum A, Stuart DI, Ravantti JJ, Huiskonen JT, Sundberg LR (2017) Virus found in a boreal lake links ssDNA and dsDNA viruses. *Proc Natl Acad Sci USA* 114:8378–8383
10. Yutin N, Backstrom D, Ettema TJG, Krupovic M, Koonin EV (2018) Vast diversity of prokaryotic virus genomes encoding double jelly-roll major capsid proteins uncovered by genomic and metagenomic sequence analysis. *Virus J* 15:67
11. Bath C, Cukalac T, Porter K, Dyal-Smith ML (2006) His1 and His2 are distantly related, spindle-shaped haloviruses belonging to the novel virus group, *Salterprovirus*. *Virology* 350:228–239
12. Bath C, Dyal-Smith ML (1998) His1, an archaeal virus of the *Fuselloviridae* family that infects *Haloarcula hispanica*. *J Virol* 72:9392–9395
13. Krupovic M, Quemin ER, Bamford DH, Forterre P, Prangishvili D (2014) Unification of the globally distributed spindle-shaped viruses of the Archaea. *J Virol* 88:2354–2358
14. Pietilä MK, Atanasova NS, Oksanen HM, Bamford DH (2013) Modified coat protein forms the flexible spindle-shaped virion of haloarchaeal virus His1. *Environ Microbiol* 15:1674–1686
15. Barylski J, Enault F, Dutilh BE, Schuller MB, Edwards RA, Gillis A, Klumpp J, Knezevic P, Krupovic M, Kuhn JH, Lavigne R, Oksanen HM, Sullivan MB, Jang HB, Simmonds P, Aiewsakun P, Wittmann J, Tolstoy I, Brister JR, Kropinski AM, Adriaenssens EM (2020) Analysis of spounaviruses as a case study for the overdue reclassification of tailed phages. *Syst Biol* 69:110–123
16. Klumpp J, Lavigne R, Loessner MJ, Ackermann HW (2010) The SPO1-related bacteriophages. *Arch Virol* 155:1547–1561
17. Aiewsakun P, Simmonds P (2018) The genomic underpinnings of eukaryotic virus taxonomy: creating a sequence-based framework for family-level virus classification. *Microbiome* 6:38
18. Rohwer F, Edwards R (2002) The phage proteomic tree: a genome-based taxonomy for phage. *J Bacteriol* 184:4529–4535
19. Wang H, Guo Z, Feng H, Chen Y, Chen X, Li Z, Hernandez-Ascencio W, Dai X, Zhang Z, Zheng X, Mora-Lopez M, Fu Y, Zhang C, Zhu P, Huang L (2018) Novel *Sulfolobus* virus with an exceptional capsid architecture. *J Virol* 92:e01717–e01727
20. Kim JG, Kim SJ, Cvirkaite-Krupovic V, Yu WJ, Gwak JH, Lopez-Perez M, Rodriguez-Valera F, Krupovic M, Cho JC, Rhee SK (2019) Spindle-shaped viruses infect marine ammonia-oxidizing thaumarchaea. *Proc Natl Acad Sci USA* 116:15645–15650
21. Prangishvili D, Krupovic M, ICTV Report Consortium (2018) ICTV virus taxonomy profile: *Ampullaviridae*. *J Gen Virol* 99:288–289
22. Prangishvili D, Krupovic M, ICTV Report Consortium (2018) ICTV virus taxonomy profile: *Bicaudaviridae*. *J Gen Virol* 99:864–865
23. Prangishvili D, Krupovic M, ICTV Report Consortium (2018) ICTV Virus Taxonomy Profile: *Globuloviridae*. *J Gen Virol* 99:1357–1358
24. Prangishvili D, Mochizuki T, Krupovic M, ICTV Report Consortium (2018) ICTV virus taxonomy profile: *Guttaviridae*. *J Gen Virol* 99:290–291
25. Prangishvili D, Mochizuki T, Krupovic M, ICTV Report Consortium (2020) ICTV virus taxonomy profile: *Spiraviridae*. *J Gen Virol*. <https://doi.org/10.1099/jgv.0.001385>
26. Prangishvili D, Rensen E, Mochizuki T, Krupovic M, ICTV Report Consortium (2019) ICTV virus taxonomy profile: *Tristromaviridae*. *J Gen Virol* 100:135–136
27. Krupovic M, ICTV Report Consortium (2018) ICTV virus taxonomy profile: *Plasmaviridae*. *J Gen Virol* 99:617–618
28. Siddell SG, Walker PJ, Lefkowitz EJ, Mushegian AR, Dutilh BE, Harrach B, Harrison RL, Junglen S, Knowles NJ, Kropinski AM, Krupovic M, Kuhn JH, Nibert ML, Rubino L, Sabanadzovic S, Simmonds P, Varsani A, Zerbini FM, Davison AJ (2019) Binomial nomenclature for virus species: a consultation. *Arch Virol*. <https://doi.org/10.1007/s00705-019-04477-6>
29. Bao Y, Amarasinghe GK, Basler CF, Bavari S, Bukreyev A, Chandran K, Dolnik O, Dye JM, Ebihara H, Formenty P, Hewson R, Kobinger GP, Leroy EM, Muhlberger E, Netesov SV, Patterson JL, Paweska JT, Smither SJ, Takada A, Towner JS, Volchkov VE, Wahl-Jensen V, Kuhn JH (2017) Implementation of objective PASC-derived taxon demarcation criteria for official classification of filoviruses. *Viruses* 9:106
30. Muhire BM, Varsani A, Martin DP (2014) SDT: a virus classification tool based on pairwise sequence alignment and identity calculation. *PLoS One* 9:e108277
31. Randhawa GS, Hill KA, Kari L (2019) ML-DSP: machine learning with digital signal processing for ultrafast, accurate, and scalable genome classification at all taxonomic levels. *BMC Genom* 20:267
32. Meier-Kolthoff JP, Goker M (2017) VICTOR: genome-based phylogeny and classification of prokaryotic viruses. *Bioinformatics* 33:3396–3404
33. Chibani CM, Farr A, Klama S, Dietrich S, Liesegang H (2019) Classifying the unclassified: a phage classification method. *Viruses* 11:195
34. Nishimura Y, Yoshida T, Kuronishi M, Uehara H, Ogata H, Goto S (2017) ViPTree: the viral proteomic tree server. *Bioinformatics* 33:2379–2380
35. Low SJ, Dzunkova M, Chaumeil PA, Parks DH, Hugenholtz P (2019) Evaluation of a concatenated protein phylogeny for classification of tailed double-stranded DNA viruses belonging to the order *Caudovirales*. *Nat Microbiol* 4:1306–1315
36. Kristensen DM, Kannan L, Coleman MK, Wolf YI, Sorokin A, Koonin EV, Mushegian A (2010) A low-polynomial algorithm for assembling clusters of orthologous groups from intergenomic symmetric best matches. *Bioinformatics* 26:1481–1487
37. Paez-Espino D, Eloe-Fadrosh EA, Pavlopoulos GA, Thomas AD, Hunteemann M, Mikhailova N, Rubin E, Ivanova NN, Kyrpidis NC (2016) Uncovering Earth's virome. *Nature* 536:425–430
38. Roux S, Brum JR, Dutilh BE, Sunagawa S, Duhaime MB, Loy A, Poulos BT, Solonenko N, Lara E, Poulain J, Pesant S, Kandels-Lewis S, Dimier C, Picheral M, Searson S, Cruaud C, Alberti A, Duarte CM, Gasol JM, Vaque D, Bork P, Acinas SG, Wincker P, Sullivan MB (2016) Ecogenomics and potential biogeochemical impacts of globally abundant ocean viruses. *Nature* 537:689–693
39. Shkoporov AN, Clooney AG, Sutton TDS, Ryan FJ, Daly KM, Nolan JA, McDonnell SA, Khokhlova EV, Draper LA, Forde A, Guerin E, Velayudhan V, Ross RP, Hill C (2019) The human gut virome is highly diverse, stable, and individual specific. *Cell Host Microbe* 26:527–541 (e525)
40. Callanan J, Stockdale SR, Shkoporov A, Draper LA, Ross RP, Hill C (2018) RNA phage biology in a metagenomic era. *Viruses* 10:386
41. Creasy A, Rosario K, Leigh BA, Dishaw LJ, Breitbart M (2018) Unprecedented Diversity of ssDNA phages from the family *Microviridae* detected within the gut of a protochordate model organism (*Ciona robusta*). *Viruses* 10:404
42. Gong Z, Liang Y, Wang M, Jiang Y, Yang Q, Xia J, Zhou X, You S, Gao C, Wang J, He J, Shao H, McMinn A (2018) Viral diversity

- and its relationship with environmental factors at the surface and deep sea of Prydz Bay, Antarctica. *Front Microbiol* 9:2981
43. Krishnamurthy SR, Janowski AB, Zhao G, Barouch D, Wang D (2016) Hyperexpansion of RNA bacteriophage diversity. *PLoS Biol* 14:e1002409
 44. Quaiser A, Dufresne A, Ballaud F, Roux S, Zivanovic Y, Colombet J, Sime-Ngando T, Francez AJ (2015) Diversity and comparative genomics of *Microviridae* in Sphagnum-dominated peatlands. *Front Microbiol* 6:375
 45. Sommers P, Fontenele RS, Kringen T, Kraberger S, Porazinska DL, Darcy JL, Schmidt SK, Varsani A (2019) Single-stranded DNA viruses in antarctic cryoconite holes. *Viruses* 11:1022
 46. Wang H, Ling Y, Shan T, Yang S, Xu H, Deng X, Delwart E, Zhang W (2019) Gut virome of mammals and birds reveals high genetic diversity of the family *Microviridae*. *Virus Evol* 5:vez013
 47. Zhong X, Guidoni B, Jacas L, Jacquet S (2015) Structure and diversity of ssDNA *Microviridae* viruses in two peri-alpine lakes (Annecy and Bourget, France). *Res Microbiol* 166:644–654
 48. Callanan J, Stockdale SR, Shkoporov A, Draper LA, Ross RP (2020) Hill C (2020) Expansion of known ssRNA phage genomes: from tens to over a thousand. *Sci Adv* 6(6):eaay5981. <https://doi.org/10.1126/sciadv.aay5981>

Publisher's Note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Affiliations

Evelien M. Adriaenssens¹ · Matthew B. Sullivan² · Petar Knezevic³ · Leonardo J. van Zyl⁴ · B. L. Sarkar⁵ · Bas E. Dutilh^{6,7} · Poliane Alfenas-Zerbini⁸ · Małgorzata Łobocka⁹ · Yigang Tong¹⁰ · James Rodney Brister¹¹ · Andrea I. Moreno Switt¹² · Jochen Klumpp¹³ · Ramy Karam Aziz¹⁴ · Jakub Barylski¹⁵ · Jumpei Uchiyama¹⁶ · Rob A. Edwards^{17,18} · Andrew M. Kropinski^{19,20} · Nicola K. Petty²¹ · Martha R. J. Clokie²² · Alla I. Kushkina²³ · Vera V. Morozova²⁴ · Siobain Duffy²⁵ · Annika Gillis²⁶ · Janis Rumnieks²⁷ · İpek Kurtböke²⁸ · Nina Chanishvili²⁹ · Lawrence Goodridge¹⁹ · Johannes Wittmann³⁰ · Rob Lavigne³¹ · Ho Bin Jang³² · David Prangishvili^{33,34} · Francois Enault³⁵ · Dann Turner³⁶ · Minna M. Poranen³⁷ · Hanna M. Oksanen³⁷ · Mart Krupovic³³

✉ Andrew M. Kropinski
Phage.Canada@gmail.com

¹ Quadram Institute Bioscience, Norwich Research Park, Norwich, Norfolk NR4 7UQ, UK

² Department of Microbiology and Civil, Environmental and Geodetic Engineering, The Ohio State University, Columbus, OH 43210, USA

³ Department of Biology and Ecology, Faculty of Sciences, University of Novi Sad, Novi Sad, Serbia

⁴ Department of Biotechnology, Institute for Microbial Biotechnology and Metagenomics (IMBM), University of the Western Cape, Bellville, Cape Town 7535, South Africa

⁵ ICMR-National Institute of Cholera and Enteric Diseases, P-33, C.I.T. Road, Scheme XM, Belegata, Kolkata 700010, India

⁶ Theoretical Biology and Bioinformatics, Utrecht University, Utrecht, The Netherlands

⁷ Centre for Molecular and Biomolecular Informatics, Radboud University Medical Centre, Nijmegen, The Netherlands

⁸ Laboratory of Industrial Microbiology, Instituto de Biotecnologia Aplicada à Agropecuária, Universidade Federal de Viçosa, Viçosa, Minas Gerais, Brazil

⁹ Department of Microbial Biochemistry, Institute of Biochemistry and Biophysics of the Polish Academy of Sciences, 02-106 Warsaw, Poland

¹⁰ College of Life Science and Technology, Beijing University of Chemical Technology, Beijing 100029, China

¹¹ National Center for Biotechnology Information, National Library of Medicine, National Institutes of Health, Bethesda, MD 20894, USA

¹² Faculty of Life Sciences, Universidad Andres Bello, Republica 440, 8370146 Santiago, Chile

¹³ Institute of Food, Nutrition and Health, ETH Zurich, 8092 Zurich, Switzerland

¹⁴ Department of Microbiology and Immunology, Faculty of Pharmacy, Cairo University, Qasr El-Ainy St, Cairo 11562, Egypt

¹⁵ Department of Molecular Virology, Institute of Experimental Biology, Adam Mickiewicz University, Poznan, Poland

¹⁶ School of Veterinary Medicine, Azabu University, Fuchinobe 1-17-71, Chuo-ku, Sagamihara, Kanagawa 252-0206, Japan

¹⁷ Department of Computer Science, San Diego State University, San Diego, CA 92182, USA

¹⁸ Department of Biology, San Diego State University, San Diego, CA 92182, USA

¹⁹ Food Science Department, University of Guelph, 50 Stone Road East, Guelph, ON N1G 2W1, Canada

²⁰ Department of Pathobiology, Ontario Veterinary College, University of Guelph, 50 Stone Road E, Guelph, ON N1G 2W1, Canada

²¹ The itree Institute, University of Technology Sydney, Sydney, NSW 2007, Australia

²² Department of Infection, Immunity and Inflammation, University of Leicester, Leicester LE1 9HN, UK

²³ Department of Bacteriophage Molecular Genetics, D.K. Zabolotny Institute of Microbiology and Virology,

- the NAS of Ukraine, 154 Acad.Zabolotny str, Kyiv 03143, Ukraine
- ²⁴ Institute of Chemical Biology and Fundamental Medicine (RAS), Novosibirsk, Russia
- ²⁵ Department of Ecology, Evolution and Natural Resources, Rutgers University, New Brunswick, NJ 08901, USA
- ²⁶ Section of Molecular Microbiology and Medical Research Council Centre for Molecular Bacteriology and Infection, Imperial College London, London SW72AZ, UK
- ²⁷ Latvian Biomedical Research and Study Center, Ratsupites 1, Riga 1067, Latvia
- ²⁸ Genecology Research Centre, School of Science and Engineering, University of the Sunshine Coast, Maroochydore DC, QLD 4558, Australia
- ²⁹ G. Eliava Institute of Bacteriophages, Microbiology and Virology, 3 Gotua Street, 0160 Tbilisi, Georgia
- ³⁰ Leibniz Institute DSMZ-German Collection of Microorganisms and Cell Cultures GmbH, 38124 Brunswick, Germany
- ³¹ Laboratory of Gene Technology, Department of Biosystems, KU Leuven, Kasteelpark Arenberg 21 box 2462, 3001 Leuven, Belgium
- ³² Department of Microbiology, The Ohio State University, Columbus, OH 43210, USA
- ³³ Archaeal Virology Unit, Department of Microbiology, Institut Pasteur, 75015 Paris, France
- ³⁴ Ivane Javakhishvili Tbilisi State University, 0179 Tbilisi, Georgia
- ³⁵ Université Clermont Auvergne, CNRS, LMGE, 63000 Clermont-Ferrand, France
- ³⁶ Faculty of Health and Applied Sciences, University of the West of England, Bristol, Frenchay Campus, Bristol BS16 1QY, UK
- ³⁷ Molecular and Integrative Biosciences Research Programme, Faculty of Biological and Environmental Sciences, University of Helsinki, Helsinki, Finland