VIROLOGY DIVISION NEWS



Taxonomy of prokaryotic viruses: 2017 update from the ICTV Bacterial and Archaeal Viruses Subcommittee

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The prokaryotic virus community is represented at the International Committee on Taxonomy of Viruses (ICTV) by the Bacterial and Archaeal Viruses Subcommittee. Since our last report [5], the committee composition has changed, and a large number of taxonomic proposals (TaxoProps) were submitted to the ICTV Executive Committee (EC) for approval.

1. Changes in subcommittee membership. During the past year we have lost two members. Dr. Hans-Wolfgang Ackermann, a life member of the ICTV, the father of caudovirus taxonomy [1] and an electron microscopist extraordinaire [2–4], lamentably died and will be gravely missed. In addition, Dr. Jens H. Kuhn, who, in spite of protestations about not being a genuine phage biologist, proved invaluable

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Table 1 List of current subcommittee members who have new responsibilities (*), along with new members of the subcommittee

Name	Country	Position
Evelien Adriaenssens*	United Kingdom	Chair, Caudovirales phage study group
Dann Turner	United Kingdom	Chair, Acinetobacter phage study group
Jakub Barylski*	Poland	Chair, Bacillus phage study group
Jochen Klumpp*	Switzerland	Chair, Listeria phage study group
Małgorzata Łobocka	Poland	Chair, Staphylococcus phage study group
Poliane Alfenas-Zerbini	Brazil	Member
Ramy Aziz	Egypt	Member
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Yigang Tong	People's Republic of China	Member
Leonardo van Zyl	South Africa	Member
Jumpei Uchiyama	Japan	Member
Nicola K. Petty	Australia	Member

to our discussions and preparation of TaxoProps and manuscripts, resigned from the Subcommittee. Both Hans and Jens are acknowledged for their significant contributions to prokaryotic virus taxonomy. Furthermore, a number of current members have new responsibilities; and, in an effort to increase the geographical diversity of members, we appointed representatives from South America, Africa, and Asia (Table 1).

2. Changing the names of prokaryotic virus genera. A significant number of prokaryotic virus genera have either unpronounceable names (e.g., *Pocjvirus*, *Rdjlvirus*) or incorporated numerals (e.g., *T4virus*, *D3112virus*). In the first case, these names contravene The International Code of Virus Classification and Nomenclature (ICVCN, April 2017) Rule 3.12, which states "Names for taxa shall be easy to use and easy to remember. Euphonious names are preferred." In the latter case, pronunciation is a problem. For example,

is *D3112virus* pronounced "Dee+three thousand one hundred and twelve+virus" or "Dee+thirty one+twelve+virus" or "Dee+three+one+one+two+virus"? In addition, this nomenclature differs drastically from that for other virus taxa; and, would be incompatible with a Linnaean system of nomenclature [13]. We identified all prokaryotic taxon names that are problematic in the ICTV Master Species List (https://talk.ictvonline.org/files/master-species-lists/m/msl/6776) and suggested alternative names (Supplementary data file S1). These changes will be proposed officially at the next meeting of the ICTV EC in 2018.

3. Re-evaluation of the SPO1-like virus taxonomy. Over the past two years, members of the subcommittee have re-evaluated the taxonomy of a subset of myoviruses related to *Bacillus* phage SPO1. This group, made up of members of the subfamily *Spounavirinae* [10] and several genera of *Bacillus*-infecting viruses, was represented as a distinct

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Table 2 Taxonomy proposals (TaxoProps) proposing new taxa (families, subfamilies, genera, species) submitted to the ICTV Executive Committee in 2017

Family	Subfamily	Genus	Type species	No. of species in genus***
Ackermannviridae	Aglimvirinae	Ag3virus	Shigella virus AG3	1 (2)
Ackermannviridae	Aglimvirinae	Limestonevirus	Dickeya virus Limestone	1 (2)
Ackermannviridae	Cvivirinae	Cba120virus	Escherichia virus CBA120	4 (9)
Ackermannviridae	Cvivirinae	Vi1virus*	Salmonella virus ViI	(5)
Ackermannviridae	unassigned	unassigned	Erwinia virus Ea2809, Serratia virus MAM1, Serratia virus IME250, Klebsiella virus 0507KN21	4
Myoviridae*		Arvunavirus	Arthrobacter virus ArVI	2
Myoviridae*		Eah2virus	Erwinia virus EaH2	2
Myoviridae*		Machinavirus	Erwinia virus Machina	1
Myoviridae*		Ntreusvirus	Salmonella virus SPN3US	1
Myoviridae*		Svunavirus	Geobacillus virus GBSVI	2
Myoviridae*	Ampvirinae	Chippewavirus	Arthrobacter virus BarretLemon	1
Myoviridae*	Ampvirinae	Jawnskivirus	Arthrobacter virus Jawnski	2
Myoviridae*	Ampvirinae	Sonnyvirus	Arthrobacter virus Sonny	3
Podoviridae*	•	Dfl12virus	Dinoroseobacter virus DFL12phi1	1
Podoviridae*		Jwalphavirus	Achromobacter virus JWAlpha	2
Podoviridae*		P22virus*	Salmonella virus P22	1 (5)
Podoviridae*		Sp58virus	Salmonella virus SP058	3
Portogloboviridae		Alphaportoglobovirus	Sulfolobus alphaportoglobovirus 1	1
Siphoviridae*		Anatolevirus	Propionibacterium virus Anatole	2
Siphoviridae*		Attisvirus	Gordonia virus Attis	1
Siphoviridae*		Doucettevirus	Propionibacterium virus Doucette	4
Siphoviridae*		Hk97virus	Escherichia virus HK97*	9 (11)
Siphoviridae*		Lambdavirus*	Escherichia virus Lambda	3 (4)
Siphoviridae*		Pfr1virus	Propionibacterium virus PFR1	1
Siphoviridae*		Tp84virus	Geobacillus virus TP84	1
Siphoviridae*		Trigintaduovirus	Mycobacterium virus 32HC	1
Siphoviridae*		Wizardvirus	Gordonia virus Wizard	2
Siphoviridae*	Chebruvirinae	Brujitavirus	Mycobacterium virus Brujita	(2)
Siphoviridae*	Chebruvirinae	Che9cvirus*	Mycobacterium virus Che9c	1 (2)
Siphoviridae*	Dclasvirinae	Hawkeyevirus	Mycobacterium virus Hawkeye	1
Siphoviridae*	Dclasvirinae	Plotvirus	Mycobacterium virus PLot	1
Siphoviridae*	Mccleskeyvirinae	Lmd1virus	Leuconostoc virus Lmd1	6
Siphoviridae*	Mccleskeyvirinae	Una4virus	Leuconostoc virus 1A4	6
Siphoviridae*	Nclasvirinae	Buttersvirus	Mycobacterium virus Butters	2
Siphoviridae*	Nclasvirinae	Charlievirus	Mycobacterium virus Charlie	2 (3)
Siphoviridae*	Nclasvirinae	Redivirus	Mycobacterium virus Redi	3 (4)
Siphoviridae*	Nymbaxtervirinae	Baxtervirus	Gordonia virus BaxterFox	2
Siphoviridae*	Nymbaxtervirinae	Nymphadoravirus	Gordonia virus Nymphadora	3
Cystoviridae*	•	Cystovirus*	Pseudomonas virus phi6	6 (7)
Tectiviridae*		Alphatectivirus**	Pseudomonas virus PRD1	1 (2)
Tectiviridae*		Betatectivirus	Bacillus virus Bam35	2 (4)

^{*}taxon established, **previously known as *Tectivirus*, ***Number in parenthesis indicates the total number of viral species in this genus



module in various network analyses published recently [8, 9]. Using a combination of genomic, proteomics, and phylogenetic approaches, we have shown that this group of phages represents a new family, comprising five subfamilies and 13 genera [7]. We therefore suggest that these viruses be moved from their current taxonomic position in the family *Myoviridae* to a new family included in the order *Caudovirales*.

- **4. New taxa.** Table 2 lists of all new taxa proposed at the ICTV EC49 meeting in Singapore in 2017. In total, two new families, eight new subfamilies, 34 new genera, and 91 new species were proposed. Two significant items are on this list. The first item is the introduction of two new families of prokaryotic viruses: *Ackermannviridae* and *Portogloboviridae*. With the acceptance of changes to ICVCN Rule 3.11, the second item is the application of the names of eminent phage scientists, specifically Hans-Wolfgang Ackermann (Université Laval) and Charles Shelton McCleskey (Louisiana State University) as prefixes for taxon name stems.
- **5.** Updates to taxonomy. As the readership may be aware, "Virus Taxonomy: The Classification and Nomenclature of Viruses The Online (10th) Report of the ICTV" is freely accessible at http://ictv.global/report. We would like to acknowledge the hard work of Hanna M. Oksanen (*Corticoviridae*), Dennis H. Bamford (*Pleolipoviridae*), and Minna M. Poranen (*Cystoviridae*) for completing updates to their sections. The family *Pleolipoviridae* is now recognized as the first virus taxon in the newly established ICTV category for ssDNA/dsDNA Viruses. The summaries of the ICTV Report chapters are published in The Journal of General Virology [6, 11, 12].

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Compliance with ethical standards

The views and conclusions contained in this document are those of the authors and should not be interpreted as necessarily representing the official policies, either expressed or implied, of the US Department of Health and Human Services or of the institutions and companies affiliated with the authors.

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