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# Summary of taxonomy changes ratified by the International Committee on Taxonomy of Viruses (ICTV) from the Bacterial Viruses Subcommittee, 2025

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## Abstract

This article summarises the activities of the International Committee on Taxonomy of Viruses Bacterial Viruses Subcommittee, detailing developments in the classification of bacterial viruses. We provide here an overview of all new, abolished, moved and renamed taxa proposed in 2024, approved by the Executive Committee, and ratified by membership vote in 2025. Through the collective efforts of 74 international contributors of taxonomy proposals in this round, 43 ratified proposals have led to the creation of one new phylum, one class, four orders, 33 families, 14 subfamilies, 194 genera and 995 species. These proposals mark significant progress in refining the taxonomy of bacterial viruses. Key updates include the creation of new orders and families that include existing taxa to better reflect genomic and evolutionary relationships. As sequencing and bioinformatics approaches continue to advance, further expansion and refinements in viral taxonomy can be anticipated in the coming years.

## INTRODUCTION

The taxonomy of bacterial viruses has undergone significant changes in recent years, notably the abolishment of the paraphyletic families *Myoviridae*, *Siphoviridae* and *Podoviridae*, along with the order *Caudovirales* [1, 2]. This reorganization left a number of orphan subfamilies and genera within the class *Caudoviricetes* without a designated family or order. The adoption of genome-based classification, which focuses on reconstructing virus evolutionary histories through nucleotide and amino-acid sequence similarity, phylogenetic analysis of core genes and increasingly, of protein folds and motifs for the demarcation of higher ranks, has enabled the identification of many new monophyletic taxa [3]. Additionally, a number of tools are now available to aid the classification of bacterial viruses across different taxonomic ranks (e.g. [4–7]).

Here, we present summaries of all ratified taxonomic proposals submitted to the Bacterial Viruses Subcommittee of the International Committee on Taxonomy of Viruses (ICTV) in 2024. These proposals were assessed and approved at the ICTV Executive Committee meeting (EC56, July 2024) and ratified by membership vote in 2025. Full versions of the proposals summarized here,

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For numbered affiliations see end of the article.

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Abbreviations: ICTV, International Committee on Taxonomy of Viruses; RdRP, RNA-directed RNA polymerase.

Supplementary files are available with the online version of this article.

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**Table 1.** Summary of taxonomic changes for bacterial viruses, ratified in 2025

Taxonomy change	Species	Genus	Subfamily	Family	Order	Class	Phylum
Abolished	22	1	2	-	-	-	-
New	995	194	14	33	3	1	1
Moved or promoted (and/or renamed)	30	120	12	4	1	-	-
Renamed	101	3	-	-	-	-	-

as well as those from previous years, are available on the ICTV website (<https://ictv.global/files/proposals/approved>). A total of 43 proposals were ratified, leading to the creation of one new phylum, one class, four orders, 33 families, 14 subfamilies, 194 genera and 995 species (Table 1).

Notable highlights include the creation of the phylum *Artimaviricota* for a group of segmented RNA viruses predicted to infect thermophilic bacteria [8]. The Lak megaphages [9, 10] were assigned to the new order *Grandevirales*, while the family *Autographiviridae* was elevated to order (*Autographivirales*), containing four newly defined families. The new order *Pantenvirales* encompasses the families *Straboviridae*, *Kyanoviridae*, and *Ackermannviridae*. Importantly, the creation of 30 new families continues the process of addressing floating taxa within the class *Caudoviricetes*. This class now contains 31 families assigned to 10 orders, with 70 families remaining unassigned to any order (Fig. 1).

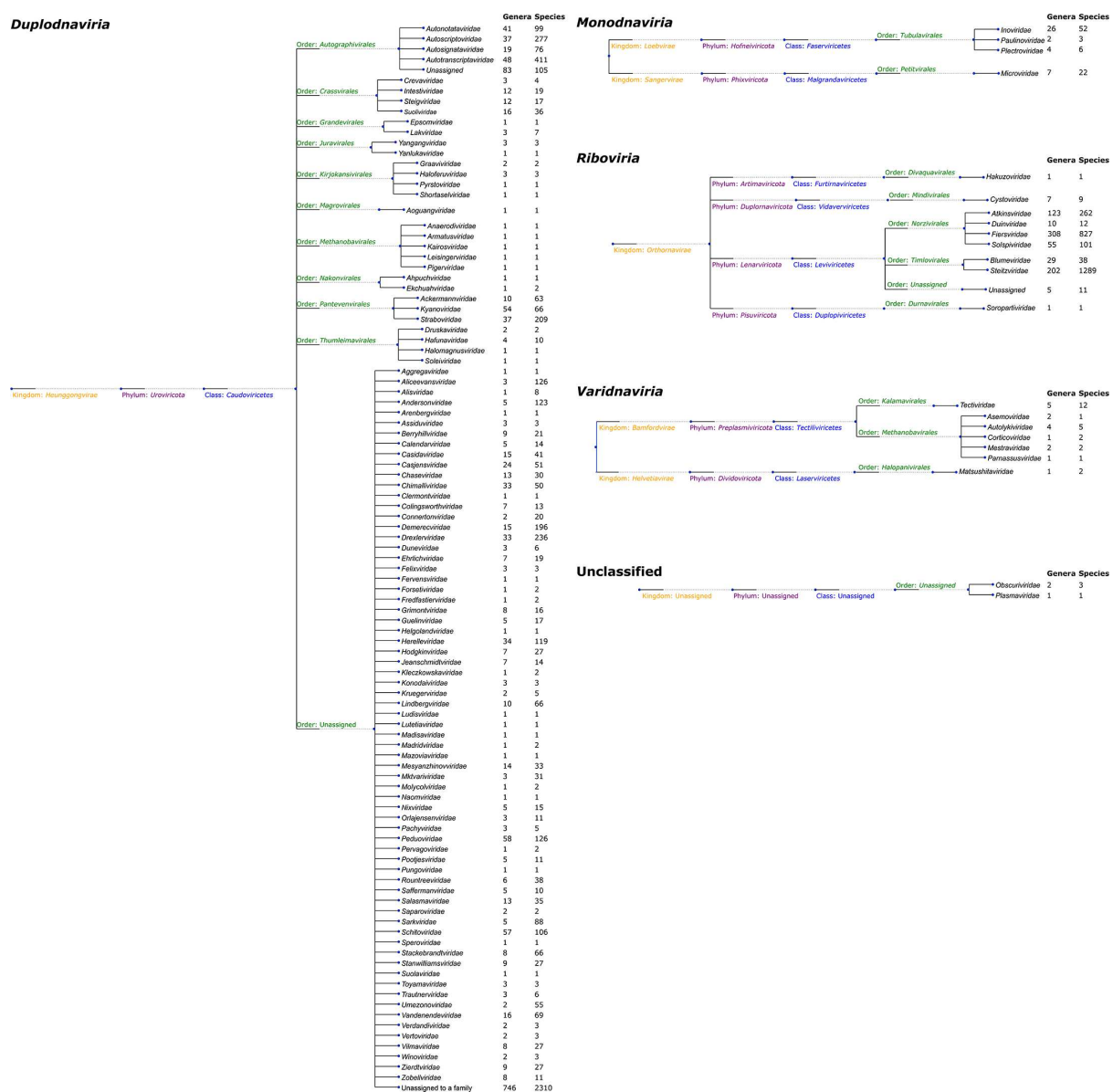
The Bacterial Viruses Subcommittee has also seen growing engagement from the wider research community, as evidenced by an increased number of proposals originating from outside the Subcommittee and its Study Groups. This is encouraged, and we invite researchers to contact the Subcommittee and relevant Study Groups for guidance ([https://ictv.global/study-groups?webform\\_submission\\_value\\_1=7](https://ictv.global/study-groups?webform_submission_value_1=7)). A file including all the Tables of taxonomic changes below is available as a supplementary file to this article.

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**Fig. 1.** Cladograms depicting the taxonomy of bacterial viruses for the realms *Duplodnaviria*, *Monodnaviria*, *Riboviria*, *Varidnaviria* and the families *Obscuriviridae* and *Plasmaviridae* which have no realm assigned. The columns adjacent to the leaf labels provide counts of the numbers of genera and species by family.

**2024.001B.Alisviridae\_Ludisviridae\_Nixviridae\_3nf\_7ng\_24ns****Title:** Create three new families (*Alisviridae*, *Ludisviridae* and *Nixviridae*) with seven new genera and 24 new species**Authors:** Matrishin CB, Kauffman KM (kmkauffm@buffalo.edu)**Summary****Taxonomic rank(s) affected**Realm *Duplodnaviria*, kingdom *Heunggongvirae*, phylum *Uroviricota*, class *Caudoviricetes*.**Description of current taxonomy**

The viruses classified in this proposal do not have a current taxonomic assignment.

**Proposed taxonomic change(s)**Creation of three new families (*Alisviridae*, *Ludisviridae* and *Nixviridae*) with seven new genera (*Honmavirus*, *Ludivirus*, *Dewhirstvirus*, *Nixvirus*, *Haasevirus*, *Excelsiorvirus* and *Schifferlevirus*) and 24 new species.**Justification**

A comprehensive analysis of publicly available NCBI *Porphyromonas gingivalis* genomes revealed three new families of viruses, containing seven new genera and 24 new species. This discovery, using a rigorous, complementary bioinformatic approach, revealed what we believe to be precise nucleotide start and end points of the prophage genomes within bacterial contigs. These novel prophages represent the first systematically described phages of *P. gingivalis*. This work, including the proposed taxonomic classifications and figures shown in this proposal, is described in Matrishin *et al.* [11]. Phage genomes are available on NCBI within BioProject PRJNA874424.

**Submitted:** 29 May 24**Table 2.** *Alisviridae*, 34 new taxa\*

Operation	Rank	New taxon name	Virus name	Exemplar
New taxon	Family	<i>Alisviridae</i>		
New taxon	Genus	<i>Honmavirus</i>		
New taxon	Species	<i>Honmavirus pging00B</i>	Porphyromonas phage phage006a_EM3	BK068089
New taxon	Species	<i>Honmavirus pging00C</i>	Porphyromonas phage phage007a_Bg4	PP754929
New taxon	Species	<i>Honmavirus pging00D</i>	Porphyromonas phage phage008a_KCOM2795	BK068090
New taxon	Species	<i>Honmavirus pging00E</i>	Porphyromonas phage phage010a_HG1691old	PP754930
New taxon	Species	<i>Honmavirus pging00F</i>	Porphyromonas phage phage011a_WW2952	BK068092
New taxon	Species	<i>Honmavirus pging00G</i>	Porphyromonas phage phage012a_381OKJP	BK068093
New taxon	Species	<i>Honmavirus pging00H</i>	Porphyromonas phage phage013a_WW2885	BK068094
New taxon	Species	<i>Honmavirus pging00I</i>	Porphyromonas phage phage014a_Kyudai4	BK068095
New taxon	Family	<i>Ludisviridae</i>		
New taxon	Genus	<i>Ludivirus</i>		
New taxon	Species	<i>Ludivirus pging00A</i>	Porphyromonas phage phage005b_ATCC49417	PP754928
New taxon	Family	<i>Nixviridae</i>		
New taxon	Genus	<i>Dewhirstvirus</i>		
New taxon	Species	<i>Dewhirstvirus pging00J</i>	Porphyromonas phage phage016a_WW2866	BK068097
New taxon	Species	<i>Dewhirstvirus pging00K</i>	Porphyromonas phage phage017a_JCVISC001	BK068098
New taxon	Species	<i>Dewhirstvirus pging00L</i>	Porphyromonas phage phage018a_AFR5B1	BK068099
New taxon	Species	<i>Dewhirstvirus pging00M</i>	Porphyromonas phage phage019b_ATCC49417	PP754931
New taxon	Genus	<i>Nixvirus</i>		
New taxon	Species	<i>Nixvirus pging00X</i>	Porphyromonas phage phage032a_KCOM2801	BK068113
New taxon	Genus	<i>Haasevirus</i>		

Continued

**Table 2.** Continued

Operation	Rank	New taxon name	Virus name	Exemplar
New taxon	Species	<i>Haasevirus pging00R</i>	Porphyromonas phage phage025a_SJD11	BK068106
New taxon	Species	<i>Haasevirus pging00T</i>	Porphyromonas phage phage027a_F0568	BK068108
New taxon	Species	<i>Haasevirus pging00U</i>	Porphyromonas phage phage028a_KCOM2799	BK068109
New taxon	Species	<i>Haasevirus pging00V</i>	Porphyromonas phage phage029a_Kyudai3	BK068110
New taxon	Species	<i>Haasevirus pging00W</i>	Porphyromonas phage phage030a_KCOM2803	BK068111
New taxon	Genus	<i>Excelsiorvirus</i>		
New taxon	Species	<i>Excelsiorvirus pging00S</i>	Porphyromonas phage phage026a_KCOM2802	BK068107
New taxon	Genus	<i>Schifferlevirus</i>		
New taxon	Species	<i>Schifferlevirus pging00N</i>	Porphyromonas phage phage020a_SJD2	BK068101
New taxon	Species	<i>Schifferlevirus pging00O</i>	Porphyromonas phage phage022a_WW2931	BK068103
New taxon	Species	<i>Schifferlevirus pging00P</i>	Porphyromonas phage phage023a_KCOM2797	BK068104
New taxon	Species	<i>Schifferlevirus pging00Q</i>	Porphyromonas phage phage024a_F0570	BK068105

\*Source/full text: [https://ictv.global/ictv/proposals/2024.001B.Alisviridae\\_Ludisviridae\\_Nixviridae\\_3nf\\_7ng\\_24ns.zip](https://ictv.global/ictv/proposals/2024.001B.Alisviridae_Ludisviridae_Nixviridae_3nf_7ng_24ns.zip).

## 2024.002B.Andersonviridae\_1nf\_2ng\_98ns

**Title:** Create a new family, *Andersonviridae* for the FelixO1-like phages (class: *Caudoviricetes*)

**Authors:** Moraru C, Tolstoy I, Kropinski AM (Phage.Canada@gmail.com)

### Summary

#### Taxonomic rank(s) affected

Realm *Duplodnaviria*, kingdom *Heunggongvirae*, phylum *Uroviricota*, class *Caudoviricetes*.

#### Description of current taxonomy

At present the following taxa exist as floating genera in the class *Caudoviricetes*: *Felixounavirus*, *Kolesnikvirus*, *Suspivirus* and *Mooglevirus*.

#### Proposed taxonomic change(s)

We propose the creation of one new family, *Andersonviridae*.

To update the genus *Felixounavirus* with 80 new species.

To update the genus *Mooglevirus* with 11 new species.

To add one new species to the genus *Kolesnikvirus*.

To create a new genus *Daniellevirus* with two species.

To create a new genus *Arnovirus* with three species.

#### Justification

We investigated the evolutionary relationships of 123 bacteriophages. Analysis of conserved genes revealed that these phages form a deeply branching monophyletic clade with a distance commensurate with the creation of a new family.

**Submitted:** 01 June 24; **Revised:** 30 September 24

**Table 3.** *Andersonviridae*, 100 new taxa\*. Table too large, see supplementary information sheet supp\_info\_tab\_3

**Table 4.** *Andersonviridae*, 1 move taxa\*

Operation	Rank	Taxon name	New parent taxon	Old parent taxon
Move taxon	Subfamily	<i>Ounavirinae</i>	<i>Andersonviridae</i>	<i>Caudoviricetes</i>

**Table 5.** Andersonviridae, 2 move; rename taxa\*

Operation	Rank	New taxon name	New parent taxon	Old taxon name
Move; rename taxon	Species	<i>Mooglevirus susp1</i>	<i>Mooglevirus</i>	<i>Suspivirus SUSP1</i>
Move; rename taxon	Species	<i>Mooglevirus susp2</i>	<i>Mooglevirus</i>	<i>Suspivirus SUSP2</i>

**Table 6.** Andersonviridae, 1 abolish taxon\*

Operation	Rank	Abolished taxon name
Abolish taxon	Genus	<i>Suspivirus</i>

\*Source/full text: [https://ictv.global/ictv/proposals/2024.002B.Andersonviridae\\_1nf\\_2ng\\_98ns.zip](https://ictv.global/ictv/proposals/2024.002B.Andersonviridae_1nf_2ng_98ns.zip).

## 2024.003B.Berryhillviridae\_1nf\_7ng\_3mg\_10ns

**Title:** Create a new family, *Berryhillviridae*, for a group of lytic *Arthrobacter* phages (class: *Caudoviricetes*)

**Authors:** Kurtböke I, Moraru C, Tolstoy I, Kropinski AM (Phage.Canada@gmail.com)

### Summary

#### Taxonomic rank(s) affected

Realm *Duplodnaviria*, kingdom *Heunggongvirae*, phylum *Uroviricota*, class *Caudoviricetes*.

#### Description of current taxonomy

At present the following taxa exist as floating genera in the class *Caudoviricetes*: genera *Marthavirus*, *Vibakivirus*, *Jawnskivivirus* and *Ayohtrevirus*.

#### Proposed taxonomic change(s)

We propose the creation of a new family, *Berryhillviridae*, including the existing genera *Marthavirus*, *Vibakivirus*, and *Ayohtrevirus* in addition to six new genera, *Jinkiesvirus*, *Jawnskivivirus*, *Lilmacvirus*, *Altadenavirus*, *Eastwestvirus* and *Sicariusvirus*.

#### Justification

We investigated the evolutionary relationships of 21 bacteriophages. Analysis of conserved genes and tBLASTx distances revealed that these phages form a deeply branching clade at a distance commensurate with the creation of a new family.

**Submitted:** 25 May 24

**Table 7.** Berryhillviridae, 18 new taxa\*

Operation	Rank	New taxon name	Virus name	Exemplar
New taxon	Family	<i>Berryhillviridae</i>		
New taxon	Genus	<i>Jawnskivivirus</i>		
New taxon	Species	<i>Jawnskivivirus king2</i>	Arthrobacter phage King2	MT776811
New taxon	Genus	<i>Jinkiesvirus</i>		
New taxon	Species	<i>Jinkiesvirus jinkies</i>	Arthrobacter phage Jinkies	MT498043
New taxon	Genus	<i>Lilmacvirus</i>		
New taxon	Species	<i>Lilmacvirus lilmac1015</i>	Arthrobacter phage Lilmac1015	OL742560
New taxon	Species	<i>Lilmacvirus bolt007</i>	Arthrobacter phage Bolt007	OP985600
New taxon	Species	<i>Lilmacvirus prairie</i>	Arthrobacter phage Prairie	MW601223
New taxon	Species	<i>Lilmacvirus klevey</i>	Arthrobacter phage Klevey	MZ747522
New taxon	Genus	<i>Altadenavirus</i>		
New taxon	Species	<i>Altadenavirus altadena</i>	Arthrobacter phage Altadena	OR521058
New taxon	Species	<i>Altadenavirus bumble</i>	Arthrobacter phage Bumble	MT498055
New taxon	Genus	<i>Eastwestvirus</i>		

Continued

**Table 7.** Continued

Operation	Rank	New taxon name	Virus name	Exemplar
New taxon	Species	<i>Eastwestvirus eastwest</i>	Arthrobacter phage EastWest	OK999980
New taxon	Genus	<i>Sicariusvirus</i>		
New taxon	Species	<i>Sicariusvirus sicarius2</i>	Arthrobacter phage Sicarius2	MW862982
New taxon	Species	<i>Sicariusvirus wyborn</i>	Arthrobacter phage Wyborn	OR475274

**Table 8.** Berryhillviridae, 4 move; rename taxa.\*

Operation	Rank	New taxon name	New parent taxon	Old taxon name
Move; rename taxon	Species	<i>Jawnskivirus jawnski</i>	<i>Jawnskivirus</i>	<i>Marthavirus jawnski</i>
Move; rename taxon	Species	<i>Jawnskivirus beans</i>	<i>Jawnskivirus</i>	<i>Marthavirus beans</i>
Move; rename taxon	Species	<i>Jawnskivirus piccoletto</i>	<i>Jawnskivirus</i>	<i>Marthavirus piccoletto</i>
Move; rename taxon	Species	<i>Jawnskivirus brent</i>	<i>Jawnskivirus</i>	<i>Marthavirus brent</i>

**Table 9.** Berryhillviridae, 3 move taxa.\*

Operation	Rank	Taxon name	New parent taxon	Old parent taxon
Move taxon	Genus	<i>Vibakivirus</i>	<i>Berryhillviridae</i>	<i>Caudoviricetes</i>
Move taxon	Genus	<i>Ayohtrevirus</i>	<i>Berryhillviridae</i>	<i>Caudoviricetes</i>
Move taxon	Genus	<i>Marthavirus</i>	<i>Berryhillviridae</i>	<i>Caudoviricetes</i>

\*Source/full text: [https://ictv.global/ictv/proposals/2024.003B.Berryhillviridae\\_1nf\\_7ng\\_3mg\\_10ns.zip](https://ictv.global/ictv/proposals/2024.003B.Berryhillviridae_1nf_7ng_3mg_10ns.zip).

## 2024.005B.Casidaviridae\_1nf\_9ng\_23ns

**Title:** Create a new family, *Casidaviridae*, for a group of *Arthrobacter-Microbacterium* phages (class: *Caudoviricetes*)

**Authors:** Kurtböke I, Moraru C, Tolstoy I, Kropinski AM (Phage.Canada@gmail.com)

### Summary

#### Taxonomic rank(s) affected

Realm *Duplodnaviria*, kingdom *Heunggongvirae*, phylum *Uroviricota*, class *Caudoviricetes*.

#### Description of current taxonomy

At present the following taxa exist as floating genera in the class *Caudoviricetes*: *Zetavirus*, *Baileybluvirus*, *Yangvirus*, *Manhattanvirus* and *Liebevirus*.

#### Proposed taxonomic change(s)

- To create a new genus, *Gardenstatevirus*, with two species
- To create a new genus, *Percivalvirus*, with two species.
- To create a new single species genus *Mabodamacavirus*.
- To create a new genus, *Barnstormervirus* with two species.
- To create a new single species genus *Honkvirus*.
- To create a new single species genus *Cenunavirus*.
- To create a new species in *Baileybluvirus*.
- To create seven new species in the genus *Yangvirus*.
- To create two new species in the genus *Manhattanvirus*.
- To create a new single species genus, *Emotionvirus*.
- To create a new single species genus, *Hilgardvirus*.
- To create a new single species genus, *Swepdovirus*.
- To create one new species in the genus *Liebevirus*.
- To create a new family, *Casidaviridae*.

## Justification

We propose the creation of a new family, *Casidaviridae*, after examination of 21 bacteriophages related to the genera *Zetavirus*, *Baileybluvirus*, *Yangvirus*, *Manhattanvirus* and *Liebevirus* based on nucleotide sequence similarity, tBLASTx distances and core gene phylogeny.

**Submitted:** 20 May 24; **Revised:** 30 September 24

**Table 10.** *Casidaviridae*, 33 new taxa\*

Operation	Rank	New taxon name	Virus name	Exemplar
New taxon	Family	<i>Casidaviridae</i>		
New taxon	Species	<i>Baileybluvirus callinallbarbz</i>	Arthrobacter phage CallinAllBarbz	OR553891
New taxon	Species	<i>Yangvirus janeemi</i>	Arthrobacter phage Janeemi	ON970616
New taxon	Species	<i>Yangvirus tuck</i>	Arthrobacter phage Tuck	OP820474
New taxon	Species	<i>Yangvirus berrie</i>	Arthrobacter phage Berrie	PP208921
New taxon	Species	<i>Yangvirus ascela</i>	Arthrobacter phage Ascela	OQ709218
New taxon	Species	<i>Yangvirus cassia</i>	Arthrobacter phage Cassia	OQ709212
New taxon	Species	<i>Yangvirus tfortroy</i>	Arthrobacter phage TfOrTroy	PP208923
New taxon	Species	<i>Yangvirus nitro</i>	Arthrobacter phage Nitro	OR553895
New taxon	Species	<i>Manhattanvirus vresidence</i>	Arthrobacter phage VResidence	OP434455
New taxon	Species	<i>Manhattanvirus wildwest</i>	Arthrobacter phage Wildwest	OR521060
New taxon	Species	<i>Liebevirus maguco</i>	Arthrobacter phage MaGuCo	OQ709203
New taxon	Genus	<i>Gardenstatevirus</i>		
New taxon	Species	<i>Gardenstatevirus gardenstate</i>	Microbacterium phage GardenState	MT952845
New taxon	Species	<i>Gardenstatevirus iamgroot</i>	Microbacterium phage IAmGroot	MK880124
New taxon	Genus	<i>Percivalvirus</i>		
New taxon	Species	<i>Percivalvirus percival</i>	Microbacterium phage Percival	MH271308
New taxon	Species	<i>Percivalvirus floof</i>	Microbacterium phage Floof	MH271298
New taxon	Genus	<i>Mabodamacavirus</i>		
New taxon	Species	<i>Mabodamacavirus mabodamaca</i>	Microbacterium phage Mabodamaca	OR613467
New taxon	Genus	<i>Barnstormervirus</i>		
New taxon	Species	<i>Barnstormervirus barnstormer</i>	Microbacterium phage Barnstormer	OQ190478
New taxon	Species	<i>Barnstormervirus caron</i>	Microbacterium phage Caron	OQ190481
New taxon	Genus	<i>Honkvirus</i>		
New taxon	Species	<i>Honkvirus honk</i>	Microbacterium phage Honk	MW862981
New taxon	Genus	<i>Cenunavivirus</i>		
New taxon	Species	<i>Cenunavirus Cen1621</i>	Microbacterium phage Cen1621	ON970568
New taxon	Genus	<i>Emotionvirus</i>		
New taxon	Species	<i>Emotionvirus emotion</i>	Arthrobacter phage Emotion	OQ709216
New taxon	Genus	<i>Hilgardvirus</i>		
New taxon	Species	<i>Hilgardvirus vroomvroom</i>	Arthrobacter phage VroomVroom	OQ938592
New taxon	Genus	<i>Swepdovirus</i>		
New taxon	Species	<i>Swepdovirus SWEP2</i>	Arthrobacter phage SWEP2	ON528933

**Table 11.** *Casidaviridae*, 5 move taxa\*

Operation	Rank	Taxon name	New parent taxon	Old parent taxon
Move taxon	Genus	<i>Zetavirus</i>	<i>Casidaviridae</i>	<i>Caudoviricetes</i>
Move taxon	Genus	<i>Baileybluvirus</i>	<i>Casidaviridae</i>	<i>Caudoviricetes</i>
Move taxon	Genus	<i>Yangvirus</i>	<i>Casidaviridae</i>	<i>Azeevirinae</i>
Move taxon	Genus	<i>Manhattanvirus</i>	<i>Casidaviridae</i>	<i>Azeevirinae</i>
Move taxon	Genus	<i>Liebevirus</i>	<i>Casidaviridae</i>	<i>Azeevirinae</i>

**Table 12.** *Casidaviridae*, 1 abolish taxon\*

Operation	Rank	Abolished taxon name
Abolish taxon	Subfamily	<i>Azeevirinae</i>

\*Source/full text: [https://ictv.global/ictv/proposals/2024.005B.Casidaviridae\\_1nf\\_9ng\\_23ns.zip](https://ictv.global/ictv/proposals/2024.005B.Casidaviridae_1nf_9ng_23ns.zip).

## 2024.006B.Cepavirus\_Suseptimavirus\_Uetakevirus\_1ng\_3ns

**Title:** Create a new genus, *Cepavirus*, with one species (*Caudoviricetes*; *Autographiviridae*; *Slopekvirinae*) and new species in the genera *Suseptimavirus* (*Caudoviricetes*; *Gordonclarkvirinae*) and *Uetakevirus* (*Caudoviricetes*).

**Authors:** Pas C, Fieseler L, Briers Y (yves.briers@ugent.be)

### Summary

#### Taxonomic rank(s) affected

Genus and species.

#### Description of current taxonomy

The bacterial viruses in this proposal are currently unclassified.

#### Proposed taxonomic change(s)

Creation of a new genus, *Cepavirus*, within the subfamily *Slopekvirinae*, family *Autographiviridae*.

Assign *Escherichia* phage vB\_EcoP\_PAS7 as a new species in the new genus, *Cepavirus*.

Assign *Escherichia* phage vB\_EcoP\_PAS59 as a new species in the genus *Suseptimavirus*, subfamily *Gordonclarkvirinae*.

Assign *Escherichia* phage vB\_EcoP\_PAS6 as a new species within the genus *Uetakevirus*.

#### Justification

These bacterial viruses fall within current genus and species demarcation criteria for inclusion within existing genera.

**Submitted:** 27 November 23

**Table 13.** *Cepavirus*, 4 new taxa\*

Operation	Rank	New taxon name	Virus name	Exemplar
New taxon	Genus	<i>Cepavirus</i>		
New taxon	Species	<i>Cepavirus PAS7</i>	<i>Escherichia</i> phage vB_EcoP_PAS7	OQ921331
New taxon	Species	<i>Suseptimavirus PAS59</i>	<i>Escherichia</i> phage vB_EcoP_PAS59	OQ921332
New taxon	Species	<i>Uetakevirus PAS61</i>	<i>Escherichia</i> phage vB_EcoP_PAS61	OQ921333

\*Source/full text: [https://ictv.global/ictv/proposals/2024.006B.Cepavirus\\_Suseptimavirus\\_Uetakevirus\\_1ng\\_3ns.zip](https://ictv.global/ictv/proposals/2024.006B.Cepavirus_Suseptimavirus_Uetakevirus_1ng_3ns.zip).

## 2024.007B.Chimalliviridae\_16mg

**Title:** Move newly classified viral genera into *Chimalliviridae* family and fix previous error

**Author:** Prichard A, Pogliano J (jpogliano@ucsd.edu)

### Summary

#### Taxonomic rank(s) affected

We propose to move fifteen genera into the *Chimalliviridae* family and remove one genus from this family.

**Description of current taxonomy**

Last year, we submitted a proposal to create a new viral family called *Chimalliviridae* and re-assigned currently classified viruses into this family. This proposal was accepted, but in the meantime, more viruses that belong in this family have been officially recognized and classified by the ICTV. Since these proposals were submitted in the same year as ours, and there was no existing family that was appropriate for them to be assigned to at the time, these viruses were not assigned to any viral families. However, we believe that these viruses belong to the now-recognized *Chimalliviridae* family.

**Proposed taxonomic change(s)**

Members of the newly created genera *Miamivirus*, *Nimduovirus*, *Meadowvirus*, *Branisovskavirus*, *Ferozepurvirus*, *Chaoshanvirus*, *Ludhianavirus*, *Siatvirus*, *Maaswegvirus*, *Eowynvirus*, *Miltoncavirus*, *Phabiovirus*, *Serwervirus*, *Tepukevirus* and *Pawinskivirus* should be added to the family *Chimalliviridae*. Additionally, the genus *Takahashivirus* should be removed from the family *Chimalliviridae*, as it was included in our proposal last year by mistake and should not belong to this family.

**Justification**

We have redone the phylogenetic analysis used to support the creation of the family *Chimalliviridae* but with the inclusion of several newly classified genera. This shows that these new genera belong in the family *Chimalliviridae*, while *Takahashivirus PBS1*, which we included by mistake, does not.

**Submitted:** 12 June 24

**Table 14.** *Chimalliviridae*, 16 move taxa\*

Operation	Rank	Taxon name	Old parent taxon	New parent taxon
Move taxon	Genus	<i>Takahashivirus</i>	<i>Chimalliviridae</i>	<i>Caudoviricetes</i>
Move taxon	Genus	<i>Branisovskavirus</i>	<i>Caudoviricetes</i>	<i>Chimalliviridae</i>
Move taxon	Genus	<i>Chaoshanvirus</i>	<i>Caudoviricetes</i>	<i>Chimalliviridae</i>
Move taxon	Genus	<i>Eowynvirus</i>	<i>Caudoviricetes</i>	<i>Chimalliviridae</i>
Move taxon	Genus	<i>Ferozepurvirus</i>	<i>Caudoviricetes</i>	<i>Chimalliviridae</i>
Move taxon	Genus	<i>Ludhianavirus</i>	<i>Caudoviricetes</i>	<i>Chimalliviridae</i>
Move taxon	Genus	<i>Maaswegvirus</i>	<i>Caudoviricetes</i>	<i>Chimalliviridae</i>
Move taxon	Genus	<i>Meadowvirus</i>	<i>Caudoviricetes</i>	<i>Chimalliviridae</i>
Move taxon	Genus	<i>Miamivirus</i>	<i>Caudoviricetes</i>	<i>Chimalliviridae</i>
Move taxon	Genus	<i>Miltoncavirus</i>	<i>Caudoviricetes</i>	<i>Chimalliviridae</i>
Move taxon	Genus	<i>Nimduovirus</i>	<i>Caudoviricetes</i>	<i>Chimalliviridae</i>
Move taxon	Genus	<i>Pawinskivirus</i>	<i>Caudoviricetes</i>	<i>Chimalliviridae</i>
Move taxon	Genus	<i>Phabiovirus</i>	<i>Caudoviricetes</i>	<i>Chimalliviridae</i>
Move taxon	Genus	<i>Serwervirus</i>	<i>Caudoviricetes</i>	<i>Chimalliviridae</i>
Move taxon	Genus	<i>Siatvirus</i>	<i>Caudoviricetes</i>	<i>Chimalliviridae</i>
Move taxon	Genus	<i>Tepukevirus</i>	<i>Caudoviricetes</i>	<i>Chimalliviridae</i>

\*Source/full text: [https://ictv.global/ictv/proposals/2024.007B.Chimalliviridae\\_16mg.zip](https://ictv.global/ictv/proposals/2024.007B.Chimalliviridae_16mg.zip).

## 2024.008B.Colingsworthviridae\_1nf\_4ng\_3mg\_8ns

**Title:** Create a new family, *Colingsworthviridae*, of *Streptomyces* temperate phages (class *Caudoviricetes*)

**Authors:** Kurtböke I, Moraru C, Tolstoy I, Kropinski AM (Phage.Canada@gmail.com)

**Summary**

**Taxonomic rank(s) affected**

Realm *Duplodnaviria*, kingdom *Heunggongvirae*, phylum *Uroviricota*, class *Caudoviricetes*.

**Description of current taxonomy**

Three taxa of temperate *Streptomyces* phages exist as floating genera in the class *Caudoviricetes*, namely, *Vashvirus*, *Tigunavirus* and *Lomovskayavirus*. Up to the present, no effort has been made to assign them to higher taxa.

**Proposed taxonomic change(s)**

Create a new single species genus *Shadyvirus*.  
 Create a new single species genus *Sycamorevirus*.  
 Create a new single species genus *Shaekyvirus*.  
 Create a new genus *Sebastisaurusvirus* with three species.  
 To add a single new species to the genus *Vashvirus*.  
 To add a single new species to the genus *Lomovskayavirus*.  
 To create a new family, *Colingsworthviridae*, for these genera.  
 To transfer *Vashvirus*, *Tigunavirus* and *Lomovskayavirus* to this new family.

#### Justification

As a result of detailed genomic, proteomic and phylogenetic analyses using VIRIDIC, ViPTree, VirClust, we propose to create four new genera of Phi-C31-like temperate siphoviruses, in a new family named in honour of the first person to isolate a *Streptomyces* bacteriophage, Dr. Donald Colingsworth.

**Submitted:** 15 May 24; **Revised:** 30 September 24

**Table 15.** *Colingsworthviridae*, 13 new taxa\*

Operation	Rank	New taxon name	Virus name	Exemplar
New taxon	Family	<i>Colingsworthviridae</i>		
New taxon	Genus	<i>Shadyvirus</i>		
New taxon	Species	<i>Shadyvirus shady</i>	Streptomyces phage Shady	MT701596
New taxon	Genus	<i>Sycamorevirus</i>		
New taxon	Species	<i>Sycamorevirus sycamore</i>	Streptomyces phage Sycamore	MT701593
New taxon	Genus	<i>Shaekyvirus</i>		
New taxon	Species	<i>Shaekyvirus shaeky</i>	Streptomyces phage Shaeky	MT701595
New taxon	Genus	<i>Sebastisaurusvirus</i>		
New taxon	Species	<i>Sebastisaurusvirus sebastisaurus</i>	Streptomyces phage Sebastisaurus	MK450433
New taxon	Species	<i>Sebastisaurusvirus heather</i>	Streptomyces phage Heather	MK686069
New taxon	Species	<i>Sebastisaurusvirus remusloopin</i>	Streptomyces phage RemusLoopin	MK686068
New taxon	Species	<i>Vashvirus euratis</i>	Streptomyces phage Euratis	MK450426
New taxon	Species	<i>Lomovskayavirus shawty</i>	Streptomyces phage Shawty	MK433266

**Table 16.** *Colingsworthviridae*, 3 move taxa\*

Operation	Rank	Taxon name	New parent taxon	Old parent taxon
Move taxon	Genus	<i>Vashvirus</i>	<i>Colingsworthviridae</i>	<i>Caudoviricetes</i>
Move taxon	Genus	<i>Lomovskayavirus</i>	<i>Colingsworthviridae</i>	<i>Caudoviricetes</i>
Move taxon	Genus	<i>Tigunavirus</i>	<i>Colingsworthviridae</i>	<i>Caudoviricetes</i>

\*Source/full text: [https://ictv.global/ictv/proposals/2024.008B.Colingsworthviridae\\_1nf\\_4ng\\_3mg\\_8ns.zip](https://ictv.global/ictv/proposals/2024.008B.Colingsworthviridae_1nf_4ng_3mg_8ns.zip).

## 2024.009B.Connertonviridae\_1nf\_2mg\_12ns

**Title:** Create a new family, *Connertonviridae* for a group of *Campylobacter* phages (class: *Caudoviricetes*)

**Authors:** Moraru C, Tolstoy I, Kropinski AM (Phage.Canada@gmail.com)

#### Summary

##### Taxonomic rank(s) affected

Realm *Duplodnaviria*, kingdom *Heunggongvirae*, phylum *Uroviricota*, class *Caudoviricetes*.

##### Description of current taxonomy

At present the following taxa exist as genera within the floating subfamily *Eucampyvirinae*, class *Caudoviricetes*: *Fletchervirus* and *Firehammerivirus*.

**Proposed taxonomic change(s)**

To create eight new species in the genus *Fletchervirus*.

To create four new species in the genus *Firehammervirus*.

To create a new family *Connertonviridae* and abolish the subfamily *Eucampyvirinae*.

To move the genera *Fletchervirus* and *Firehammervirus* into this family.

**Justification**

We propose the abolishment of the subfamily *Eucampyvirinae* and the creation of a new family *Connertonviridae* based on analysis of the genera *Fletchervirus* and *Firehammervirus* using VIRIDIC, ViPTree, VirClust and phylogeny of 16 core proteins shared between the member species.

**Submitted:** 30 May 24

**Table 17.** *Connertonviridae*, 13 new taxa\*

Operation	Rank	New taxon name	Virus name	Exemplar
New taxon	Family	<i>Connertonviridae</i>		
New taxon	Species	<i>Fletchervirus F341</i>	Campylobacter phage F341	OQ864999
New taxon	Species	<i>Fletchervirus QDYZ</i>	Campylobacter phage vB_Cj_QDYZ	OQ515481
New taxon	Species	<i>Fletchervirus PC5</i>	Campylobacter phage PC5	KX229736
New taxon	Species	<i>Fletchervirus F336</i>	Campylobacter phage F336	MT863715
New taxon	Species	<i>Fletchervirus CJLB7</i>	Campylobacter phage CJLB-7	MW057933
New taxon	Species	<i>Fletchervirus CJLB10</i>	Campylobacter phage CJLB-10	MW074124
New taxon	Species	<i>Fletchervirus F372</i>	Campylobacter phage F372	MT863729
New taxon	Species	<i>Fletchervirus F207</i>	Campylobacter phage F207	MT863714
New taxon	Species	<i>Firehammervirus CJLB15</i>	Campylobacter phage CJLB-15	MW365733
New taxon	Species	<i>Firehammervirus F379</i>	Campylobacter phage F379	MT932329
New taxon	Species	<i>Firehammervirus CJLB12</i>	Campylobacter phage CJLB-12	MW074125
New taxon	Species	<i>Firehammervirus CJLB14</i>	Campylobacter phage CJLB-14	MW074126

**Table 18.** *Connertonviridae*, 2 move taxa\*

Operation	Rank	Taxon name	New parent taxon	Old parent taxon
Move taxon	Genus	<i>Fletchervirus</i>	<i>Connertonviridae</i>	<i>Eucampyvirinae</i>
Move taxon	Genus	<i>Firehammervirus</i>	<i>Connertonviridae</i>	<i>Eucampyvirinae</i>

**Table 19.** *Connertonviridae*, 1 abolish taxon\*

Operation	Rank	Abolished taxon name
Abolish taxon	Subfamily	<i>Eucampyvirinae</i>

\*Source/full text: [https://ictv.global/ictv/proposals/2024.009B.Connertonviridae\\_1nf\\_2mg\\_12ns.zip](https://ictv.global/ictv/proposals/2024.009B.Connertonviridae_1nf_2mg_12ns.zip).

**2024.010B.Dovevirinae\_1nsf\_1ng\_12ns**

**Title:** Create a new subfamily, *Dovevirinae*, with two genera for a group of lytic *Gordonia* phages (class: *Caudoviricetes*)

**Authors:** Kurtböke I, Moraru C, Tolstoy I, Kropinski AM (Phage.Canada@gmail.com)

**Summary****Taxonomic rank(s) affected**

Realm *Duplodnaviria*, kingdom *Heunggongvirae*, phylum *Uroviricota*, class *Caudoviricetes*.

**Description of current taxonomy**

Currently phages grouped within cluster DV in the Actinobacteriophage Database are recognized in the genus *Lambovirus*. These are lytic siphophages with circularly permuted genomes infecting *Gordonia* species.

**Proposed taxonomic change(s)**

The creation of a new subfamily, *Dovevirinae*, including two genera, *Lambovirus* and *Xeniaduovirus*, is proposed.

**Justification**

Constituent species in these genera exhibit greater than 50% nucleotide sequence similarity supporting their assignment to a new subfamily.

**Submitted:** 01 June 24

**Table 20.** *Dovevirinae*, 14 new taxa\*

Operation	Rank	New taxon name	Virus name	Exemplar
New taxon	Subfamily	<i>Dovevirinae</i>		
New taxon	Genus	<i>Xeniaduovirus</i>		
New taxon	Species	<i>Xeniaduovirus xenia2</i>	Gordonia phage Xenia2	PP725409
New taxon	Species	<i>Lambovirus wojtek</i>	Gordonia phage Wojtek	OL455890
New taxon	Species	<i>Lambovirus genamy16</i>	Gordonia phage Genamy16	ON755185
New taxon	Species	<i>Lambovirus novasharks</i>	Gordonia phage NovaSharks	ON755187
New taxon	Species	<i>Lambovirus zany</i>	Gordonia phage Zany	OL455887
New taxon	Species	<i>Lambovirus erutan</i>	Gordonia phage Erutan	OR475273
New taxon	Species	<i>Lambovirus jalebi</i>	Gordonia phage Jalebi	OL455895
New taxon	Species	<i>Lambovirus birthdayboy</i>	Gordonia phage BirthdayBoy	OR475261
New taxon	Species	<i>Lambovirus fulcrum</i>	Gordonia phage Fulcrum	OR521071
New taxon	Species	<i>Lambovirus parvustarda</i>	Gordonia phage ParvusTarda	OP172868
New taxon	Species	<i>Lambovirus otterstedtS21</i>	Gordonia phage OtterstedtS21	OP172870
New taxon	Species	<i>Lambovirus patos</i>	Gordonia phage Patos	OP172876

**Table 21.** *Dovevirinae*, 1 move taxon\*

Operation	Rank	Taxon name	New parent taxon	Old parent taxon
Move taxon	Genus	<i>Lambovirus</i>	<i>Dovevirinae</i>	<i>Caudoviricetes</i>

\*Source/full text: [https://ictv.global/ictv/proposals/2024.010B.Dovevirinae\\_1nsf\\_1ng\\_12ns.zip](https://ictv.global/ictv/proposals/2024.010B.Dovevirinae_1nsf_1ng_12ns.zip).

**2024.011B.Durnavirales\_1nf\_1ng\_1ns**

**Title:** Create a new family, *Soropartiviridae*, within the order *Durnavirales* for classification of partiti-like virus infecting thermoacidophilic bacteria

**Authors:** Syun-ichi Urayama (urayama.shunichi.gn@u.tsukuba.ac.jp), Akihito Fukudome, Eugene V. Koonin, Takuro Nunoura, Mart Krupovic (mart.krupovic@pasteur.fr)

**Summary****Taxonomic rank(s) affected**

*Riboviria*, *Orthornavirae*, *Pisuviricota*, *Duplopiviricetes*, *Durnavirales*.

**Description of current taxonomy**

Order *Durnavirales* includes six families of viruses with double-stranded RNA genomes. Most of the durnavirals infect fungal hosts, with the exception of partitivirids, which besides fungi, infect plants and protozoa, and picobirnavirids, which appear to infect bacteria.

**Proposed taxonomic change(s)**

Create a new family, *Soropartiviridae*, with a genus, *Caliparnavirus*, within the order *Durnavirales* to classify partiti-like viruses discovered in the hot spring samples and infecting thermoacidophilic bacteria.

**Justification**

Phylogenetic analysis based on the RNA-directed RNA polymerase (RdRP) placed the new group of bacterial partiti-like viruses outside of the established *Partitiviridae* genera. Furthermore, unlike all other classified partivirids, one of the two segments of soropartivirids is bicistronic.

**Submitted:** 21 June 24

**Table 22.** *Durnavirales*, 3 new taxa\*

Operation	Rank	New taxon name	Virus name	Exemplar
New taxon	Family	<i>Soropartiviridae</i>		
New taxon	Genus	<i>Caliparnavirus</i>		
New taxon	Species	<i>Caliparnavirus acidus</i>	hot spring partiti-like virus 1	RNA1: BTCP01000001; RNA2: BTCP01000004

\*Source/full text: [https://ictv.global/ictv/proposals/2024.011B.Durnavirales\\_1nf\\_1ng\\_1ns.zip](https://ictv.global/ictv/proposals/2024.011B.Durnavirales_1nf_1ng_1ns.zip).

**2024.012B.Ehrlichviridae\_1nf\_6ng\_1mg\_9ns**

**Title:** Create a new family, *Ehrlichviridae*, for a group of *Bacillus* Andromeda-like phages (class: *Caudoviricetes*)

**Authors:** Barylski J (b54026@amu.edu.pl), Moraru C, Tolstoy I, Kropinski AM

**Summary****Taxonomic rank(s) affected**

Realm *Duplodnaviria*, kingdom *Heunggongvirae*, phylum *Uroviricota*, class *Caudoviricetes*.

**Description of current taxonomy**

The genus *Andromedavirus* currently exists as a floating genus in the class *Caudoviricetes*.

**Proposed taxonomic change(s)**

To create a new genus *Suttonboningtonvirus* with one species.

To create a new genus *Gettysburgvirus* with three species.

To add three new species to the genus *Andromedavirus*.

To create a new single species genus *Anathvirus*.

To create a new single species genus *Dazunavirus*.

To create a new single species genus *Chennaivirus*.

To create a new single species genus *Nairobivirus*.

To create a new family, *Ehrlichviridae*, for the above-mentioned taxa.

**Justification**

The phages comprising these taxa form a deep branching clade using tBLASTx distances and single gene phylogeny. Core gene analysis shows the presence of 15 proteins conserved across all members of the proposed family.

**Submitted:** 19 April 24

**Table 23.** *Ehrlichviridae*, 16 new taxa\*

Operation	Rank	New taxon name	Virus name	Exemplar
New taxon	Family	<i>Ehrlichviridae</i>		
New taxon	Species	<i>Andromedavirus leo2</i>	<i>Bacillus phage Leo2</i>	KU836751
New taxon	Genus	<i>Suttonboningtonvirus</i>		
New taxon	Species	<i>Suttonboningtonvirus sv1ICo2020</i>	<i>Bacillus phage 1_ICo-2020</i>	MT700412
New taxon	Genus	<i>Gettysburgvirus</i>		
New taxon	Species	<i>Gettysburgvirus gv056SW001B</i>	<i>Bacillus phage 056SW001B</i>	MN176230
New taxon	Species	<i>Gettysburgvirus gv268TH004</i>	<i>Bacillus phage 268TH004</i>	MW394467
New taxon	Species	<i>Gettysburgvirus gv019DV002</i>	<i>Bacillus phage 019DV002</i>	MN176220
New taxon	Genus	<i>Anathvirus</i>		
New taxon	Species	<i>Anathvirus anath</i>	<i>Bacillus phage Anath</i>	MG983742

Continued

**Table 23.** Continued

Operation	Rank	New taxon name	Virus name	Exemplar
New taxon	Genus	<i>Dazunavirus</i>		
New taxon	Species	<i>Dazunavirus DZ1</i>	Bacillus phage DZ1	OR338916
New taxon	Genus	<i>Chennaivirus</i>		
New taxon	Species	<i>Chennaivirus MVCVPHSA1</i>	Staphylococcus phage MVC_VPHSA1	OR670591
New taxon	Genus	<i>Nairovirus</i>		
New taxon	Species	<i>Nairovirus nv36</i>	Bacillus phage vB_BpsS-36	MH884513

**Table 24.** *Ehrlichviridae*, 1 move taxon\*

Operation	Rank	Taxon name	New parent taxon	Old parent taxon
Move taxon	Genus	<i>Andromedavirus</i>	<i>Ehrlichviridae</i>	<i>Caudoviricetes</i>

\*Source/full text: [https://ictv.global/ictv/proposals/2024.012B.Ehrlichviridae\\_1nf\\_6ng\\_1mg\\_9ns.zip](https://ictv.global/ictv/proposals/2024.012B.Ehrlichviridae_1nf_6ng_1mg_9ns.zip).

## 2024.013B.Ferrettivirinae\_1nsf\_3ng\_38ns

**Title:** To create a new subfamily, *Ferrettivirinae*, for *Streptococcus* prophages (class: *Caudoviricetes*)

**Authors:** Tolstoy I, Moraru C, Kropinski AM (Phage.Canada@gmail.com)

### Summary

#### Taxonomic rank(s) affected

Realm *Duplodnaviria*, kingdom *Heunggongvirae*, phylum *Uroviricota*, class *Caudoviricetes*.

#### Description of current taxonomy

The viruses classified in this proposal do not have a current taxonomic assignment.

#### Proposed taxonomic change(s)

We propose a new subfamily in the class *Caudoviricetes*, named in honour of Professor Joseph J. Ferretti, of *Streptococcus* temperate siphophages including three genera: *Spinunavirus*, *Norfolkplacevirus* and *Hinxtonvirus*.

#### Justification

Collectively, these phages share ≥29.7% DNA sequence similarity and 14 homologous proteins (22.2% in common).

**Submitted:** 07 May 24

**Table 25.** *Ferrettivirinae*, 42 new taxa\*. Table too large, see supplementary information sheet supp\_info\_tab\_25

\*Source/full text: [https://ictv.global/ictv/proposals/2024.013B.Ferrettivirinae\\_1nsf\\_3ng\\_38ns.zip](https://ictv.global/ictv/proposals/2024.013B.Ferrettivirinae_1nsf_3ng_38ns.zip)

## 2024.014B.Grandevirales\_1no\_2nf\_3nsf\_4ng\_8ns

**Title:** Create one new order *Grandevirales* (*Duplodnaviria*)

**Authors:** Cook R, Pye HV, Crisci MA, Telatin A, Santini JM (j.santini@ucl.ac.uk), Adriaenssens EM (Evelien.adriaenssens@quadram.ac.uk)

### Summary

#### Taxonomic rank(s) affected

Realm *Duplodnaviria*, kingdom *Heunggongvirae*, phylum *Uroviricota*, class *Caudoviricetes*.

#### Description of current taxonomy

The viruses classified in this proposal do not have a current taxonomic assignment.

#### Proposed taxonomic change(s)

We propose the creation of one new order, *Grandevirales*, including two new families, *Lakviridae* and *Epsomviridae*, three sub-families (*Quingentivirinae*, *Quadringentisvirinae* and *Sescentorumvirinae*) and four genera (*Vetruanivirus*, *Hatfieldvirus*, *Amboselivirus* and *Wendovervirus*). We also propose the creation of eight novel species within these genera.

### Justification

We investigated the evolutionary relationships of 23 megaphage genomes with sizes greater than 400 kb and propose a taxonomy for their classification. Analysis of their putative proteins revealed that Lak phages form a deeply branching monophyletic clade within the class *Caudoviricetes* apart from all other genomes, and hence justifies the creation of a new order *Grandevirales*. One of the interesting features of this clade is that all current members are characterized by an alternative genetic code, where the TAG stop codon is repurposed to encode an amino acid.

**Submitted:** 04 June 23; **Revised:** 07 October 24

**Table 26.** *Grandevirales*, 18 new taxa\*

Operation	Rank	New taxon name	Virus name	Exemplar
New taxon	Order	<i>Grandevirales</i>		
New taxon	Family	<i>Lakviridae</i>		
New taxon	Subfamily	<i>Quingentivirinae</i>		
New taxon	Genus	<i>Vetruanivirus</i>		
New taxon	Species	<i>Vetruanivirus primi</i>	Prevotella phage Lak-A1	MK250015
New taxon	Species	<i>Vetruanivirus secundi</i>	Prevotella phage Lak-A2	MK250019
New taxon	Species	<i>Vetruanivirus dhakaense</i>	Prevotella phage Lak-C1	MK250029
New taxon	Species	<i>Vetruanivirus porcinprimi</i>	uncultured phage RVC AP1_GC26	OR769218
New taxon	Species	<i>Vetruanivirus porcinsecundi</i>	uncultured phage RVC AP3_GC26	OR769219
New taxon	Genus	<i>Amboselivirus</i>		
New taxon	Species	<i>Amboselivirus simi</i>	Prevotella phage Lak-B1	MK250020
New taxon	Subfamily	<i>Quadrangentivirinae</i>		
New taxon	Genus	<i>Hatfieldvirus</i>		
New taxon	Species	<i>Hatfieldvirus porci</i>	uncultured phage RVC_JS4_GC31	OR769222
New taxon	Family	<i>Epsomviridae</i>		
New taxon	Subfamily	<i>Sescentorumvirinae</i>		
New taxon	Genus	<i>Wendovervirus</i>		
New taxon	Species	<i>Wendovervirus sonii</i>	uncultured phage HB1	OR769223

\*Source/full text: [https://ictv.global/ictv/proposals/2024.014B.Grandevirales\\_1no\\_2nf\\_3nsf\\_4ng\\_8ns.zip](https://ictv.global/ictv/proposals/2024.014B.Grandevirales_1no_2nf_3nsf_4ng_8ns.zip).

### 2024.015B.Grimontviridae\_2ng\_1mg\_2ns

**Title:** To update the family *Grimontviridae* through the addition of three genera (class: *Caudoviricetes*)

**Authors:** Dechesne A, Moraru C, Parra B, Tolstoy I, Kropinski AM (Phage.Canada@gmail.com)

#### Summary

##### Taxonomic rank(s) affected

Realm *Duplodnaviria*, kingdom *Heunggongvirae*, phylum *Uroviricota*, class *Caudoviricetes*, family *Grimontviridae*.

##### Description of current taxonomy

The C3 group of podoviruses are a very rare morphotype possessing elongated capsids, the ‘type virus’ of which is *Escherichia* phage phiEco32. The family *Grimontviridae* currently consists of five genera: *Crifsvirus*, *Dalianvirus*, *Libingvirus*, *Moazamivirus* and *Privateervirus*.

##### Proposed taxonomic change(s)

Create a new single species genus *Lundtoftevirus* Transfer the genus *Lahexavirus* to the family *Grimontviridae*.

**Justification**

The family is represented by a cohesive and monophyletic group in the main predicted proteome-based clustering tools (VirClust, ViPTree and vConTACT2). Members of the family share 12 core proteins.

**Submitted:** 06 May 24; **Revised:** 30 June 24

**Table 27.** *Grimontviridae*, 2 new taxa.\*

Operation	Rank	New taxon name	Virus name	Exemplar
New taxon	Genus	<i>Lundtoftevirus</i>		
New taxon	Species	<i>Lundtoftevirus Lu221</i>	IncN phage Lu221	OQ829281

**Table 28.** *Grimontviridae*, 2 move taxa\*

Operation	Rank	Taxon name	New parent taxon	Old parent taxon
Move taxon	Genus	<i>Lahexavirus</i>	<i>Grimontviridae</i>	<i>Caudoviricetes</i>
Move taxon	Genus	<i>Aquaneticvirus</i>	<i>Grimontviridae</i>	<i>Caudoviricetes</i>

\*Source/full text: [https://ictv.global/ictv/proposals/2024.015B.Grimontviridae\\_2ng\\_1mg\\_2ns.zip](https://ictv.global/ictv/proposals/2024.015B.Grimontviridae_2ng_1mg_2ns.zip).

**2024.016B.Hodgkinviridae\_1nf\_2ng\_4mg\_5ns**

**Title:** Create a new family, *Hodgkinviridae*, for a group of lytic *Microbacterium* phages (class: *Caudoviricetes*)

**Authors:** Kurtböke I, Moraru C, Tolstoy I, Kropinski AM (Phage.Canada@gmail.com)

**Summary****Taxonomic rank(s) affected**

Realm *Duplodnaviria*, kingdom *Heunggongvirae*, phylum *Uroviricota*, class *Caudoviricetes*.

**Description of current taxonomy**

The genera *Mementomorivirus*, *Quhwahvirus* and *Meganvirus* exist as floating genera in the class *Caudoviricetes*.

**Proposed taxonomic change(s)**

To create a new single-species genus, *Fuzzbustervirus*

To add a single new species to the genus *Kozievirus*.

To split the genus *Mementomorivirus* in two, creating *Margaeryvirus*.

To add a single new species to the genus *Meganvirus*.

To add two species to the genus *Quhwahvirus*.

To create a new family, *Hodgkinviridae*, for these genera and *Metamorphoovirus*.

**Justification**

Using VIRIDIC, ViPTree, VIRCLUST and vConTACT v.3.0, we have established that this is a cohesive group of lytic *Microbacterium* siphoviruses which share ≥12.2% DNA sequence similarity and 14 common proteins.

**Submitted:** 27 May 24; **Revised:** 30 September 24

**Table 29.** *Hodgkinviridae*, 8 new taxa\*

Operation	Rank	New taxon name	Virus name	Exemplar
New taxon	Family	<i>Hodgkinviridae</i>		
New taxon	Species	<i>Kozievirus MO526</i>	<i>Microbacterium</i> phage MO526	OR941552
New taxon	Species	<i>Meganvirus nichole72</i>	<i>Microbacterium</i> phage Nicole72	OR159674
New taxon	Species	<i>Quhwahvirus pulchra</i>	<i>Microbacterium</i> phage Pulchra	MW601217
New taxon	Species	<i>Quhwahvirus littlefortune</i>	<i>Microbacterium</i> phage LittleFortune	OR475280
New taxon	Genus	<i>Fuzzbustervirus</i>		
New taxon	Species	<i>Fuzzbustervirus fuzzbuster</i>	<i>Microbacterium</i> phage FuzzBuster	MN062720
New taxon	Genus	<i>Margaeryvirus</i>		

**Table 30.** *Hodgkinviridae*, 4 move taxa\*

Operation	Rank	Taxon name	New parent taxon	Old parent taxa
Move taxon	Subfamily	<i>Kutznervirinae</i>	<i>Hodgkinviridae</i>	<i>Caudoviricetes</i>
Move taxon	Genus	<i>Meganvirus</i>	<i>Hodgkinviridae</i>	<i>Kutznervirinae</i>
Move taxon	Genus	<i>Quhwahvirus</i>	<i>Hodgkinviridae</i>	<i>Kutznervirinae</i>
Move taxon	Genus	<i>Metamorphoovirus</i>	<i>Hodgkinviridae</i>	<i>Kutznervirinae</i>

**Table 31.** *Hodgkinviridae*, 2 move; rename taxa\*

Operation	Rank	New taxon name	New parent taxon	Old taxon name
Move; rename taxon	Species	<i>Margaeryvirus margaery</i>	<i>Margaeryvirus</i>	<i>Mementomorivivirus margaery</i>
Move; rename taxon	Species	<i>Margaeryvirus terij</i>	<i>Margaeryvirus</i>	<i>Mementomorivivirus terij</i>

\*Source/full text: [https://ictv.global/ictv/proposals/2024.016B.Hodgkinviridae\\_1nf\\_2ng\\_4mg\\_5ns.zip](https://ictv.global/ictv/proposals/2024.016B.Hodgkinviridae_1nf_2ng_4mg_5ns.zip).

## 2024.017B.Jeanschmidtviridae\_1nf\_3ng\_4mg\_6ns

**Title:** Create a new family, *Jeanschmidtviridae* for a group of *Caulobacter* and *Brevundimonas* phages (class: *Caudoviricetes*)

**Authors:** Millard A, Moraru C, Tolstoy I, Kropinski AM (Phage.Canada@gmail.com)

### Summary

#### Taxonomic rank(s) affected

Realm *Duplodnaviria*, kingdom *Heunggongviriae*, phylum *Uroviricota*, class *Caudoviricetes*.

#### Description of current taxonomy

The taxa *Colossusvirus*, *Bertelyvirus*, *Shapirovirus* and *Poindextervirus* are floating genera in the class *Caudoviricetes*.

#### Proposed taxonomic change(s)

To create a new genus, *Kikimoravirus*, with two species.

To create a new genus, *Marchewkavirus*, with three species.

To create a single-species genus, *Bajunvirus*. Abolish the subfamily *Dolichocephalovirinae*.

To create a new family, *Jeanschmidtviridae*, for these genera and *Colossusvirus*, *Bertelyvirus*, *Shapirovirus* and *Poindextervirus*.

#### Justification

The proposed members share ≥10.3% DNA sequence similarity and share 38 protein homologs.

**Submitted:** 10 June 24; **Revised:** 30 September 24

**Table 32.** *Jeanschmidtviridae*, 10 new taxa\*

Operation	Rank	New taxon name	Virus name	Exemplar
New taxon	Family	<i>Jeanschmidtviridae</i>		
New taxon	Genus	<i>Kikimoravirus</i>		
New taxon	Species	<i>Kikimoravirus kikimora</i>	Brevundimonas phage vB_BpoS-Kikimora	ON529857
New taxon	Species	<i>Kikimoravirus gurke</i>	Brevundimonas phage vB_BpoS-Gurke	ON529850
New taxon	Genus	<i>Marchewkavirus</i>		
New taxon	Species	<i>Marchewkavirus marchewka</i>	Brevundimonas phage vB_BpoS-Marchewka	ON529851
New taxon	Species	<i>Marchewkavirus kabachok</i>	Brevundimonas phage vB_BpoS-Kabachok	ON529852
New taxon	Species	<i>Marchewkavirus domovoi</i>	Brevundimonas phage vB_BpoS-Domovoi	ON529855
New taxon	Genus	<i>Bajunvirus</i>		
New taxon	Species	<i>Bajunvirus bajun</i>	Brevundimonas phage vB_BpoS-Bajun	ON529858

**Table 33.** Jeanschmidtviridae, 4 move taxa\*

Operation	Rank	Taxon name	New parent taxon	Old parent taxon
Move taxon	Genus	<i>Colossusvirus</i>	<i>Jeanschmidtviridae</i>	<i>Dolichocephalovirinae</i>
Move taxon	Genus	<i>Bertelyvirus</i>	<i>Jeanschmidtviridae</i>	<i>Dolichocephalovirinae</i>
Move taxon	Genus	<i>Shapirovirus</i>	<i>Jeanschmidtviridae</i>	<i>Dolichocephalovirinae</i>
Move taxon	Genus	<i>Poindextervirus</i>	<i>Jeanschmidtviridae</i>	<i>Dolichocephalovirinae</i>

**Table 34.** Jeanschmidtviridae, 1 abolish taxon\*

Operation	Rank	Abolished taxon name
Abolish taxon	Subfamily	<i>Dolichocephalovirinae</i>

\*Source/full text: [https://ictv.global/ictv/proposals/2024.017B.Jeanschmidtviridae\\_1nf\\_3ng\\_4mg\\_6ns.zip](https://ictv.global/ictv/proposals/2024.017B.Jeanschmidtviridae_1nf_3ng_4mg_6ns.zip).

## 2024.018B.Kronosvirus\_1ng\_3ns

**Title:** Create one new genus (*Kronosvirus*) with three species (*Caudoviricetes*).

**Author:** Ely B (ely@sc.edu)

### Summary

#### Taxonomic rank(s) affected

Realm *Duplodnaviria*, kingdom *Heunggongvirae*, phylum *Uroviricota*, class *Caudoviricetes*.

#### Description of current taxonomy

The viruses classified in this proposal do not have a current taxonomic assignment.

#### Proposed taxonomic change(s)

Caulobacter bacteriophage Kronos is a unique siphovirus with a 50 nm icosahedral head and a long (150 nm) flexible tail. Aside from two phages, there are no close matches of its genome sequence to any other bacteriophage in the GenBank database. Additional phages that are closely related to Kronos can be grouped into three distinct species based on the ICTV criteria.

#### Justification

The creation of the genus *Kronosvirus* complies with current demarcation criteria [2].

**Submitted:** 27 May 24

**Table 35.** *Kronosvirus*, 4 new taxa\*

Operation	Rank	New taxon name	Virus name	Exemplar
New taxon	Genus	<i>Kronosvirus</i>		
New taxon	Species	<i>Kronosvirus pelion</i>	Caulobacter phage Kronos	MH884648
New taxon	Species	<i>Kronosvirus pomeria</i>	Caulobacter phage TMCBR2	OQ269668
New taxon	Species	<i>Kronosvirus elgin</i>	Caulobacter phage TMCBR4	OQ330850

\*Source/full text: [https://ictv.global/ictv/proposals/2024.018B.Kronosvirus\\_1ng\\_3ns.zip](https://ictv.global/ictv/proposals/2024.018B.Kronosvirus_1ng_3ns.zip).

## 2024.019B.Kruegerviridae\_1nf\_1ng\_1mg\_4ns

**Title:** Create a new family, *Kruegerviridae*, for a group of *Gordonia* phages (class: *Caudoviricetes*)

**Authors:** Kurtböke I, Moraru C, Tolstoy I, Kropinski AM (Phage.Canada@gmail.com)

### Summary

#### Taxonomic rank(s) affected

Realm *Duplodnaviria*, kingdom *Heunggongvirae*, phylum *Uroviricota*, class *Caudoviricetes*.

#### Description of current taxonomy

*Vanleevirus* currently exists as a floating genus in the class *Caudoviricetes*.

**Proposed taxonomic change(s)**

To create a new genus *Cafassovirus* with four species.

To create a new family, *Kruegerviridae*, for the genera *Cafassovirus* and *Vanleevirus*.

**Justification**

By VIRIDIC analysis, members of these two genera share ≥18.8% DNA sequence similarity and also share 46 protein homologs. The genera *Vanleevirus* and *Cafassovirus* form a deep-branching clade using tBLASTx distances, commensurate with the establishment of a new family of bacterial viruses.

**Submitted:** 28 May 24; **Revised:** 30 September 24

**Table 36.** *Kruegerviridae*, 6 new taxa\*

Operation	Rank	New taxon name	Virus name	Exemplar
New taxon	Family	<i>Kruegerviridae</i>		
New taxon	Genus	<i>Cafassovirus</i>		
New taxon	Species	<i>Cafassovirus cafasso</i>	Gordonia phage Cafasso	MZ322021
New taxon	Species	<i>Cafassovirus morgana</i>	Gordonia phage Morgana	PP537962
New taxon	Species	<i>Cafassovirus aleemily</i>	Gordonia phage Aleemily	ON970578
New taxon	Species	<i>Cafassovirus obladi</i>	Gordonia phage ObLaDi	OP297535

**Table 37.** *Kruegerviridae*, 1 move taxon\*

Operation	Rank	Taxon name	New parent taxon	Old parent taxon
Move taxon	Genus	<i>Vanleevirus</i>	<i>Kruegerviridae</i>	<i>Caudoviricetes</i>

\*Source/full text: [https://ictv.global/ictv/proposals/2024.019B.Kruegerviridae\\_1nf\\_1ng\\_1mg\\_4ns.zip](https://ictv.global/ictv/proposals/2024.019B.Kruegerviridae_1nf_1ng_1mg_4ns.zip).

**2024.020B.Lindbergviridae\_1nf\_3ng\_7mg\_21ns**

**Title:** Create a new family, *Lindbergviridae*, for PB1-like phages (class: *Caudoviricetes*)

**Authors:** Moraru C, Tolstoy I, Kropinski AM (Phage.Canada@gmail.com)

**Summary****Taxonomic rank(s) affected**

Realm *Duplodnaviria*, kingdom *Heunggongvirae*, phylum *Uroviricota*, class *Caudoviricetes*.

**Description of current taxonomy**

PB1-like phages have been classified into the following genera: *Kylevirus*, *Tabernariusvirus*, *Bcepfunavirus*, *Pbunavirus*, *Wifcevirus*, *Myosmarvirus* and *Carpasinavirus*. All these are myoviruses infecting members of the Betaproteobacteria and Gammaproteobacteria.

**Proposed taxonomic change(s)**

To create ten new species in the genus *Pbunavirus*.

To create one new species in the genus *Myosmarvirus*.

To add six new species to the genus *Wifcevirus*.

To add one new species to the genus *Carpasinavirus*.

To create a new single species genus *Gladiolivirus*.

To create a new single species genus *Irusalimvirus*.

To create a new single species genus *Plutovirus*.

To create a new family, *Lindbergviridae*, for the above-mentioned taxa as well as *Kylevirus*, *Tabernariusvirus* and *Bcepfunavirus*.

**Justification**

All our genomic and proteomic analyses reveal that the previously established genera *Kylevirus* (2020.086B.R.Kylevirus), *Tabernariusvirus* (2018.099B.A.v1.Tabernariusvirus), *Bcepfunavirus* (2020.116B.R.Pbunavirus), *Pbunavirus*, *Wifcevirus*, *Myosmarvirus* and *Carpasinavirus* together with the three new genera listed above belong to a new family which we have named in honour of Alf A. Lindberg. The bacteriophages share 12 common proteins.

**Submitted:** 27 April 24; **Revised:** 30 September 24

**Table 38.** *Lindbergviridae*, 25 new taxa\*

Operation	Rank	New taxon name	Virus name	Exemplar
New taxon	Family	<i>Lindbergviridae</i>		
New taxon	Species	<i>Pbunavirus SG1</i>	Pseudomonas phage SG1	OQ594965
New taxon	Species	<i>Pbunavirus pv109</i>	Pseudomonas phage 109	OQ831730
New taxon	Species	<i>Pbunavirus FBPa14</i>	Pseudomonas phage vB_PaeM_FBPa14	ON375839
New taxon	Species	<i>Pbunavirus wadjak13</i>	Pseudomonas phage Kara-mokiny kep-wari Wadjak_13	OP310979
New taxon	Species	<i>Pbunavirus TH15</i>	Pseudomonas phage TH15	MW406974
New taxon	Species	<i>Pbunavirus PSA09</i>	Pseudomonas phage PSA09	MZ089730
New taxon	Species	<i>Pbunavirus ph0031</i>	Pseudomonas phage PhL_UNISO_PA-DSM_ph0031	MW526258
New taxon	Species	<i>Pbunavirus PSA25</i>	Pseudomonas phage PSA25	MZ089736
New taxon	Species	<i>Pbunavirus FBPa35</i>	Pseudomonas phage vB_PaeM_FBPa35	ON857938
New taxon	Species	<i>Pbunavirus victoria</i>	Pseudomonas phage Victoria	OR805296
New taxon	Species	<i>Myosmarvirus SMP</i>	Serratia phage SMP	OP490597
New taxon	Species	<i>Wifcevirus SP13</i>	Escherichia phage vB_EcoM_SP13	OP352608
New taxon	Species	<i>Wifcevirus AV128</i>	Escherichia phage AV128	OR352958
New taxon	Species	<i>Wifcevirus Ro157lw</i>	Escherichia phage vB_EcoM-Ro157lw	MH051335
New taxon	Species	<i>Wifcevirus EC150</i>	Escherichia phage EC150	ON210137
New taxon	Species	<i>Wifcevirus mansfield</i>	Escherichia phage Mansfield	MK903282
New taxon	Species	<i>Wifcevirus ECO71P1</i>	Escherichia phage ECO71P1	OP172789
New taxon	Species	<i>Carpasinavirus FoX6</i>	Xanthomonas phage FoX6	MT161386
New taxon	Genus	<i>Gladiolivivirus</i>		
New taxon	Species	<i>Gladiolivivirus maja</i>	Burkholderia phage Maja	MT708549
New taxon	Genus	<i>Irusalimvirus</i>		
New taxon	Species	<i>Irusalimvirus BCSR52</i>	Burkholderia phage BCSR52	MW460246
New taxon	Genus	<i>Plutovirus</i>		
New taxon	Species	<i>Plutovirus pluto</i>	Luteibacter phage vB_LfIM-Pluto	ON529861

**Table 39.** *Lindbergviridae*, 7 move taxa\*

Operation	Rank	Taxon name	New parent taxon	Old parent taxa
Move taxon	Genus	<i>Kylevirus</i>	<i>Lindbergviridae</i>	<i>Caudoviricetes</i>
Move taxon	Genus	<i>Tabernariusvirus</i>	<i>Lindbergviridae</i>	<i>Caudoviricetes</i>
Move taxon	Genus	<i>Bcepfunavirus</i>	<i>Lindbergviridae</i>	<i>Caudoviricetes</i>
Move taxon	Genus	<i>Pbunavirus</i>	<i>Lindbergviridae</i>	<i>Caudoviricetes</i>
Move taxon	Genus	<i>Myosmarvirus</i>	<i>Lindbergviridae</i>	<i>Caudoviricetes</i>
Move taxon	Genus	<i>Wifcevirus</i>	<i>Lindbergviridae</i>	<i>Caudoviricetes</i>
Move taxon	Genus	<i>Carpasinavirus</i>	<i>Lindbergviridae</i>	<i>Caudoviricetes</i>

\*Source/full text: [https://ictv.global/ictv/proposals/2024.020B.Lindbergviridae\\_1nf\\_3ng\\_7mg\\_21ns.zip](https://ictv.global/ictv/proposals/2024.020B.Lindbergviridae_1nf_3ng_7mg_21ns.zip).

## 2024.021B.Malkevirus\_1ng\_5ns

**Title:** To create a new genus, *Malkevirus*, for *Streptococcus* prophages (class: *Caudoviricetes*)

**Authors:** Tolstoy I, Moraru C, Kropinski AM (Phage.Canada@gmail.com)

### Summary

#### Taxonomic rank(s) affected

Realm *Duplodnaviria*, kingdom *Heunggongvirae*, phylum *Uroviricota*, class *Caudoviricetes*.

#### Description of current taxonomy

The viruses classified in this proposal do not have a current taxonomic assignment.

#### Proposed taxonomic change(s)

We propose a new genus, named in honour of Professor Dr. Horst Malke, comprised of temperate siphoviruses infecting *Streptococcus spp.*

#### Justification

The viruses fall into a genus based on current demarcation criteria.

**Submitted:** 07 May 24

**Table 40.** *Malkevirus*, 6 new taxa\*

Operation	Rank	New taxon name	Virus name	Exemplar
New taxon	Genus	<i>Malkevirus</i>		
New taxon	Species	<i>Malkevirus ARI02853</i>	Streptococcus phage phiARI0285-3	KT337347
New taxon	Species	<i>Malkevirus IPP67</i>	Streptococcus phage IPP67	KY065503
New taxon	Species	<i>Malkevirus IPP45</i>	Streptococcus phage IPP45	KY065485
New taxon	Species	<i>Malkevirus mv23782</i>	Streptococcus phage 23782	FR671408
New taxon	Species	<i>Malkevirus mv11865</i>	Streptococcus phage 11865	FR671409

\*Source/full text: [https://ictv.global/ictv/proposals/2024.021B.Malkevirus\\_1ng\\_5ns.zip](https://ictv.global/ictv/proposals/2024.021B.Malkevirus_1ng_5ns.zip).

## 2024.022B.Markadamsvirinae\_1ng\_1ns

**Title:** Create a new genus, *Kononvirus*, with a single species in the subfamily *Markadamsvirinae* (*Caudoviricetes: Demerecviridae*)

**Authors:** Wójcicki M (michal.wojcicki@hirsfeld.pl), Shymialevich D, Średnicka P, Gientka I, Błażejak S, Sokołowska B

### Summary

#### Taxonomic rank(s) affected

Proposal to create a new genus (*Kononvirus*) with a single species (tailed phages from class *Caudoviricetes*, family *Demerecviridae* and subfamily *Markadamsvirinae*).

#### Description of current taxonomy

In 2022, significant changes to the taxonomy of bacterial viruses were introduced: the paraphyletic morphological families *Podoviridae*, *Siphoviridae* and *Myoviridae* as well as the order *Caudovirales* were abolished, which is replaced by the class *Caudoviricetes* to group all tailed bacterial and archaeal viruses with icosahedral capsids and a double-stranded DNA genome. Moreover, a binomial system of nomenclature for species was implemented. Currently, the family *Demerecviridae* includes three subfamilies (*Ermolyevavirinae*, *Markadamsvirinae* and *Mccorquodalevirinae*) and six separate genera (*Keyvirus*, *Novosibivirus*, *Pogseptimavirus*, *Priunavirus*, *Shenzenvirus* and *Sugarlandvirus*) not classified at the subfamily level. The subfamily *Markadamsvirinae* currently includes two genera: *Epseptimavirus* and *Tequintavirus*.

#### Proposed taxonomic change(s)

We performed genomic analysis of newly isolated *Enterobacter* phage KKP\_3711. Genome and protein analyses suggest that this bacteriophage belongs to the subfamily *Markadamsvirinae*. Still, the differences are too great to assign it to one of the two genera within this subfamily. Therefore, we propose to create a new genus, *Kononvirus*, including a single species, *Kononvirus KKP3711*.

#### Justification

Based on DNA and protein similarity, this is a cohesive genus.

**Submitted:** 16 June 24

**Table 41.** *Markadamsvirinae*, 2 new taxa\*

Operation	Rank	New taxon name	Virus name	Exemplar
New taxon	Genus	<i>Kononvirus</i>		
New taxon	Species	<i>Kononvirus KKP3711</i>	Enterobacter siphophage KKP_3711	PP579741

\*Source/full text: [https://ictv.global/ictv/proposals/2024.022B.Markadamsvirinae\\_1ng\\_1ns.zip](https://ictv.global/ictv/proposals/2024.022B.Markadamsvirinae_1ng_1ns.zip).

## 2024.023B.Mcshanvirinae\_1nsf\_3ng\_25ns

**Title:** To create a new subfamily, *Mcshanvirinae*, for *Streptococcus* prophages (class: *Caudoviricetes*)

**Authors:** Tolstoy I, Moraru C, Kropinski AM (Phage.Canada@gmail.com)

### Summary

#### Taxonomic rank(s) affected

Realm *Duplodnaviria*, kingdom *Heunggongvirae*, phylum *Uroviricota*, class *Caudoviricetes*.

#### Description of current taxonomy

The viruses classified in this proposal do not have a current taxonomic assignment.

#### Proposed taxonomic change(s)

We propose a new subfamily, named in honour of Professor W. Michael McShan, of *Streptococcus* temperate siphoviruses containing three newly established genera: *Adrianbuildvirus*, *Medawarvirus* and *Phadecaviru*s.

#### Justification

The proposed taxa conform to the demarcation criteria employed by the ICTV Bacterial Viruses Subcommittee.

**Submitted:** 06 May 2024

**Table 42.** *Mcshanvirinae*, 29 new taxa\*

Operation	Rank	New taxon name	Virus name	Exemplar
New taxon	Subfamily	<i>Mcshanvirinae</i>		
New taxon	Genus	<i>Adrianbuildvirus</i>		
New taxon	Species	<i>Adrianbuildvirus SpSL1</i>	Streptococcus phage SpSL1	KM882824
New taxon	Species	<i>Adrianbuildvirus IPP5</i>	Streptococcus phage IPP5	KY065449
New taxon	Species	<i>Adrianbuildvirus IPP44</i>	Streptococcus phage IPP44	KY065484
New taxon	Species	<i>Adrianbuildvirus IPP42</i>	Streptococcus phage IPP42	KY065482
New taxon	Species	<i>Adrianbuildvirus IPP51</i>	Streptococcus phage IPP51	KY065489
New taxon	Species	<i>Adrianbuildvirus ARI0923</i>	Streptococcus phage phiARI0923	KT337370
New taxon	Species	<i>Adrianbuildvirus IPP41</i>	Streptococcus phage IPP41	KY065481
New taxon	Species	<i>Adrianbuildvirus IPP43</i>	Streptococcus phage IPP43	KY065483
New taxon	Genus	<i>Medawarvirus</i>		
New taxon	Species	<i>Medawarvirus IPP12</i>	Streptococcus phage IPP12	KY065454
New taxon	Species	<i>Medawarvirus IPP22</i>	Streptococcus phage IPP22	KY065463
New taxon	Species	<i>Medawarvirus IPP18</i>	Streptococcus phage IPP18	KY065459
New taxon	Species	<i>Medawarvirus IPP57</i>	Streptococcus phage IPP57	KY065494
New taxon	Species	<i>Medawarvirus IPP20</i>	Streptococcus phage IPP20	KY065461
New taxon	Species	<i>Medawarvirus IPP21</i>	Streptococcus phage IPP21	KY065462
New taxon	Species	<i>Medawarvirus IPP30</i>	Streptococcus phage IPP30	KY065471
New taxon	Species	<i>Medawarvirus IPP11</i>	Streptococcus phage IPP11	KY065453
New taxon	Species	<i>Medawarvirus IPP29</i>	Streptococcus phage IPP29	KY065470

Continued

**Table 42.** Continued

Operation	Rank	New taxon name	Virus name	Exemplar
New taxon	Species	<i>Medawarvirus IPP19</i>	Streptococcus phage IPP19	KY065460
New taxon	Species	<i>Medawarvirus ARI01312</i>	Streptococcus phage phiARI0131-2	KT337342
New taxon	Species	<i>Medawarvirus IPP63</i>	Streptococcus phage IPP63	KY065499
New taxon	Species	<i>Medawarvirus IPP17</i>	Streptococcus phage IPP17	KY065458
New taxon	species	<i>Medawarvirus IPP28</i>	Streptococcus phage IPP28	KY065469
New taxon	Genus	<i>Phadecavirus</i>		
New taxon	Species	<i>Phadecavirus PH10</i>	Streptococcus phage PH10	FN391954
New taxon	Species	<i>Phadecavirus pv23TH</i>	Streptococcus phage 23TH	MT900487
New taxon	Species	<i>Phadecavirus olisA1</i>	Streptococcus phage OlisA1	OL774868

\*Source/full text: [https://ictv.global/ictv/proposals/2024.023B.Mcshanvirinae\\_1nsf\\_3ng\\_25ns.zip](https://ictv.global/ictv/proposals/2024.023B.Mcshanvirinae_1nsf_3ng_25ns.zip).

## 2024.024B.Mtkvariviridae\_1nf\_1msf\_10ns

**Title:** Create a new family, *Mtkvariviridae*, for PhiEco32-like phages (class: *Caudoviricetes*)

**Authors:** Moraru C, Tolstoy I, Kropinski AM (Phage.Canada@gmail.com)

### Summary

#### Taxonomic rank(s) affected

Realm *Duplodnaviria*, kingdom *Heunggongvirae*, phylum *Uroviricota*, class *Caudoviricetes*.

#### Description of current taxonomy

The genera *Kuravirus*, *Nieuwеканаalvirus* and *Suseptimavirus* are currently classified within the subfamily *Gordonclarkvirinae*, class *Caudoviricetes*.

#### Proposed taxonomic change(s)

Create eight new species in the genus *Kuravirus*.

Create two new species in the genus *Suseptimavirus*.

Create a new family, *Mtkvariviridae*.

#### Justification

Species classified to the genera *Kuravirus*, *Nieuwеканаalvirus* and *Suseptimavirus* within the subfamily *Gordonclarkvirinae* share 12 core proteins and form a deep branching clade in ViPTree tBLASTx distance analysis.

**Submitted:** 06 May 24; **Revised:** 30 September 24

**Table 43.** *Mtkvariviridae*, 11 new taxa\*

Operation	Rank	New taxon name	Virus name	Exemplar
New taxon	Family	<i>Mtkvariviridae</i>		
New taxon	Species	<i>Kuravirus myPSH1131</i>	Escherichia phage myPSH1131	MG983840
New taxon	Species	<i>Kuravirus myPSH2311</i>	Escherichia phage myPSH2311	MG976803
New taxon	Species	<i>Kuravirus XT18</i>	Escherichia phage vB-EcoP-XT18	OR757434
New taxon	Species	<i>Kuravirus LAMP</i>	Escherichia phage LAMP	MG673519
New taxon	Species	<i>Kuravirus SDYTW1F1223</i>	Escherichia phage SDYTW1-F1-2-2_3	OR296290
New taxon	Species	<i>Kuravirus pECN12032Af1</i>	Escherichia phage pEC-N1203-2Af.1	OQ540978
New taxon	Species	<i>Kuravirus YF01</i>	Escherichia phage vB_EcoP_YF01	OQ025076
New taxon	Species	<i>Kuravirus SR02</i>	Escherichia phage SR02	OQ870566
New taxon	Species	<i>Suseptimavirus PAS59</i>	Escherichia phage vB_EcoP_PAS59	OQ921332
New taxon	Species	<i>Suseptimavirus sv4E8</i>	Escherichia phage 4E8	OQ689734

**Table 44.** *Mtkvariviridae*, 1 move taxon\*

Operation	Rank	Taxon name	New parent taxon	Old parent taxon
Move taxon	Subfamily	<i>Gordonclarkvirinae</i>	<i>Mtkvariviridae</i>	<i>Caudoviricetes</i>

\*Source/full text: [https://ictv.global/ictv/proposals/2024.024B.Mtkvariviridae\\_1nf\\_1msf\\_10ns.zip](https://ictv.global/ictv/proposals/2024.024B.Mtkvariviridae_1nf_1msf_10ns.zip).

## 2024.025B.Obscuriviridae\_1nf\_2ng\_3ns

**Title:** Create a new family, *Obscuriviridae*

**Authors:** Bartlau N (nina.bartlau@univie.ac.at), Moraru C, Wichels A, Holmfeldt K, Amann RI

### Summary

#### Taxonomic rank(s) affected

New, unassigned family.

#### Description of current taxonomy

The viruses described in this proposal do not have a current taxonomic assignment.

#### Proposed taxonomic change(s)

A new family, *Obscuriviridae*, is proposed, including two new genera, *Omtjevirus* and *Cebaduodecimvirus*.

#### Justification

The *Obscuriviridae* family was delineated with two genera. The genus *Omtjevirus* has *Omtjevirus Omtje* as a species and the genus *Cebaduodecimvirus* has *Cebaduodecimvirus phi12duo* and *Cebaduodecimvirus phi12auna* as species. The proposed taxonomic assignments are based on six different methods including nucleotide-, protein-, amino acid- and core protein-based analysis.

**Submitted:** 21 June 24; **Revised:** 30 September 24

**Table 45.** *Obscuriviridae*, 6 new taxa\*

Operation	Rank	New taxon name	Virus name	Exemplar
New taxon	Family	<i>Obscuriviridae</i>		
New taxon	Genus	<i>Omtjevirus</i>		
New taxon	Species	<i>Omtjevirus Omtje</i>	Cellulophaga phage Omtje_1	MT732445
New taxon	Genus	<i>Cebaduodecimvirus</i>		
New taxon	Species	<i>Cebaduodecimvirus phi12duo</i>	Cellulophaga phage phi12:2	KC821606
New taxon	Species	<i>Cebaduodecimvirus phi12auna</i>	Cellulophaga phage phi12a:1	KC821623

\*Source/full text: [https://ictv.global/ictv/proposals/2024.025B.Obscuriviridae\\_1nf\\_2ng\\_3ns.zip](https://ictv.global/ictv/proposals/2024.025B.Obscuriviridae_1nf_2ng_3ns.zip).

## 2024.026B.Pantenvirales\_1no\_3mf

**Title:** Create one new order *Pantenvirales* (*Duplodnaviria*)

**Authors:** Adriaenssens EM (Evelien.adriaenssens@quadram.ac.uk), Cook R, Millard AD, Turner D

### Summary

#### Taxonomic rank(s) affected

Realm: *Duplodnaviria*; kingdom: *Heunggongvira*; phylum: *Uroviricota*; class: *Caudoviricetes*.

#### Description of current taxonomy

The families *Straboviridae*, *Kyanoviridae* and *Ackermannviridae* are unassigned at the order rank within the class *Caudoviricetes*.

#### Proposed taxonomic change(s)

The order *Pantenvirales* is created for these families.

#### Justification

The families *Straboviridae*, *Kyanoviridae* and *Ackermannviridae* form a monophyletic cluster in proteome-based analyses. Members of these families share a number of core orthologous genes.

**Submitted:** 21 June 24

**Table 46.** Pantevenvirales, 3 move taxa\*

Operation	Rank	Taxon name	New parent taxon	Old parent taxon
Move taxon	Family	Straboviridae	Pantevenvirales	Caudoviricetes
Move taxon	Family	Kyanoviridae	Pantevenvirales	Caudoviricetes
Move taxon	Family	Ackermannviridae	Pantevenvirales	Caudoviricetes

**Table 47.** Pantevenvirales, 1 new taxon\*

Operation	Rank	New taxon name
New taxon	Order	Pantevenvirales

\*Source/full text: [https://ictv.global/ictv/proposals/2024.026B.Pantevenvirales\\_1no\\_3mf.zip](https://ictv.global/ictv/proposals/2024.026B.Pantevenvirales_1no_3mf.zip).

## 2024.028B.Philemonvirus\_1ns

**Title:** To create one new species in the genus *Bifilivirus*, family *Paulinoviridae*

**Authors:** Deptula P (deptula@food.ku.dk), Sha Y, Potipimpanon S, Vogensen FK, Nielsen DS, Knezevic P

### Summary

#### Taxonomic rank(s) affected

This is a proposal for creating a new species within the genus *Bifilivirus*.

#### Description of current taxonomy

The genus *Bifilivirus* currently includes one species.

#### Proposed taxonomic change(s)

Addition of a new species *Bifilivirus philemonii*.

#### Justification

Filamentous phage Philemon was isolated from raw milk Emmental cheese on a dairy-associated strain of *Propionibacterium freudenreichii* PB4. The obtained TEM confirmed filamentous morphology. Philemon forms plaques on three *P. freudenreichii* strains in our collection, including the strain TL18, which was used for characterization of phage B5 (though no plaquing was reported for phage B5) [12]. The phage genome was sequenced with the Illumina platform after the formation of the complementary strand with the MDA technique. The genome is 5802 nucleotides, with 63 G+C %mol. Comparison with the only other representative of genus *Bifilivirus*, *Propionibacterium virus B5* (AF428260), revealed that the genome of Philemon is 92.8% identical to the genome of the *Propionibacterium virus B5*, thus fulfilling the criteria for the creation of a separate species. The similarity of morphogenesis protein and CoA B confirms that Philemon belongs to the genus *Bifilivirus*. We propose the name of the species – *Bifilivirus philemonii*.

**Submitted:** 23 April 24

**Table 48.** Philemonvirus, 1 new taxon\*

Operation	Rank	New taxon name	Virus name	Exemplar
New taxon	Species	<i>Bifilivirus philemonii</i>	<i>Propionibacterium phage Philemon</i>	PP693361

\*Source/full text: [https://ictv.global/ictv/proposals/2024.028B.Philemonvirus\\_1ns.zip](https://ictv.global/ictv/proposals/2024.028B.Philemonvirus_1ns.zip).

## 2024.029B.Rhodococcus\_siphoviruses\_7ng\_7ns

**Title:** Create seven new genera for *Rhodococcus* siphoviruses (class: *Caudoviricetes*)

**Authors:** Kurtböke I, Moraru C, Tolstoy I, Kropinski AM (Phage.Canada@gmail.com)

### Summary

#### Taxonomic rank(s) affected

Realm *Duplodnaviria*, kingdom *Heunggongvirae*, phylum *Uroviricota*, class *Caudoviricetes*.

#### Description of current taxonomy

The viruses classified in this proposal do not have a current taxonomic assignment.

#### Proposed taxonomic change(s)

We propose the creation of seven new taxa for unique *Rhodococcus* siphoviruses: *Edwardsroadvirus*, *Wodongavirus*, *Reqipinevirus*, *Melbournevirus*, *Togglehumpervirus*, *Mboduovirus*, *Reynauldvirus*.

#### Justification

The taxa proposed conform to demarcation criteria specified by the ICTV Bacterial Viruses Subcommittee.

**Submitted:** 09 May 2024

**Table 49.** *Rhodococcus* siphoviruses, 14 new taxa\*

Operation	Rank	New taxon name	Virus name	Exemplar
New taxon	Genus	<i>Edwardsroadvirus</i>		
New taxon	Species	<i>Edwardsroadvirus RRH1</i>	Rhodococcus phage RRH1	JN116822
New taxon	Genus	<i>Wodongavirus</i>		
New taxon	Species	<i>Wodongavirus REQ3</i>	Rhodococcus phage REQ3	JN116824
New taxon	Genus	<i>Reqipinevirus</i>		
New taxon	Species	<i>Reqipinevirus reqipine5</i>	Rhodococcus phage ReqiPine5	GU580943
New taxon	Genus	<i>Melbournevirus</i>		
New taxon	Species	<i>Melbournevirus REQ2</i>	Rhodococcus phage REQ2	JN116823
New taxon	Genus	<i>Togglehumpervirus</i>		
New taxon	Species	<i>Togglehumpervirus togglehumper</i>	Rhodococcus phage Togglehumper	OQ709222
New taxon	Genus	<i>Mboduovirus</i>		
New taxon	Species	<i>Mboduovirus mbo2</i>	Rhodococcus phage Mbo2	ON191531
New taxon	Genus	<i>Reynauldvirus</i>		
New taxon	Species	<i>Reynauldvirus reynaud</i>	Rhodococcus phage Reynaud	OR159659

\*Source/full text: [https://ictv.global/ictv/proposals/2024.029B.Rhodococcus\\_siphoviruses\\_7ng\\_7ns.zip](https://ictv.global/ictv/proposals/2024.029B.Rhodococcus_siphoviruses_7ng_7ns.zip).

## 2024.030B.Trautnerviridae\_1nf\_1nsf\_3ng\_6ns

**Title:** Create a new family, *Trautnerviridae*, subfamily *Polsinellivirinae* and two genera (*Rivavivirus*, and *Splendidredvirus*, class *Caudoviricetes*)

**Authors:** Cook R, Tavares P, Lurz R, Barylski J, Moraru C, Tolstoy I, Kropinski AM (Phage.Canada@gmail.com)

#### Summary

##### Taxonomic rank(s) affected

Realm *Duplodnaviria*, kingdom *Heunggongvirae*, phylum *Uroviricota*, class *Caudoviricetes*.

##### Description of current taxonomy

The viruses classified in this proposal do not have a current taxonomic assignment.

##### Proposed taxonomic change(s)

To create a new genus, *Rivavivirus*, with three species.

To create a new genus, *Splendidredvirus*, with two species.

To create a new subfamily, *Polsinellivirinae*, with these two genera (*Rivavivirus* and *Splendidredvirus*).

To create a new single-species genus, *Prospekt naukivivirus*.

To create a new family, *Trautnerviridae*, for these taxa.

#### Justification

Bacillus phage SPP1 was isolated in 1966; sequenced in 1997 (corrected in 2018); and has been the subject of numerous morphological and physiological studies; yet has remained unclassified. In this proposal, it has been assigned to a new genus, *Rivavivirus*, together with Bacillus phage vB\_BspS\_SplendidRed (assigned to the new genus *Splendidredvirus*), to form a new subfamily, *Polsinellivirinae*. The members of this taxon are siphoviruses which have genomes of 42.8–46.3 kb (43.7–44.6 mol% G+C) and encode 74–77 proteins and no tRNAs. As a result of detailed genomic, proteomic and phylogenetic analyses using VIRIDIC, ViPTree and VirClust, we further propose to create a new family for these genera named *Trautnerviridae* named in honour of Thomas A. Trautner.

**Submitted:** 25 May 24; **Revised:** 30 September 24

**Table 50.** Trautnerviridae, 11 new taxa\*

Operation	Rank	New taxon name	Virus name	Exemplar
New taxon	Family	<i>Trautnerviridae</i>		
New taxon	Subfamily	<i>Polsinellivirinae</i>		
New taxon	Genus	<i>Rivavirus</i>		
New taxon	Species	<i>Rivavirus SPP1</i>	Bacillus phage SPP1	X97918
New taxon	Species	<i>Rivavirus rv000TH010</i>	Bacillus phage 000TH010	MN176219
New taxon	Species	<i>Rivavirus rv049ML001</i>	Bacillus phage 049 ml001	MN176227
New taxon	Genus	<i>Splendidredvirus</i>		
New taxon	Species	<i>Splendidredvirus splendidred</i>	Bacillus phage vB_BspS_SplendidRed	MN013088
New taxon	Species	<i>Splendidredvirus ray17</i>	Bacillus phage Ray17	MH752385
New taxon	Genus	<i>Prospektaukivirus</i>		
New taxon	Species	<i>Prospektaukivirus sam112</i>	Bacillus phage vB_BcM_Sam112	MN604230

\*Source/full text: [https://ictv.global/ictv/proposals/2024.030B.Trautnerviridae\\_1nf\\_1nsf\\_3ng\\_6ns.zip](https://ictv.global/ictv/proposals/2024.030B.Trautnerviridae_1nf_1nsf_3ng_6ns.zip).

## 2024.031B.Sarkviridae\_1nf\_1msf\_2mg

**Title:** Create a new family, *Sarkviridae* for the Jersey-like siphophages (class: *Caudoviricetes*)

**Authors:** Moraru C, Tolstoy I, Kropinski AM (Phage.Canada@gmail.com)

### Summary

#### Taxonomic rank(s) affected

Realm *Duplodnaviria*, kingdom *Heunggongvirae*, phylum *Uroviricota*, class *Caudoviricetes*.

#### Description of current taxonomy

The genera *Jerseyvirus*, *Cornellvirus* and *Kagunavirus* in the subfamily *Guernseyvirinae* and among the *Serratia* phages the genera *Seretavirus* and *Otakuvirus*.

#### Proposed taxonomic change(s)

To create a new family *Sarkviridae*.

#### Justification

Genomic, proteomic and phylogenetic data indicate that this group of phages is a family. In addition, Taxonomy Proposal 2023.068B.Caudoviricetes\_Serratia\_3ng suggested a higher level relationship between the taxa *Otakuvirus* and *Guernseyvirinae*.

**Submitted:** 04 June 24; **Revised:** 30 September 24

**Table 51.** *Sarkviridae*, 3 move taxa\*

Operation	Rank	Taxon name	New parent taxon	Old parent taxon
Move taxon	Subfamily	<i>Guernseyvirinae</i>	<i>Sarkviridae</i>	<i>Caudoviricetes</i>
Move taxon	Genus	<i>Seretavirus</i>	<i>Sarkviridae</i>	<i>Caudoviricetes</i>
Move taxon	Genus	<i>Otakuvirus</i>	<i>Sarkviridae</i>	<i>Caudoviricetes</i>

**Table 52.** *Sarkviridae*, 1 new taxon\*

Operation	Rank	New taxon name
New taxon	Family	<i>Sarkviridae</i>

\*Source/full text: [https://ictv.global/ictv/proposals/2024.031B.Sarkviridae\\_1nf\\_1msf\\_2mg.zip](https://ictv.global/ictv/proposals/2024.031B.Sarkviridae_1nf_1msf_2mg.zip).

## 2024.032B.Sepahanvirus\_1ng\_2ns

**Title:** Create a new genus, *Sepahanvirus*, containing two species (*Caudoviricetes*)

**Authors:** Ganjoor MS, Bouzari M (bouzari@sci.ui.ac.ir), Soleimani-Delfan A

### Summary

#### Taxonomic rank(s) affected

Genus.

#### Description of current taxonomy

These phages are currently unclassified.

#### Proposed taxonomic change(s)

To create a new genus, *Sepahanvirus*, within the class *Caudoviricetes* comprising two species, for Yersinia phage vB\_Yru\_GN1 and Yersinia phage YerA41.

#### Justification

Yersinia phage vB\_Yru\_GN1 and Yersinia phage YerA41 exhibit nucleotide sequence similarity that falls within the demarcation threshold for the creation of a new genus.

**Submitted:** 09 December 23

**Table 53.** *Sepahanvirus*, 3 new taxa\*

Operation	Rank	New taxon name	Virus name	Exemplar
New taxon	Genus	<i>Sepahanvirus</i>		
New taxon	Species	<i>Sepahanvirus GN1</i>	Yersinia phage vB_Yru_GN1	LC779065
New taxon	Species	<i>Sepahanvirus yerA41</i>	Yersinia phage YerA41	MW570730

\*Source/full text: [https://ictv.global/ictv/proposals/2024.032B.Sepahanvirus\\_1ng\\_2ns.zip](https://ictv.global/ictv/proposals/2024.032B.Sepahanvirus_1ng_2ns.zip).

## 2024.033B.Mazoviaviridae\_1nf\_1ng\_1ns

**Title:** Create a new family, *Mazoviaviridae*, and a new genus, *Dabrowskivirus*, with a single species (class *Caudoviricetes*)

**Authors:** Shymialevich D, Wójcicki M (michal.wojcicki@hirsfeld.pl), Sokołowska B

### Summary

#### Taxonomic rank(s) affected

Proposal to create a new family, *Mazoviaviridae*, and a new genus, *Dabrowskivirus*, with a single species (class *Caudoviricetes*).

#### Description of current taxonomy

In 2022, significant changes to the taxonomy of bacterial viruses were introduced: the paraphyletic morphological families *Podoviridae*, *Siphoviridae* and *Myoviridae* as well as the order *Caudovirales* were abolished, which is replaced by the class *Caudoviricetes* to group all tailed bacterial and archaeal viruses with icosahedral capsids and a double-stranded DNA genome. Moreover, a binomial system of nomenclature for species was established. Based on the morphology and the comparative analysis of its predicted proteins, *Alicyclobacillus myophage vB\_Aac\_IAFB\_3916* was assigned to viruses with complex structures (class *Caudoviricetes*).

#### Proposed taxonomic change(s)

Analyses of the phylogenetic relationship of *Alicyclobacillus myophage vB\_Aac\_IAFB\_3916* were unable to provide an unambiguous assignment to a specific family and genus. The weak similarity with other phage genomes deposited in the databases suggests that the isolated bacteriophage may be a representative of a new genus and new family of tailed bacteriophages.

#### Justification

The genome of newly isolated *Alicyclobacillus myophage vB\_Aac\_IAFB\_3916* possesses no DNA homologues. At the protein level, this virus is unique. Therefore, we propose the creation of a new species (*Dabrowskivirus KKP3916*), genus (*Dabrowski-virus*) and family (*Mazoviaviridae*) for viruses of this type.

**Submitted:** 09 June 24; **Revised:** 09 October 24

**Table 54.** *Mazoviaviridae*, 3 new taxa\*

Operation	Rank	New taxon name	Virus name	Exemplar
New taxon	Family	<i>Mazoviaviridae</i>		
New taxon	Genus	<i>Dabrowskivirus</i>		
New taxon	Species	<i>Dabrowskivirus KKP3916</i>	Alicyclobacillus myophage vB_Aac_IAFB_3916	OQ846916

\*Source/full text: [https://ictv.global/ictv/proposals/2024.033B.Mazoviaviridae\\_1nf\\_1ng\\_1ns.zip](https://ictv.global/ictv/proposals/2024.033B.Mazoviaviridae_1nf_1ng_1ns.zip).

## 2024.034B.*Stackebrandtviridae*\_1nf\_2nsf\_8mg\_8ns

**Title:** Create a new family, *Stackebrandtviridae*, for a group of *Gordonia* phages (class: *Caudoviricetes*)

**Authors:** Kurtbøke I, Moraru C, Tolstoy I, Kropinski AM (Phage.Canada@gmail.com)

### Summary

#### Taxonomic rank(s) affected

Realm *Duplodnaviria*, kingdom *Heunggongvirae*, phylum *Uroviricota*, class *Caudoviricetes*.

#### Description of current taxonomy

At present, the following taxa exist as floating genera in the class *Caudoviricetes*; *Wizardvirus*, *Clownvirus*, *Vividuovirus*, *Dexdertvirus*, *Zitchvirus*, *Kroosvirus* and *Leonardvirus*.

#### Proposed taxonomic change(s)

To create one new species in the genus *Wizardvirus*.

To create a new subfamily, *Frickvirinae* with two genera (*Clownvirus* and *Wizardvirus*).

To add one new species to the genus *Vividuovirus*.

To add one new species to the genus *Dexdertvirus*.

To add four new species to the genus *Zitchvirus*.

To add one new species to the genus *Leonardvirus*.

To create a new subfamily, *Schenleyvirinae*, for the above four genera and *Kroosvirus*.

To create a new family, *Stackebrandtviridae*, for the above-mentioned taxa.

#### Justification

Members of the Actinobacteriophage Database Cluster DC (<https://phagesdb.org/clusters/DC/>) are temperate *Gordonia* phages for which we have created two genera. The related lytic viruses of Cluster DE (<https://phagesdb.org/clusters/DE/>) have previously been placed in five genera.

**Submitted:** 15 June 24; **Revised:** 30 September 24

**Table 55.** *Stackebrandtviridae*, 11 new taxa\*

Operation	Rank	New taxon name	Virus name	Exemplar
New taxon	Family	<i>Stackebrandtviridae</i>		
New taxon	Subfamily	<i>Frickvirinae</i>		
New taxon	Species	<i>Wizardvirus halo3</i>	<i>Gordonia</i> phage Halo3	OR521081
New taxon	Subfamily	<i>Schenleyvirinae</i>		
New taxon	Species	<i>Vividuovirus sitar</i>	<i>Gordonia</i> phage Sitar	MH153809
New taxon	Species	<i>Dexdertvirus kwekel</i>	<i>Gordonia</i> phage Kwekel	OR521074
New taxon	Species	<i>Zitchvirus tardus</i>	<i>Gordonia</i> phage Tardus	ON392159
New taxon	Species	<i>Zitchvirus viaconlectus</i>	<i>Gordonia</i> phage ViaConlectus	OP068342
New taxon	Species	<i>Zitchvirus sampson</i>	<i>Gordonia</i> phage Sampson	ON456337
New taxon	Species	<i>Zitchvirus apunk</i>	<i>Gordonia</i> phage APunk	ON755186
New taxon	Species	<i>Leonardvirus phauci</i>	<i>Gordonia</i> phage Phauci	ON456349

**Table 56.** Stackebrandtviridae, 8 move taxa\*

Operation	Rank	Taxon name	New parent taxon	Old parent taxon
Move taxon	Genus	<i>Clownvirus</i>	<i>Frickvirinae</i>	<i>Caudoviricetes</i>
Move taxon	Genus	<i>Wizardvirus</i>	<i>Stackebrandtviridae</i>	<i>Caudoviricetes</i>
Move taxon	Genus	<i>Kroosvirus</i>	<i>Schenleyvirinae</i>	<i>Caudoviricetes</i>
Move taxon	Genus	<i>Vividuovirus</i>	<i>Schenleyvirinae</i>	<i>Caudoviricetes</i>
Move taxon	Genus	<i>Dexdertvirus</i>	<i>Schenleyvirinae</i>	<i>Caudoviricetes</i>
Move taxon	Genus	<i>Zitchvirus</i>	<i>Schenleyvirinae</i>	<i>Caudoviricetes</i>
Move taxon	Genus	<i>Leonardvirus</i>	<i>Schenleyvirinae</i>	<i>Caudoviricetes</i>
Move taxon	Genus	<i>Lilbeanievirus</i>	<i>Stackebrandtviridae</i>	<i>Caudoviricetes</i>

\*Source/full text: [https://ictv.global/ictv/proposals/2024.034B.Stackebrandtviridae\\_1nf\\_2nsf\\_8mg\\_8ns.zip](https://ictv.global/ictv/proposals/2024.034B.Stackebrandtviridae_1nf_2nsf_8mg_8ns.zip).

## 2024.036B.Caudoviricetes\_Faserviricetes\_Name\_Corrections

**Title:** Corrections to species names in the classes *Caudoviricetes* and *Faserviricetes*

**Authors:** Turner D (dann2.turner@uwe.ac.uk)

### Summary

#### Taxonomic rank(s) affected

Genera and species in realm *Duplodnaviria*, kingdom *Heunggongvirae*, phylum *Uroviricota*, class *Caudoviricetes*.

#### Description of current taxonomy

All of these species are currently classified within the latest release of the ICTV taxonomy.

#### Proposed taxonomic change(s)

Rename species to conform to the binomial species epithet.

Correct spelling errors.

Create genus to contain a floating species in the subfamily *Tevenvirinae*.

Change to genus name and constituent species in the genus *Roskildevirus* as ‘Roskilde virus’ refers to Norovirus in Danish.

Correction of misspelt genus names in binomial species epithets.

Move two species that were incorrectly classified within the genus *Tequattrovirus*.

Abolish species *Campylobacter virus IBB35*.

#### Justification

To ensure that the naming of viruses is consistent with the guidelines for binomial species names, to remove any names that could cause confusion in native languages and to remove any genome records that do not represent coding complete sequences.

**Submitted:** 19 June 24; **Revised:** 30 September 24

**Table 57.** Caudoviricetes, 96 rename taxa\*. Table too large, see supplementary information sheet supp\_info\_tab\_57**Table 58.** Caudoviricetes, 3 move; rename taxa\*

Operation	Rank	New taxon name	Old parent taxon	New parent taxon	Old taxon name
Move; rename taxon	Species	<i>Centumtrigintavivirus cv133</i>	<i>Tevenvirinae</i>	<i>Centumtrigintavivirus</i>	<i>Acinetobacter virus 133</i>
Move; rename taxon	Species	<i>Mosigvirus jaykay</i>	<i>Tequattrovirus</i>	<i>Mosigvirus</i>	<i>Tequattrovirus jaykay</i>
Move; rename taxon	Species	<i>Mosigvirus efftwo</i>	<i>Tequattrovirus</i>	<i>Mosigvirus</i>	<i>Tequattrovirus efftwo</i>

**Table 59.** Caudoviricetes, 1 abolish taxon\*

Operation	Rank	Abolished taxon name
Abolish taxon	Species	<i>Campylobacter virus IBB35</i>

**Table 60.** Caudoviricetes, 1 new taxon\*

Operation	Rank	New taxon name
New taxon	Genus	<i>Centumtrigintavirüs</i>

\*Source/full text: [https://ictv.global/ictv/proposals/2024.036B.Caudoviricetes\\_Faserviricetes\\_Name\\_Corrections.zip](https://ictv.global/ictv/proposals/2024.036B.Caudoviricetes_Faserviricetes_Name_Corrections.zip).

## 2024.037B.Vandenendeviridae\_1nf\_2msf\_8ng\_1mg\_11ns

**Title:** Create a new family, *Vandenendeviridae*, for a group of lytic *Pseudomonas* phages (class: *Caudoviricetes*)

**Authors:** Moraru C, Tolstoy I, Kropinski AM (Phage.Canada@gmail.com)

### Summary

#### Taxonomic rank(s) affected

Realm *Duplodnaviria*, kingdom *Heunggongvirae*, phylum *Uroviricota*, class *Caudoviricetes*.

#### Description of current taxonomy

Five genera are currently classified; *Baldwinivirus*, *Nankokuvirus*, *Otagovirus*, *Flaumdravivirus* and *Pakpunavirus*.

#### Proposed taxonomic change(s)

To create seven new single-species genera: *Weillhallivirus*, *Omahavirus*, *Torinovirus*, *Yunamivivirus*, *Ventosusvirus*, *Uavernvirus* and *Chemalvirus*.

To create a new genus, *Tartuvirus*, with four species.

To create two new species in the genus *Kremarvirus*.

To create a new family, *Vandenendeviridae*, for these genera and *Baldwinivirus*, *Kremarvirus*, *Nankokuvirus*, *Otagovirus*, *Flaumdravivirus*, *Pakpunavirus* and *Shenlongvirus*.

#### Justification

Using VIRIDIC, ViPTree, VIRCLUST and vConTACT v.3.0, we have established that this is a cohesive group of lytic *Pseudomonas* myoviruses which share ≥12.2% DNA sequence similarity and 15 core proteins. The new family is named in honour of Marius van de Ende of South Africa.

**Submitted:** 25 May 24; **Revised:** 30 September 24

**Table 61.** *Vandenendeviridae*, 20 new taxa\*

Operation	Rank	New taxon name	Virus name	Exemplar
New taxon	Family	<i>Vandenendeviridae</i>		
New taxon	Genus	<i>Weillhallivirus</i>		
New taxon	Species	<i>Weillhallivirus wv16Q</i>	Pseudomonas phage 16Q	OR001909
New taxon	Genus	<i>Omahavirus</i>		
New taxon	Species	<i>Omahavirus UNOG1W1</i>	Pseudomonas phage UNO-G1W1	PP551948
New taxon	Genus	<i>Torinovirus</i>		
New taxon	Species	<i>Torinovirus K7A1</i>	Pseudomonas phage phiK7A1	MT740307
New taxon	Genus	<i>Yunamivivirus</i>		
New taxon	Species	<i>Yunamivivirus Y1MI</i>	Pseudomonas phage vB_PF_Y1-MI	OR500437
New taxon	Genus	<i>Ventosusvirus</i>		
New taxon	Species	<i>Ventosusvirus ventosus</i>	Pseudomonas phage ventosus	MG018930
New taxon	Genus	<i>Uavernvirus</i>		
New taxon	Species	<i>Uavernvirus uavern</i>	Pseudomonas phage UAVERN	MZ605293
New taxon	Genus	<i>Tartuvirus</i>		
New taxon	Species	<i>Tartuvirus amme3</i>	Pseudomonas phage vB_PpuM-Amme-3	PP496413
New taxon	Species	<i>Tartuvirus nopa</i>	Pseudomonas phage vB_PpuM-NoPa	PP496415
New taxon	Species	<i>Tartuvirus kopa4</i>	Pseudomonas phage vB_PpuM-KoPa-4	PP496414

Continued

**Table 61.** Continued

Operation	Rank	New taxon name	Virus name	Exemplar
New taxon	Species	<i>Tartuvirus roomu2</i>	Pseudomonas phage vB_PpuM-Roomu-2	PP496417
New taxon	Genus	<i>Chemalvirus</i>		
New taxon	Species	<i>Chemalvirus PseuGes254</i>	Pseudomonas phage PseuGes_254	OR575930

**Table 62.** *Vandenendeviridae*, 3 move taxa\*

Operation	Rank	Taxon name	New parent taxon	Old parent taxon
Move taxon	Subfamily	<i>Skurskavirinae</i>	<i>Vandenendeviridae</i>	<i>Caudoviricetes</i>
Move taxon	Subfamily	<i>Gorskivirinae</i>	<i>Vandenendeviridae</i>	<i>Caudoviricetes</i>
Move taxon	Genus	<i>Nankokuvirus</i>	<i>Vandenendeviridae</i>	<i>Caudoviricetes</i>

\*Source/full text: [https://ictv.global/ictv/proposals/2024.037B.Vandenendeviridae\\_1nf\\_2msf\\_8ng\\_1mg\\_11ns.zip](https://ictv.global/ictv/proposals/2024.037B.Vandenendeviridae_1nf_2msf_8ng_1mg_11ns.zip).

## 2024.038B.Vinavirales\_3nf\_1mf\_7ng\_5ns\_3ms

**Title:** Create three new families *Mestraviridae*, *Asemoviridae* and *Parnassusviridae*, and move the family *Autolykiviridae* into the order *Vinavirales* (*Tectiliviricetes*, *Preplasmiviricota*, *Bamfordvirae*, *Varidnaviria*)

**Authors:** Bardy P, Fogg PCM, Kalatzis PG, Middelboe M, Oksanen HM (hanna.oksanen@helsinki.fi)

### Summary

#### Taxonomic rank(s) affected

The taxonomic ranks affected are the genus *Corticovirus* in the family *Corticoviridae*, the order *Vinavirales* and the family *Autolykiviridae*. In addition, some species are affected.

#### Description of current taxonomy

Currently, the order *Vinavirales* includes one family *Corticoviridae* with one genus *Corticovirus* (two species). The family *Autolykiviridae* belongs to the class *Tectiliviricetes* but is not assigned to any order. The family *Autolykiviridae* includes two genera *Livvievirus* (two species) and *Paulavirus* (three species). The realm *Varidnaviria* to which these taxa are classified has been reorganized in proposal 2024.010D.

#### Proposed taxonomic change(s)

The genus *Corticovirus* (family *Corticoviridae*) is renamed as the genus *Merivirus* and its two species are renamed accordingly. The *Vinavirales* order is rearranged so that the family *Autolykiviridae* is placed in the order along with the family *Corticoviridae*. Two new genera, *Oliviavirus* and *Ameliavirus*, are created in the family *Autolykiviridae*, each with a single species, *Oliviavirus viph1020o* and *Ameliavirus viph1008o*, respectively. The genus of the virus species *Paulavirus viph1044o* is moved from *Livvievirus* to *Paulavirus* (family *Autolykiviridae*). In addition, three new families *Mestraviridae*, *Asemoviridae* and *Parnassusviridae* are created in the order *Vinavirales*. Two new genera, *Anticleavirus* and *Polymedevirus*, are created in the family *Mestraviridae*. One new species is created in each of two genera: *Anticleavirus jorvik* and *Polymedevirus YY*. Two new genera, *Elsinorevirus* and *Rumoivirus*, are created in the family *Asemoviridae*. One new species is created in each of two genera: *Elsinorevirus NO16* and *Rumoivirus VruC*. One new species *Corycianvirus MfV* is created in a new genus *Corycianvirus* in the family *Parnassusviridae*.

#### Justification

The inclusion/creation of four families in the order *Vinavirales*, together with the family *Corticoviridae*, is based on the nine signature genes shared by their members, which corresponds to approximately 50% of their genome. This demonstrates their common evolutionary origin. Classification of *Paulavirus viph1044o* into the genus *Livvievirus* instead of *Paulavirus* was a mistake. To make the names of the family *Corticoviridae* and the genus *Corticovirus* based on a different word stem, the genus *Corticovirus* is renamed *Merivirus*.

**Submitted:** 20 June 24; **Revised:** 30 September 24

**Table 63.** Vinavirales, 15 new taxa\*

Operation	Rank	New taxon name	Virus name	Exemplar
New taxon	Genus	<i>Olivaviirus</i>		
New taxon	Genus	<i>Ameliavirus</i>		
New taxon	Family	<i>Mestraviridae</i>		
New taxon	Genus	<i>Anticleavivirus</i>		
New taxon	Species	<i>Anticleavivirus jorvik</i>	Rhodobacter phage Jorvik	OP588643
New taxon	Genus	<i>Polymedevirus</i>		
New taxon	Species	<i>Polymedevirus YY</i>	Marinomonas phage YY	MH105080
New taxon	Family	<i>Asemoviridae</i>		
New taxon	Genus	<i>Elsinorevirus</i>		
New taxon	Species	<i>Elsinorevirus NO16</i>	Vibrio phage fNo16	MH730557
New taxon	Genus	<i>Rumoivirus</i>		
New taxon	Species	<i>Rumoivirus VruC</i>	Vibrio phage vB_VruC_PG21	OM867525
New taxon	Family	<i>Parnassusviridae</i>		
New taxon	Genus	<i>Corycianvirus</i>		
New taxon	Species	<i>Corycianvirus MfV</i>	Marinomonas phage MfV	MW618650

**Table 64.** Vinavirales, 2 move taxa and 2 move rename taxa\*

Operation	Rank	Taxon name	New parent taxon	Old parent taxon	Old taxon name
Move taxon	Family	<i>Autolykiviridae</i>	Vinavirales	Tectiliviricetes	
Move taxon	Species	<i>Paulavirus viph1044o</i>	Paulavirus	Livvievirus	
Move; rename taxon	Species	<i>Olivaviirus viph1020o</i>	Olivaviirus	Paulavirus	<i>Paulavirus viph1020o</i>
Move; rename taxon	Species	<i>Ameliavirus viph1008o</i>	Ameliavirus	Paulavirus	<i>Paulavirus viph1008o</i>

**Table 65.** Vinavirales, 3 rename taxa\*

Operation	Rank	New taxon name	Previous taxon name
Rename taxon	Genus	<i>Merivirus</i>	<i>Corticovirus</i>
Rename taxon	Species	<i>Merivirus Cr39582</i>	<i>Corticovirus Cr39582</i>
Rename taxon	Species	<i>Merivirus PM2</i>	<i>Corticovirus PM2</i>

\*Source/full text: [https://ictv.global/ictv/proposals/2024.038B.Vinavirales\\_3nf\\_1mf\\_7ng\\_5ns\\_3ms.zip](https://ictv.global/ictv/proposals/2024.038B.Vinavirales_3nf_1mf_7ng_5ns_3ms.zip).

## 2024.039B.Artimaviricota\_np

**Title:** Create new phylum, *Artimaviricota* in the kingdom *Orthornavirae* (realm *Riboviria*) for classification of a hyperthermophilic RNA virus

**Authors:** Syun-ichi Urayama (urayama.shunichi.gn@u.tsukuba.ac.jp), Akihito Fukudome, Eugene V. Koonin, Takuro Nunoura, Mart Krupovic (mart.krupovic@pasteur.fr)

### Summary

#### Taxonomic rank(s) affected

*Riboviria*, *Orthornavirae*.

#### Description of current taxonomy

Realm *Riboviria* includes two kingdoms, *Orthornavirae* and *Pararnavirae*, which include highly diverse viruses that encode RNA-directed RNA polymerases (RdRP) and reverse transcriptases (RT), respectively. Kingdom *Orthornavirae* includes six

phyla which were established based on phylogenetic analysis of the RdRP and comparative analysis of the viral genomes and proteins.

#### **Proposed taxonomic change(s)**

We propose to create a new phylum in the kingdom *Orthornavirae* for classification of a group of RNA viruses discovered in hot springs that are characterized by unusual RdRPs.

#### **Justification**

The RdRPs of hot spring RNA virus 1 and its relatives seem to deviate from the RdRP consensus farther than any of the other recently discovered putative phyla, with none of which they appear to be affiliated, and possess unusual structural features that appear to link them to viral RTs.

**Submitted:** 21 June 2024

**Table 66.** *Artimaviricota*, 6 new taxa\*

Operation	Rank	New taxon name	Virus name	Exemplar
New taxon	Phylum	<i>Artimaviricota</i>		
New taxon	Class	<i>Furtirnaviricetes</i>		
New taxon	Order	<i>Divaquavirales</i>		
New taxon	Family	<i>Hakuzoviridae</i>		
New taxon	Genus	<i>Atsuirnavirus</i>		
New taxon	Species	<i>Atsuirnavirus caloris</i>	hot spring RNA virus 1	RNA1: BTCN01000001; RNA2: BTCN01000005

\*Source/full text: [https://ictv.global/ictv/proposals/2024.039B.Artimaviricota\\_np.zip](https://ictv.global/ictv/proposals/2024.039B.Artimaviricota_np.zip).

## **2024.040B.Sharonstreetvirus\_1ns**

**Title:** Create a new species *Sharonstreetvirus xiamensis* (*Caudoviricetes*)

**Author:** Liu H-T (15844493757@163.com)

#### **Summary**

##### **Taxonomic rank(s) affected**

Species.

##### **Description of current taxonomy**

Not currently classified.

##### **Proposed taxonomic change(s)**

Create a new species *Sharonstreetvirus xiamensis* for phage phiA034 (OP792756) isolated from aquaculture water in Xiamen.

#### **Justification**

ViPTree analysis classifies phiA034 as a new species the genus *Sharonstreetvirus*.

**Submitted:** 25 April 24

**Table 67.** *Sharonstreetvirus*, 1 new taxon\*

Operation	Rank	New taxon name	Virus name	Exemplar
New taxon	Species	<i>Sharonstreetvirus xiamensis</i>	Aeromonas phage phiA034	OP792756

\*Source/full text: [https://ictv.global/ictv/proposals/2024.040B.Sharonstreetvirus\\_1ns.zip](https://ictv.global/ictv/proposals/2024.040B.Sharonstreetvirus_1ns.zip).

## **2024.041B.Camvirus\_2ns**

**Title:** Create two new species – *Camvirus vanseggelen* and *Camvirus verabelle* (subfamily *Arquatrovirinae*, class *Caudoviricetes*).

**Authors:** Kempff A (annabelkempff@gmail.com), van Neer V (vanneervera@gmail.com), Ongenae V (v.m.a.ongenae@biology.leidenuniv.nl), Rozen DE, Briegel A, Claessen D

## Summary

### Taxonomic rank(s) affected

Realm *Duplodnaviria*, kingdom *Heunggongvirae*, phylum *Uroviricota*, class *Caudoviricetes*.

### Description of current taxonomy

The genus *Camvirus* is classified within the subfamily *Arquattrovirinae*, class *Caudoviricetes*.

### Proposed taxonomic change(s)

Create two new species in the genus *Camvirus*.

### Justification

We have defined two new species, *Camvirus vanseggelen* and *Camvirus verabelle*, based upon phages isolated for host *Streptomyces coelicolor* from soil samples in the Netherlands at longitude N52°23'31" and latitude E4°34'49". *Camvirus vanseggelen* contains a linear dsDNA genome of 50 426 bp (65.5% G+C) encoding 73 proteins. *Camvirus verabelle* contains a linear dsDNA genome of 49 832 bp (65.0% G+C) encoding 73 proteins. These new species can be included in the genus *Camvirus* together with Streptomyces phage Alsaber, Streptomyces phage Amela, Streptomyces phage phiCAM, Streptomyces phage Endor1, Streptomyces phage Endor2, Streptomyces phage Joe, Streptomyces phage Saftant and Streptomyces phage Sitrop.

**Submitted:** 02 April 24

**Table 68.** *Camvirus*, 2 new taxa\*

Operation	Rank	New taxon name	Virus name	Exemplar
New taxon	Species	<i>Camvirus vanseggelen</i>	Streptomyces phage Vanseggelen	OQ970438
New taxon	Species	<i>Camvirus verabelle</i>	Streptomyces phage Verabelle	OQ970439

\*Source/full text: [https://ictv.global/ictv/proposals/2024.041B.Camvirus\\_2ns.zip](https://ictv.global/ictv/proposals/2024.041B.Camvirus_2ns.zip).

## 2024.042B.Lacfervirus\_1ng\_1ns

**Title:** Create one new genus, *Lacfervirus*, in the class *Caudoviricetes*

**Authors:** Qiannan Wen, Xia Chen (chenxia8280@163.com)

## Summary

### Taxonomic rank(s) affected

Realm *Duplodnaviria*, kingdom *Heunggongvirae*, phylum *Uroviricota*, class *Caudoviricetes*.

### Description of current taxonomy

The virus classified in this proposal does not have a current taxonomic assignment.

### Proposed taxonomic change(s)

We propose a new genus, *Lacfervirus*, including a single species for the virus Lactobacillus phage LFP01 in the class *Caudoviricetes*.

### Justification

The genome of Lactobacillus phage LFP01 has very low homology to all other bacteriophage sequences in the GenBank database. A complete genome sequence comparison using the BLASTn method revealed that the genome of Lactobacillus phage LFP01 had a maximum nucleotide identity of 92.83 and 55% coverage with Lactobacillus phage LF1. The low homology indicated that the newly sequenced bacteriophage likely represents a new genus with a single species.

**Submitted:** 05 June 23

**Table 69.** *Lacfervirus*, 2 new taxa\*

Operation	Rank	New taxon name	Virus name	Exemplar
New taxon	Genus	<i>Lacfervirus</i>		
New taxon	Species	<i>Lacfervirus LFP01</i>	Lactobacillus virus LFP01	OR048821

\*Source/full text: [https://ictv.global/ictv/proposals/2024.042B.Lacfervirus\\_1ng\\_1ns.zip](https://ictv.global/ictv/proposals/2024.042B.Lacfervirus_1ng_1ns.zip).

**2024.043B.Cystoviridae\_6ng\_2nsp\_1rng\_7rnsp**

**Title:** Rename and split an existing genus of the family *Cystoviridae* (*Vidaverviricetes: Mindivirales*), rename seven virus species, create two new species and genera

**Authors:** Poranen MM (minna.poranen@helsinki.fi), Mäntynen S

**Summary****Taxonomic rank(s) affected**

The proposal affects species and genus ranks under the family *Cystoviridae*.

**Description of current taxonomy**

The family *Cystoviridae* currently includes one genus *Cystovirus* and seven species, *Cystovirus phi6*, *Cystovirus phi8*, *Cystovirus phi12*, *Cystovirus phi13*, *Cystovirus phi2954*, *Cystovirus phiNN* and *Cystovirus phiYY*. *Cystoviridae* is the only family of the order *Mindivirales* and the class *Vidaverviricetes* that belongs to the phylum *Duplornaviricota* (*Orthornavirae*, *Riboviria*). The other classes in this phylum are *Resentoviricetes* and *Chrymotiviricetes*.

**Proposed taxonomic change(s)**

We propose a new name for the genus *Cystovirus* and its splitting into five genera. Due to the introduction of the new genera, we propose the renaming of all the current species. In addition, we propose to create two new species and two additional new genera in the family *Cystoviridae*.

**Justification**

Seven new dsRNA bacteriophage isolates have been identified and are now proposed to be taxonomically classified to create two new species. Sequence comparisons of these viruses and previously classified dsRNA bacteriophages of the genus *Cystovirus* justify the splitting of the genus *Cystovirus* and the creation of seven genera in the family *Cystoviridae*. The genus *Cystovirus* is renamed to distinguish the name stems of genus and family rank.

**Submitted:** 21 June 24; **Revised:** 30 September 24

**Table 70.** *Cystoviridae*, 5 move; rename taxa\*

Operation	Rank	New taxon name	Old parent taxon	New parent taxon	Old taxon name
Move; rename taxon	Species	<i>Alphacystovirus phi8</i>	<i>Cystovirus</i>	<i>Alphacystovirus</i>	<i>Cystovirus phi8</i>
Move; rename taxon	Species	<i>Betacystovirus phi12</i>	<i>Cystovirus</i>	<i>Betacystovirus</i>	<i>Cystovirus phi12</i>
Move; rename taxon	Species	<i>Gammacystovirus phi13</i>	<i>Cystovirus</i>	<i>Gammacystovirus</i>	<i>Cystovirus phi13</i>
Move; rename taxon	Species	<i>Gammacystovirus phiYY</i>	<i>Cystovirus</i>	<i>Gammacystovirus</i>	<i>Cystovirus phiYY</i>
Move; rename taxon	Species	<i>Deltacystovirus phi2954</i>	<i>Cystovirus</i>	<i>Deltacystovirus</i>	<i>Cystovirus phi2954</i>

**Table 71.** *Cystoviridae*, 4 new taxa\*

Operation	Rank	New taxon name	Virus name	Exemplar
New taxon	Genus	<i>Epsiloncystovirus</i>		
New taxon	Species	<i>Epsiloncystovirus phiNY</i>	Microvirgula phage phiNY	MW471133; MW471134; MW471135
New taxon	Genus	<i>Zetacystovirus</i>		
New taxon	Species	<i>Zetacystovirus CAP</i>	Acinetobacter phage CAP7	MZ558516; MZ558517; MZ558518

**Table 72.** *Cystoviridae*, 3 rename taxa\*

Operation	Rank	New taxon name	Previous taxon name
Rename taxon	Genus	<i>Orthocystovirus</i>	<i>Cystovirus</i>
Rename taxon	Species	<i>Orthocystovirus phi6</i>	<i>Cystovirus phi6</i>
Rename taxon	Species	<i>Orthocystovirus phiNN</i>	<i>Cystovirus phiNN</i>

**Table 73.** Cystoviridae, 1 split taxa, 4 new taxa\*

Operation	Rank	Old taxon	New taxon 1	New taxon 2	New taxon 3	New taxon 4
Split taxon	Genus	<i>Cystovirus</i>	<i>Alphacystovirus</i>	<i>Betacystovirus</i>	<i>Gammacystovirus</i>	<i>Deltacystovirus</i>

\*Source/full text: [https://ictv.global/ictv/proposals/2024.043B.Cystoviridae\\_6ng\\_2ns\\_1rng\\_7rnsp.zip](https://ictv.global/ictv/proposals/2024.043B.Cystoviridae_6ng_2ns_1rng_7rnsp.zip).

## 2024.044B.Felixviridae\_1nf\_1nsf\_2ng\_1mg\_2ns

**Title:** Create one new family (*Felixviridae*), including one new subfamily (*Maevirinae*), three genera (two new: *Nakavirus*, *Chronivirus*; one existent: *Certevirus*), including two new species (*Nakavirus sapi* and *Chronivirus chronis*).

**Authors:** Nobrega F.L. (f.nobrega@soton.ac.uk), Rothschild-Rodriguez, D., Lambon, K.

### Summary

#### Taxonomic rank(s) affected

*Duplodnaviria*; *Heunggongviriae*; *Uroviricota*; *Caudoviricetes*.

#### Description of current taxonomy

Unclassified *Caudoviricetes*.

#### Proposed taxonomic change(s)

We propose a new family, *Felixviridae*, which includes the new subfamily, *Maevirinae*, and three genera: *Nakavirus* (new), *Chronivirus* (new) and *Certevirus* (already existent). The *Nakavirus* and *Chronivirus* genera include at least one newly proposed species each, *Nakavirus sapi* (for Klebsiella phage Roth C and Klebsiella phage Roth D), and *Chronivirus chronis* (for Klebsiella phage vB\_Kpn\_Chronis), respectively.

#### Justification

Phages assigned to this new family have not been taxonomically characterised, with only one previously cultured *Klebsiella* phage relative, vB\_Kpn\_Chronis, and a *Protoea* phage, PdC23. We isolated 53 phages as part of the Klebsiella Phage Collection, two of which, RothC and RothD, could not be assigned to any existing viral families, leading us to propose a new family, *Felixviridae*. Additionally, we found representatives of this family to be present in metagenomes originating from human stool samples, a proxy for the human gut microbiota, and in metagenomic sequences within the Gut Phage Database. Genomic analyses suggest that the family *Felixviridae* includes at least one subfamily, *Maevirinae*, and three genera, *Certevirus* and two new genera *Nakavirus* and *Chronivirus*. The *Nakavirus* genus includes species for Klebsiella phage Roth C and Klebsiella phage Roth D. High similarity between these viruses places them both in the proposed species *Nakavirus sapi*. The *Chronivirus* genus includes the species *Chronivirus chronis* for phage vB\_Kpn\_Chronis.

**Submitted:** 19 June 24; **Revised:** 27 June 24

**Table 74.** *Felixviridae*, 6 new taxa\*

Operation	Rank	New taxon name	Virus name	Exemplar
New taxon	Family	<i>Felixviridae</i>		
New taxon	Subfamily	<i>Maevirinae</i>		
New taxon	Genus	<i>Chronivirus</i>		
New taxon	Species	<i>Chronivirus chronis</i>	vB_Kpn_Chronis	MN013086
New taxon	Genus	<i>Nakavirus</i>		
New taxon	Species	<i>Nakavirus sapi</i>	Klebsiella phage RothC	PP934563

**Table 75.** *Felixviridae*, 1 move taxon\*

Operation	Rank	Taxon name	New parent taxon	Old parent taxon
Move taxon	Genus	<i>Certevirus</i>	<i>Felixviridae</i>	<i>Caudoviricetes</i>

\*Source/full text: [https://ictv.global/ictv/proposals/2024.044B.Felixviridae\\_1nf\\_1nsf\\_2ng\\_1mg\\_2ns.zip](https://ictv.global/ictv/proposals/2024.044B.Felixviridae_1nf_1nsf_2ng_1mg_2ns.zip).

## 2024.045B. Autographivirales

**Title:** Promoting the family *Autographiviridae* to create one new order, *Autographivirales*, with four new families, four new subfamilies, 93 new genera and 607 new species (*Duplodnaviria*, *Caudoviricetes*).

**Authors:** Turner D (dann2.Turner@uwe.ac.uk), Carrillo D, Lood C, Ely B, Lehman SM, Dutilh B, Kropinski AM, Lavigne R, Adriaenssens EM, Millard AD

### Summary

#### Taxonomic rank(s) affected

Realm: *Duplodnaviria*; kingdom: *Heunggongvirae*; phylum: *Uroviricota*; class: *Caudoviricetes*.

#### Description of current taxonomy

The family *Autographiviridae* was established under taxonomic proposal 2019.103B.

#### Proposed taxonomic change(s)

We propose

The establishment of a new order, *Autographivirales*, containing four new families.

The creation of four new subfamilies.

The creation of 93 new genera.

The creation of 610 new species.

Abolition of 21 species.

#### Justification

The proposed order forms a single deep-branching clade in tBLASTx distance analysis, reflected in core gene maximum-likelihood phylogeny. The proposed families form monophyletic clusters in proteome-based analyses and each share a number of core orthologous genes.

**Submitted:** 21 June 24; **Revised:** 30 September 24

**Table 76.** *Autographivirales*, 708 new taxa\*. Table too large, see supplementary information sheet supp\_info\_tab\_76

**Table 77.** *Autographivirales*, 60 move taxa\*. Table too large, see supplementary information sheet supp\_info\_tab\_77

**Table 78.** *Autographivirales*, 21 abolish taxa\*

Operation	Rank	Abolished taxon name
Abolish taxon	Species	<i>Friunavirus SWHAb1</i>
Abolish taxon	Species	<i>Friunavirus SWHAb3</i>
Abolish taxon	Species	<i>Phikmvirus NFS</i>
Abolish taxon	Species	<i>Phikmvirus PT2</i>
Abolish taxon	Species	<i>Vectrevirus cee</i>
Abolish taxon	Species	<i>Maculvirus OWB</i>
Abolish taxon	Species	<i>Murciavirus CB5A</i>
Abolish taxon	Species	<i>Kaohsiungvirus AS51</i>
Abolish taxon	Species	<i>Cuernavacavivirus RHEph09</i>
Abolish taxon	Species	<i>Atuphduovirus atuph03</i>
Abolish taxon	Species	<i>Przondovirus KpV766</i>
Abolish taxon	Species	<i>Teseptimavirus YpsPG</i>
Abolish taxon	Species	<i>Teseptimavirus YpPY</i>
Abolish taxon	Species	<i>Helsettivirus fPS53</i>
Abolish taxon	Species	<i>Teetrevirus T7M</i>
Abolish taxon	Species	<i>Berlinvirus Yepe2</i>

Continued

**Table 78.** Continued

Operation	Rank	Abolished taxon name
Abolish taxon	Species	<i>Berlinvirus Yepf</i>
Abolish taxon	Species	<i>Kayfunavirus EcpYZU01</i>
Abolish taxon	Species	<i>Pektosvirus PP81</i>
Abolish taxon	Species	<i>Ghunavirus Psa17</i>
Abolish taxon	Species	<i>Friunavirus AB3</i>

**Table 79.** Autographivirales, 14 move; rename taxa\*

Operation	Rank	New taxon name	New parent taxon	Old taxon name
Move; rename taxon	Species	<i>Rodentiumvirus LL11</i>	<i>Rodentiumvirus</i>	<i>Vectrevirus LL11</i>
Move; rename taxon	Species	<i>Rodentiumvirus CrRp3</i>	<i>Rodentiumvirus</i>	<i>Vectrevirus CrRp3</i>
Move; rename taxon	Species	<i>Nerthusvirus achelous</i>	<i>Nerthusvirus</i>	<i>Uliginvirus achelous</i>
Move; rename taxon	Species	<i>Nerthusvirus nerthus</i>	<i>Nerthusvirus</i>	<i>Uliginvirus nerthus</i>
Move; rename taxon	Species	<i>Nerthusvirus alpheus</i>	<i>Nerthusvirus</i>	<i>Uliginvirus alpheus</i>
Move; rename taxon	Species	<i>Njordvirus njord</i>	<i>Njordvirus</i>	<i>Uliginvirus njord</i>
Move; rename taxon	Species	<i>Ebriosvirus ebrios</i>	<i>Ebriosvirus</i>	<i>Teseptimavirus ebrios</i>
Move; rename taxon	Species	<i>Ebriosvirus IME15</i>	<i>Ebriosvirus</i>	<i>Teseptimavirus IME15</i>
Move; rename taxon	Species	<i>Hennigervirus shl2</i>	<i>Hennigervirus</i>	<i>Ghunavirus shl2</i>
Move; rename taxon	Species	<i>Hennigervirus PPPL1</i>	<i>Hennigervirus</i>	<i>Ghunavirus PPPL1</i>
Move; rename taxon	Species	<i>Hennigervirus henninger</i>	<i>Hennigervirus</i>	<i>Ghunavirus henninger</i>
Move; rename taxon	Species	<i>Unosvirus UNOSLW1</i>	<i>Unosvirus</i>	<i>Pifdecavivirus UNOSLW1</i>
Move; rename taxon	Species	<i>Pfluivirus PFP1</i>	<i>Pfluivirus</i>	<i>Pifdecavivirus PFP1</i>
Move; rename taxon	Species	<i>Pfluivirus pv22PfluR64PP</i>	<i>Pfluivirus</i>	<i>Pifdecavivirus pv22PfluR64PP</i>

**Table 80.** Autographivirales, 2 rename taxa\*

Operation	Rank	New taxon name	Previous taxon name
Rename taxon	Species	<i>Ghunavirus gv17A</i>	<i>Ghunavirus 17A</i>
Rename taxon	Species	<i>Warsawvirus wv3MF5</i>	<i>Warsawvirus 3MF5</i>

**Table 81.** Autographivirales, 1 promote taxon\*

Operation	New taxon name	Old rank	New rank
Promote taxon	<i>Autographivirales</i>	Family	Order

\*Source/full text: <https://ictv.global/ictv/proposals/2024.045B.Autographivirales.zip>.

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**Keywords:** Abidjanvirus Ab18; Abidjanvirus Ab19; Abidjanvirus PaMx11; Acadenvirus bigMiraUFV01; Acadenvirus premi; Acinetobacter virus Acj61; Actaeavirus; Actaeavirus CRP125; Actinidiaviruses; Actinidiaviruses Ep4; Adrianbuildvirus; Adrianbuildvirus ARI0923; Adrianbuildvirus IPP41; Adrianbuildvirus IPP42; Adrianbuildvirus IPP43; Adrianbuildvirus IPP44; Adrianbuildvirus IPP5; Adrianbuildvirus IPP51; Adrianbuildvirus SpSL1; Aequorvirus; Aequorvirus HTVC041P; Aeromonas virus 43; Ahdouovirus AH2; Aphunavirus A014L; Aphunavirus AHPMCC7; Aphunavirus LAH1; Aphunavirus P2; Aphunavirus ST21; Aphunavirus yong1; Alfirivirus; Alfirivirus alfirin; Alisviridae; Alphacystovirus; Alphacystovirus ph18; Ameliavirus viph1008o; Altadenavirus; Altadenavirus altadena; Altadenavirus bumble; Amboselivirus; Amboselivirus simi; Ameliavirus; Ameliavirus viph1008o; Anathivirus; Anathivirus anath; Andersonviridae; Andromedavirus leo2; Angmobivirus; Angmobivirus SCBP1; Antileavirus; Antileavirus jorvik; Apdecimavirus K12P11; Aristophanesvirus; Aristophanesvirus aristophanes; Armandavirus; Armandavirus PT2; Armandavirus PZLAh152; Armandavirus PZLAh8; Armandavirus T7Ah; Arnovirus; Arnovirus Wc4; Arnovirus arno162; Arnovirus arno18; Arthrobacter virus Liebe; Artimaviricota; Asemoviridae; Atoyacvirus; Atoyacvirus atoyac1; Atoyacvirus atoyac15; Atsuirnavirus; Atsuirnavirus caloris; Atuphduovirus atuph03; Autographivirales; Autonotataviridae; Autoscriptoviridae; Autosignataviridae; Autotranscriptaviridae; Axyvirus; Axyvirus 1932Axy09; Axyvirus 1932Axy21; Axyvirus 1932Axy23; Azeevirinae; Baileybluvirus callinalbarbz; Bajunvirus; Bajunvirus bajun; Bamvirus; Bamvirus bam; Barnstormervirus; Barnstormervirus barnstormer; Barnstormervirus caron; Baxterfoxvirus baxterfox; Baxterfoxvirus yeezy; Baxtervirus baxterfox; Baxtervirus yeezy; Benllochvirus; Benllochvirus K10PH82C1; Benllochvirus VLCpiA3a; Benllochvirus cmc355D; Benllochvirus cp31; Berlinvirus D226; Berlinvirus JSS1; Berlinvirus JSS2; Berlinvirus P151; Berlinvirus PC127; Berlinvirus PZJ0206; Berlinvirus PcCB251; Berlinvirus SEqdws315; Berlinvirus SPLA4; Berlinvirus SWJM03; Berlinvirus SalMLPST153; Berlinvirus V1; Berlinvirus Yepe2; Berlinvirus Yefp; Berlinvirus carlspitteler; Berlinvirus ernstbeyeler; Berlinvirus pEA5NUABM57; Berlinvirus p0103; Berryhillviridae; Bertilvirus; Bertilvirus bertil; Betacystovirus; Betacystovirus ph12; Bifilivirus philemonii; Bifseptvirus BIMBV45; Bifseptvirus SoKa; Boesrvirus; Boesrvirus BOESR1; Bolekvirus; Bolekvirus bolek; Bolekvirus lolek; Bonaevitae bonaevitae; Bonaevitae bonaevitae; Bonnellvirus Kc261; Bonnellvirus RZ4; Bonnellvirus altidsur; Bonnellvirus glasur; Bonnellvirus mellemur; Bonnellvirus smaasur; Bonnellvirus usur; Bordetella virus CN1; Bordetella virus CN2; Bordetella virus FP1; Bordetella virus MW2; Bronvirus bron; Bronvirus joedirt; Burkholderia virus AH2; Burkholderia virus BcepNazgul; Cafassovirus; Cafassovirus aleemili; Cafassovirus cafasso; Cafassovirus morganii; Cafassovirus obladi; Caliparnavirus; Caliparnavirus acidus; Campylobacter virus IBB35; Camvirus vansegeleni; Camvirus verabelle; Cankvirus; Cankvirus cv10P302A; Caparsinavirus FoX6; Casidaviridae; Catalonavirus NF1; Caulobacter virus Sansa; Cebaduodecimvirus; Cebaduodecimvirus ph12auna; Cebaduodecimvirus ph12duo; Ceetrepovirus C3PO; Ceetrepovirus darwin; Ceetrepovirus zion; Centumtrigintavirus; Centumtrigintavirus cv133; Cenunavirus; Cenunavirus Cen1621; Cepavirus; Cepavirus PAS7; Ceskevirus; Ceskevirus SB4; Chamipavirus; Chamipavirus RHEph21; Chemalvirus; Chemalvirus PseuGes254; Chennavirus; Chennavirus MVCVPHSA1; Cheoctovirus PMC; Cheoctovirus SG4; Cheoctovirus ardmore; Cheoctovirus boomer; Cheoctovirus che8; Cheoctovirus deadp; Cheoctovirus dlane; Cheoctovirus dorothy; Cheoctovirus dotproduct; Cheoctovirus drago; Cheoctovirus fruitloop; Cheoctovirus gumbie; Cheoctovirus ihubesi; Cheoctovirus llj; Cheoctovirus mozy; Cheoctovirus mutaforma13; Cheoctovirus pacc40; Cheoctovirus ramsey; Cheoctovirus rockyhorror; Cheoctovirus shauna1; Cheoctovirus shilan; Cheoctovirus spartacus; Cheoctovirus taj; Cheoctovirus tweety; Cheoctovirus wee; Chimalliviridae; Chivirus BP12C; Chronisvirus; Chronisvirus chronis; Chunghsingvirus P1201; Colingsworthviridae; Conareevirus; Conareevirus Cd1; Conareevirus babayka; Conareevirus doublea; Connertonviridae; Cornievirus cornie; Corticovirus; Corticovirus Cr39582; Corticovirus PM2; Corycianvirus; Corycianvirus MFV; Corycianvirus; Corycianvirus A014S; Corynebacterium virus C3PO; Corynebacterium virus Darwin; Corynebacterium virus P1201; Corynebacterium virus Zion; Cotavirus; Cotavirus cota; Cronosvirus EspYZU05; Cronosvirus EspYZU13; Cronosvirus GY3; Cronosvirus Kc318; Cuernavacavirus RHEph09; Cuernavacavirus RHph138; Cuernavacavirus RHphN37; Cuernavacavirus RHphTM33; Cullenvirus; Cullenvirus 6937; Cullenvirus K59PH2; Cullenvirus KYP; Cystovirus; Cystovirus phi6; Cystovirus phiNN; Dabrowskivirus; Dabrowskivirus KKP3916; Daeravirus; Daeravirus MA13; Daniellevirus; Daniellevirus Zyzzx; Daniellevirus danielle; Daolivirus; Daolivirus MJG; Dazunavirus; Dazunavirus DZ1; Dcmvirus; Dcmvirus DCM; Deltacytovirus; Deltacytovirus phi2954; Dewhirstvirus; Dewhirstvirus pgmg00J; Dewhirstvirus pgmg00K; Dewhirstvirus pgmg00L; Dewhirstvirus pgmg00M; Dexdertvirus kwekel; Dishuivirus; Dishuivirus DSLLC07; Divaquavirales; Dolichocephalovirinae; Dovevirinae; Drulisvirus BHU1; Drulisvirus BHU2; Drulisvirus BHU3; Drulisvirus BUCT631; Drulisvirus BUCT86; Drulisvirus Bp5; Drulisvirus CX1; Drulisvirus Dlv622; Drulisvirus FBKp18; Drulisvirus FK1979; Drulisvirus IME308; Drulisvirus IME337; Drulisvirus JK2P; Drulisvirus K15PH90; Drulisvirus K1PH164C1; Drulisvirus K24PH164C1; Drulisvirus K25PH129C1; Drulisvirus K39PH122C2; Drulisvirus K40PH129C1; Drulisvirus K51PH129C1; Drulisvirus K66PH128C1; Drulisvirus K71PH129C1; Drulisvirus K72PH164C2; Drulisvirus KA; Drulisvirus KMI3; Drulisvirus KMI6; Drulisvirus KPPK1081; Drulisvirus KPPK1082; Drulisvirus KPR2; Drulisvirus KXP; Drulisvirus KpV2883; Drulisvirus Kpn13; Drulisvirus LLY; Drulisvirus M21221; Drulisvirus NER40; Drulisvirus P1010; Drulisvirus P929; Drulisvirus PWKp1; Drulisvirus Pone; Drulisvirus QL; Drulisvirus SCNJ1Z; Drulisvirus SKP1; Drulisvirus SRD2021; Drulisvirus VAC25; Drulisvirus VLC1; Drulisvirus VLC3; Drulisvirus VLC4; Drulisvirus VLC5; Drulisvirus VLC6; Drulisvirus VLCpiA1b; Drulisvirus VLCpiA1c; Drulisvirus VLCpiA1d; Drulisvirus VLCpiA1e; Drulisvirus VLCpiA1f; Drulisvirus VLCpiA1g; Drulisvirus VLCpiA1h; Drulisvirus VLCpiA1i; Drulisvirus VLCpiA1j; Drulisvirus VLCpiA1k; Drulisvirus VLCpiA1l; Drulisvirus VLCpiA1m; Drulisvirus VLCpiA1n; Drulisvirus VLCpiA1o;

Drulisvirus VLCpiA1q; Drulisvirus VLCpiA1r; Drulisvirus ZH5; Drulisvirus ZX11; Drulisvirus ZX6; Drulisvirus tk2018; Drulisvirus cp48; Drulisvirus dv6993; Drulisvirus dv6995; Drulisvirus fHeKpn01; Drulisvirus pKPM18622; Drulisvirus pKp11; Drulisvirus pokalde001; Drulisvirus tk2018; Drulisvirus xx20; Dunnvirinae; Dynamenevirus; Dynamenevirus CRP114; Dynamenevirus CRP227; Dynamenevirus CRP361; Eastwestvirus; Eastwestvirus eastwest; Ebriosvirus; Ebriosvirus IME15; Ebriosvirus ebrios; Edwardsroadvirus; Edwardsroadvirus RRH1; Efekovirus efeko; Efekovirus efeko; Ehrlichviridae; Elsinorevirus; Elsinorevirus N016; Elunavirus PagPSK1; Elunavirus stepyanka; Emotionvirus; Emotionvirus emotion; Eseptimavir KKP3831; Epsiloncystovirus; Epsiloncystovirus phiNY; Epsomviridae; Eracentumvirus Nifs112; Escherichia phage ESC013; Escherichia virus CF2; Escherichia virus DE3; Escherichia virus ESC05; Escherichia virus Schickermooser; Escherichia virus phAPEC8; Eucampyvirinae; Euvesivirus; SB3; Excelsiorvirus; Excelsiorvirus pgng00S; Felixounavirus AS01A; Felixounavirus BPSEL1; Felixounavirus CL1; Felixounavirus CRP22; Felixounavirus CapYZU01; Felixounavirus D12; Felixounavirus DE17; Felixounavirus DE7; Felixounavirus DR094; Felixounavirus EC106; Felixounavirus ECOH1; Felixounavirus EF202P1; Felixounavirus ESC045; Felixounavirus ESC049; Felixounavirus ESC050; Felixounavirus GSP193; Felixounavirus IME338; Felixounavirus JK55; Felixounavirus JLBYU28; Felixounavirus JLBYU32; Felixounavirus JN01; Felixounavirus KhF1; Felixounavirus L27; Felixounavirus LMP25; Felixounavirus MBP496116; Felixounavirus NBEC004; Felixounavirus NBEC005; Felixounavirus NBSal004; Felixounavirus NJ12; Felixounavirus OPTSAL01; Felixounavirus PHB11; Felixounavirus Pr103Blw; Felixounavirus REP5; Felixounavirus REP8; Felixounavirus RP3; Felixounavirus Ro111lw; Felixounavirus S19cd; Felixounavirus SEP1; Felixounavirus SME50; Felixounavirus SPJ41; Felixounavirus ST11; Felixounavirus SUTS720; Felixounavirus SWJM02; Felixounavirus Sp3Shan2021; 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Grandeviridae; Grimontviridae; Guangxiviruses; Guangxivirus PSTM2; Gujervirinae; Gundecimvirus; Gundecimvirus MG11; Gyeongsanvirus PPSG11; Haasevirus; Haasevirus pgng00R; Haasevirus pgng00T; Haasevirus pgng00U; Haasevirus pgng00V; Haasevirus pgng00W; Hakuzoviridae; Hapakavirus; Hapakavirus Nufs112; Hatfieldvirus; Hatfieldvirus porci; Helsettvirus YpEc11; Helsettvirus fPS53; Hennigervirus; Hennigervirus MR1; Hennigervirus MR2; Hennigervirus PPPL1; Hennigervirus henninger; Hennigervirus shl2; Higashivirus BHDTSo9; Hilgardivirus; Hilgardivirus vroomvroom; Hinxtionvirus; Hinxtionvirus ARI0004; Hinxtionvirus ARI0031; Hinxtionvirus ARI02851; Hinxtionvirus ARI0462; Hinxtionvirus ARI04681; Hinxtionvirus ARI0468b3; Hinxtionvirus ARI0831b; Hinxtionvirus DCC1738; Hinxtionvirus IC1; Hinxtionvirus IPP34; Hinxtionvirus IPP46; Hinxtionvirus IPP64; Hinxtionvirus IPP69; Hinxtionvirus K13; Hinxtionvirus V22; Hinxtionvirus hv2167; Hinxtionvirus hv34117; Hinxtionvirus hv8140; Hodgkinviridae; Hongshanvirus; 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Lindbergviridae; Linggongvirus; Linggongvirus VH5; Lovomskayavirus shawty; Loughboroughvirus ZCSE2; Lucadorvirus gail; Lucadorvirus jeeves; Lucadorvirus luchador; Luchadorvirus gail; Luchadorvirus jeeves; Luchadorvirus luchador; Ludisvirus; Ludisvirus pgng00A; Lullwatervirus quill52; Lundtoftevirus; Lundtoftevirus Lu221; Luzcentumvirus LUZ100; Mabodamacavirus; Mabodamacavirus mabodamac; Maculvirus AC2; Maculvirus BUCT233; Maculvirus DE10; Maculvirus DE17; Maculvirus DE18; Maculvirus F23s2; Maculvirus FE11; Maculvirus GHSM17; Maculvirus H256D1; Maculvirus HA1; Maculvirus HA5; Maculvirus MG01; Maculvirus OWB; Maculvirus SrVc9; Maculvirus VP9; Maevirinae; Maklayavirus; Maklayavirus Q19; Malkevirus; Malkevirus ARI02853; Malkevirus IPP45; Malkevirus IPP67; Malkevirus mv11865; Malkevirus mv23782; Mallvirus; Mallvirus ParMal1; Maltophivirus; Maltophivirus BUCT609; Manhattanvirus vresidence; Manhattanvirus wildwest; Marchewkaviruses; Marchewkaviruses domovoii; Marchewkaviruses kabachok; Marchewkaviruses marchewka; 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Mycobacterium virus Mozy; Mycobacterium virus Mutaforma13; Mycobacterium virus PMC; Mycobacterium virus Pacc40; Mycobacterium virus Ramsey; Mycobacterium virus Renaud18; Mycobacterium virus RockyHorror; Mycobacterium virus SG4; Mycobacterium virus Shauna1; Mycobacterium virus Shilan; Mycobacterium virus Spartacus; Mycobacterium virus Squirty; Mycobacterium virus TChen; Mycobacterium virus TM4; Mycobacterium virus Taj; Mycobacterium virus ThetaBob; Mycobacterium virus Tortellini; Mycobacterium virus Tweety; Mycobacterium virus Wee; Mycobacterium virus Wildcat; Myosmarivirus SMP; Myranavirus phabba; Myrnavirus phabba; Nairobiivirus; Nairobiivirus nv36; Nakavirus; Nakavirus sapi; Natansvirus; Natansvirus SnaPR1; Nazgulvirus bcepnazgul; Nerivirus; Nerivirus SSRP01; Nerthusvirus; Nerthusvirus BUCT553; Nerthusvirus achelous; Nerthusvirus alpheus; Nerthusvirus nerthus; Ningirsuvirus DchS19; Ningirsuvirus W2B; Ningirsuvirus nahilimali; Ningirsuvirus pEpSNUABM12; Ningirsuvirus sibilus; Nixviridae; 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Sicariusvirus wyborn; Smasvirus; Smasvirus BUCT598; Smasvirus BUCT700; Smasvirus P15; Smasvirus SB1; Smasvirus c9N; Snaubvirus; Snaubvirus pEpSNUABM09; Solymavirus; Solymavirus PSTCR2; Solymavirus PSTCR120; Songlingvirus; Songlingvirus SI01; Soropartiviridae; Spinunavirus; Spinunavirus AR101311; Spinunavirus AR10399; Spinunavirus AR104601; Spinunavirus AR104602; Spinunavirus AR104682; Spinunavirus IPP10; Spinunavirus IPP31; Spinunavirus IPP32; Spinunavirus IPP36; Spinunavirus IPP37; Spinunavirus IPP50; Spinunavirus IPP60; Spinunavirus IPP61; Spinunavirus IPP8; Spinunavirus IPP9; Spinunavirus SF39; Spinunavirus Spn1; Spinunavirus sv040922; Spiovirus; Spiovirus sbp1; Splendidredvirus; Splendidredvirus ry17; Splendidredvirus splendidred; Squiryvirus squirty; Stackebrandtviridae; Stentvirinae; Streptomyces virus Karimac; Streptomyces virus LukeCage; Streptomyces virus StarPlatinum; Streptomyces virus Wolford; Streptomyces virus Yaboi; Sumtervirus; Sumtervirus S2B; Suseptimavirus PAS59; Suseptimavirus sv4E8; Suspivirus; Suttonboningtonvirus; Suttonboningtonvirus sv11Co2020; Swepdovirus; Swepdovirus SWEP2; Sycamorevirus; Sycamorevirus sycamore; Tartuvirus; Tartuvirus amme3; Tartuvirus kopa4; Tartuvirus nopa; Tartuvirus roomu2; Teetrevirus CT02; Teetrevirus EF2; Teetrevirus Emp27; Teetrevirus F1M1C; Teetrevirus F2M1C; Teetrevirus F5M1C; Teetrevirus HMGUsm2; Teetrevirus IME305; Teetrevirus K19PH14C4P1; Teetrevirus KP13MC52; Teetrevirus Kp31; Teetrevirus Kpn11mx; Teetrevirus KpnPVAC1; Teetrevirus NLZS3; Teetrevirus Seera; Teetrevirus SALSA; Teetrevirus T7M; Teetrevirus bumbleweed; Teetrevirus glastosback; Teetrevirus keithsmous; Teetrevirus keithstache; Teetrevirus megaducksbill; Teetrevirus mtp14; Teetrevirus mtp15; Teetrevirus mtp4; Teetrevirus mtp8; Teetrevirus patroon; Teetrevirus sekstaphage1; Teetrevirus tv066044; Teetrevirus tv31; Teetrevirus tv6996; Teetrevirus tv6998; Teetrevirus vB8388; Tepoztlanvirus; Tepoztlanvirus RHphTM34; Tepoztlanvirus RHphY120; Tequatorvirus; Tequatorvirus CF2; Teseptimavirus A1122; Teseptimavirus C2; Teseptimavirus EFA2; Teseptimavirus JB01; Teseptimavirus PHB19; Teseptimavirus PVN09; Teseptimavirus SYGE1; Teseptimavirus UFV01; Teseptimavirus YpPY; Teseptimavirus YpYe09; Teseptimavirus YpsPG;

Teseptimavirus jacobburckhardt; Teseptimavirus jeantinguely; Teseptimavirus p0111; Teseptimavirus pila; Teseptimavirus tv04922B; Teseptimavirus yanou; Tevenvirinae; Thetabovirus renaud18; Thetabovirus tchen; Thetabovirus thetabob; Thoosavirus; Thoosavirus HTVC025P; Timquatrovirus TM4; Tiranvirus; Tiranvirus STIP28; Tolavirus; Tolavirus tola; Torinorumvirus; Torinovirus; Torinorumvirus B11; Torinovirus K7A1; Tortellinivirus tortellini; Trautnerviridae; Treelivelvirus; Treelivelvirus VAC51; Tritejavirus; Tritejavirus CRP804; Troedvirus stalingrad; Trogglehumpervirus; Trogglehumpervirus trogglehumper; Tuaanevirus ime13; Tulanevirus ime13; Tunggulvirus GM223; Uavernvirus; Uavernvirus uavern; Uetakevirus PAS61; Ulipvirus; Ulipvirus IME184; Ulipvirus K1ULIP33; Ulipvirus K44PH129C1; Ulipvirus cp4; Ulipvirus pKPM18621; Unosvirus; Unosvirus PCS4; Unosvirus UNOSLW1; Unyawovirus PC2B6; Vandenedenieviridae; Vashvirus euratis; Vectrevirus EC120; Vectrevirus EEC2; Vectrevirus KAW1A4500; Vectrevirus M1B1P10; Vectrevirus P101101UKE1; Vectrevirus PUT189UKE2; Vectrevirus ULINTec4; Vectrevirus ULINTec6; Vectrevirus ULINTec7; Vectrevirus cee; Vectrevirus vv6948; Ventosavirus; Ventosavirus ventosus; Vetruaniviruses; Vetruaniviruses porcinsecundi; Vetruaniviruses dhakaense; Vetruaniviruses porcinprimi; Vetruaniviruses primi; Vetruaniviruses secundi; Vinavirales; Vistulavirus; Vistulavirus BB8; Vividuovirus lennon; Vividuovirus sitar; Vojvodinavirus CN1; Waldovirus; Vojvodinavirus CN2; Vojvodinavirus FP1; Vojvodinavirus MW2; Waldovirus plaquesplease; Waldovirus waldo5; Warsawvirus 3MF5; Warsawvirus wv3MF5; Weillhallvirus; Weillhallvirus wv16Q; Wendovervirus; Wendovervirus sonii; Wifcevirus AV128; Wifcevirus EC150; Wifcevirus EC071P1; Wifcevirus Ro157lw; Wifcevirus SP13; Wifcevirus mansfield; Wildcatvirus wildcat; Wizardvirus halo3; Wodongavirus; Wodongavirus REQ3; Wuhanvirus PS07; Xeniaduovirus; Xeniaduovirus xenia2; Xylella virus Sano; Yangvirus ascela; Yangvirus berries; Yangvirus cassia; Yangvirus janeemi; Yangvirus nitro; Yangvirus tfortroy; Yangvirus tuck; Yautepécovirus; Yautepécovirus RHPhX39; Yinyavirus; Yinyavirus AerP220; Yuanmingyuanvirus; Yuanmingyuanvirus NJ2; Yuavirus LK04; Yuavirus M6; Yuavirus MP1412; Yuavirus PAE1; Yuavirus yua; Yunamivirus; Yunamivirus Y1M1; Yvonnevirus yvonneastaic; Zetacystovirus; Zetacystovirus CAP; Zindervirus GRNsp51; Zindervirus kenzaK30; Zitchvirus apunk; Zitchvirus sampson; Zitchvirus tardus; Zitchvirus viaconlectus; Zoomievirus; Zoomievirus zoomie.

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#### Conflicts of interest

The authors declare no conflict of interest.

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