

This form should be used for all taxonomic proposals. Please complete all those modules that are applicable (and then delete the unwanted sections). For guidance, see the notes written in blue and the separate document "Help with completing a taxonomic proposal"

Please try to keep related proposals within a single document; you can copy the modules to create more than one genus within a new family, for example.

MODULE 1: TITLE, AUTHORS, etc

| Code assigned: | 2016.027a-dB | | | (to be completed by ICTV officers) | | | |
|---|---------------------|------------|--------------------|------------------------------------|-----------------------|--|--|
| Short title: To create one (1) refamily Siphoviridae. (e.g. 6 new species in the genus 2 Modules attached (modules 1 and 10 are required) | rvinvirus, 1 🔀 6 🗌 | including | two (2) n 3 ⊠ 8 □ | ew species 4 | in the 5 □ 10 ⊠ | | |
| Author(s): | | | | | | | |
| Andrew M. Kropinski—University of Guelph (Canada) Jens H. Kuhn—NIH/NIAID/IRF-Frederick, Maryland (USA) Evelien M. Adriaenssens—University of Pretoria (South Africa) | | | | | | | |
| Corresponding author with e | e-mail address | : | | | | | |
| Andrew M. Kropinski Phage.C | Canada@gmail. | com | | | | | |
| List the ICTV study group(s) | that have see | n this pro | posal: | | | | |
| list of study groups and contacts is provided at http://www.ictvonline.org/subcommittees.asp . If a doubt, contact the appropriate subcommittee hair (fungal, invertebrate, plant, prokaryote or ertebrate viruses) ICTV Bacterial and Archaeal Virus Subcommittee | | | | | Viruses | | |
| ICTV Study Group comments (if any) and response of the proposer: | | | | | | | |
| | | | | | | | |
| Date first submitted to ICTV: Date of this revision (if different | | | | | | | |
| ICTV-EC comments and response of the proposer: | | | | | | | |
| | | | | | | | |

MODULE 2: NEW SPECIES

creating and naming one or more new species.

If more than one, they should be a group of related species belonging to the same genus. All new species must be placed in a higher taxon. This is usually a genus although it is also permissible for species to be "unassigned" within a subfamily or family. Wherever possible, provide sequence accession number(s) for **one** isolate of each new species proposed.

| Code 2016.027aB | | | (assigned by IC | (assigned by ICTV officers) | | | |
|--|-----------------|---|-----------------|--|--|--|--|
| To creat | te 2 no | ew species within: | | | | | |
| Genus: <i>Marvinvirus</i> (new) Subfamily: | | | v) | Fill in all that apply. • If the higher taxon has yet to be created (in a later module, below) write | | | |
| Fa | mily: Drder: | Siphoviridae Caudovirales | | "(new)" after its proposed name. If no genus is specified, enter "unassigned" in the genus box. | | | |
| Name of new species: | | Representative isol per species please) | late: (only 1 | GenBank sequence accession number(s) | | | |
| Mycobacterium virus Marvin Mycobacterium virus Mosmoris | | Mycobacterium pha Mycobacterium pha | _ | JF704100 KJ538721 | | | |

Reasons to justify the creation and assignment of the new species:

- Explain how the proposed species differ(s) from all existing species.
 - o If species demarcation criteria (see module 3) have previously been defined for the genus, **explain how the new species meet these criteria**.
 - o If criteria for demarcating species need to be defined (because there will now be more than one species in the genus), please state the proposed criteria.
- Further material in support of this proposal may be presented in the Appendix, Module 9

We have chosen 95% DNA sequence identity as the criterion for demarcation of species in this new genus. The members of each of the proposed species differ from those of other species by more than 5% at the DNA level as confirmed with the BLASTN algorithm.

MODULE 3: NEW GENUS

creating a new genus

Ideally, a genus should be placed within a higher taxon.

| Code | 201 | 6.027bB | (assigned by ICTV officers) | | | |
|-----------|-------|---------------|-----------------------------|---|--|--|
| To create | a new | genus within: | | Fill in all that apply. | | |
| Subfa | mily: | | | If the higher taxon has yet to be created | | |
| Fai | mily: | Siphoviridae | | (in a later module, below) write "(new)" after its proposed name. | | |
| 0 | rder: | Caudovirales | | If no family is specified, enter "unassigned" in the family box | | |

naming a new genus

| Code | 2016.027cB | (assigned by ICTV officers) |
|-----------|---------------------------|-----------------------------|
| To name t | he new genus: Marvinvirus | |

Assigning the type species and other species to a new genus

| Code | 2016.027dB | (assigned by ICTV officers) | | | |
|-----------|-----------------------------------|---|--|--|--|
| To desig | gnate the following as the type s | pecies of the new genus | | | |
| Mycobac | cterium virus Marvin | Every genus must have a type species. This should be a well characterized species although not necessarily the first to be discovered | | | |
| are being | • | w species created and assigned to it (Module 2) and any that Please enter here the TOTAL number of species aus will contain: | | | |

Reasons to justify the creation of a new genus:

Additional material in support of this proposal may be presented in the Appendix, Module 9

Mycobacterium phage Marvin was isolated by enrichment with *Mycobacterium smegmatis* mc²155 from soil from Radnor, PA, USA in 2009. The phage's genome has 11bp 3' sticky overhangs (sequence not given). This phage is the first fully sequenced member of Cluster S of Mycobacterium phages (http://phagesdb.org/phages/Marvin//). The next mostly closely related phages are Mycobacterium phages MosMoris and Catdawg (KF017002.1), which share 94 and 3% DNA sequence relatedness, respectively (Fig 2, 3).

Origin of the new genus name:

Based upon the name of the first sequenced member of this genus.

Reasons to justify the choice of type species:

The first sequenced member of this genus.

Species demarcation criteria in the new genus:

If there will be more than one species in the new genus, list the criteria being used for species demarcation and explain how the proposed members meet these criteria.

We have chosen 95% DNA sequence identity as the criterion for demarcation of species in this new genus. The members of each of the proposed species differ from those of other species by

more than 5% at the DNA level as confirmed with the BLASTN algorithm.

MODULE 10: **APPENDIX**: supporting material

additional material in support of this proposal

References:

- 1. Dereeper A, Guignon V, Blanc G, Audic S, Buffet S, Chevenet F, Dufayard JF, Guindon S, Lefort V, Lescot M, Claverie JM, Gascuel O. Phylogeny.fr: robust phylogenetic analysis for the non-specialist. Nucleic Acids Res. 2008; 36(Web Server issue):W465-9.
- 2. Darling AE, Mau B, Perna NT. progressiveMauve: multiple genome alignment with gene gain, loss and rearrangement. PLoS One. 2010; 5(6):e11147.

Annex:

Include as much information as necessary to support the proposal, including diagrams comparing the old and new taxonomic orders. The use of Figures and Tables is strongly recommended but direct pasting of content from publications will require permission from the copyright holder together with appropriate acknowledgement as this proposal will be placed on a public web site. For phylogenetic analysis, try to provide a tree where branch length is related to genetic distance.

Fig. 1. Electron micrograph of negatively stained Mycobacterium phage Marvin (http://phagesdb.org/phages/Marvin/) - Limited permission was granted by The Actinobacteriophages Database, funded by the Howard Hughes Medical Institute, to use this electron micrograph for this taxonomy proposal; it cannot be reused without permission of The Actinobacteriophages Database.

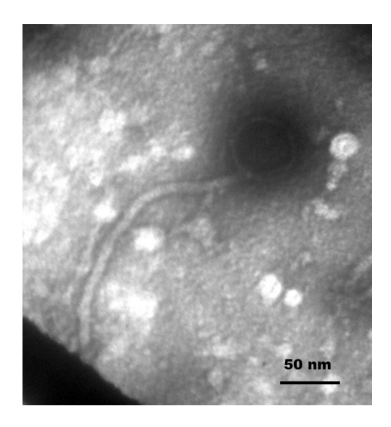


Table 1. Properties of phages belonging to the *Marvinvirus*.

| Mycobacterium phage | RefSeq | INSDC | Genome length (kb) | GC% | Protein | - | % homologous proteins** |
|---------------------|-------------|----------|--------------------------|------|---------|-----|-------------------------------|
| Marvin | None | JF704100 | 65.10 | 63.4 | 107 | 100 | 100 |
| MosMoris*** | NC_024138.1 | KJ538721 | 62.24 | 64.3 | 111 | 94 | 94.4 |

^{*} determined using BLASTN at NCBI; ** determined using CoreGenes; ***11-bp 3'cohesive overhang (GCGCGCAGCGC)

Fig. 2. Phylogenetic analysis of large subunit terminase proteins of a variety of Mycobacterium phages constructed using "one click" at phylogeny.fr [1]. "The "One Click mode" targets users that do not wish to deal with program and parameter selection. By default, the pipeline is already set up to run and connect programs recognized for their accuracy and speed (MUSCLE for multiple alignment and PhyML for phylogeny) to reconstruct a robust phylogenetic tree from a set of sequences." It also includes the use of Gblocks to eliminate poorly aligned positions and divergent regions. "The usual bootstrapping procedure is replaced by a new confidence index that is much faster to compute. See: Anisimova M., Gascuel O. Approximate likelihood ratio test for branches: A fast, accurate and powerful alternative (Syst Biol. 2006;55(4):539-52.) for details". The TerL from Gaia contained a intein which was removed before the phylogenetic tree was constructed. The members of the genus *Marvinvirus* are boxed in red.

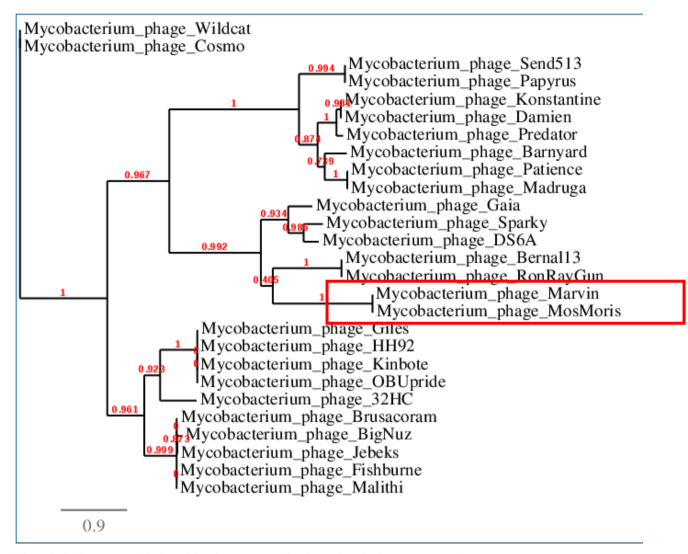


Figure 1: Phylogenetic tree (the branch length is proportional to the number of substitutions per site).

Fig. 3. progressiveMauve alignment (2) of the genomes of the two members of the genus *Marvinvirus*, Mycobacterium phages Marvin and MosMoris, and the closest relatives Mycobacterium phages CatDawg, RonRayGun, and Bernal13. Colored blocks indicate the regions of 1 to 1 best alignment with rearrangement breakpoints in a different random color. The degree of sequence similarity between regions is given by a similarity plot within the colored blocks with the height of the plot proportional to the average nucleotide identity (Aaron Darling, personal communication).

