



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:	2016.022a-dM	(to be completed by ICTV officers)
Short title: Three (3) new species in one new genus (<i>Goukovirus</i>) to be included in the proposed family <i>Phenuiviridae</i> in the proposed order <i>Bunyavirales</i> (e.g. 6 new species in the genus <i>Zetavirus</i>)		
Modules attached (modules 1 and 11 are required)	6 <input type="checkbox"/> 7 <input type="checkbox"/> 8 <input type="checkbox"/> 9 <input type="checkbox"/> 10 <input type="checkbox"/> 2 <input checked="" type="checkbox"/> 3 <input checked="" type="checkbox"/> 4 <input type="checkbox"/> 5 <input type="checkbox"/>	

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List the ICTV study group(s) that have seen this proposal:

A list of study groups and contacts is provided at <http://www.ictvonline.org/subcommittees.asp> . If in doubt, contact the appropriate subcommittee chair (fungal, invertebrate, plant, prokaryote or vertebrate viruses)

ICTV *Bunyaviridae* Study Group

ICTV Study Group comments (if any) and response of the proposer:

The ICTV *Bunyaviridae* Study Group has seen and discussed this proposal, and agreed to its submission to the ICTV Executive Committee based on votes of support by individual Study Group members or the absence of dissenting votes.

Date first submitted to ICTV:

July 18, 2016

Date of this revision (if different to above):

September 21, 2016

ICTV-EC comments and response of the proposer:

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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	2016.022aM	(assigned by ICTV officers)
To create 3 new species within:		
Genus:	<i>Goukovirus</i> (NEW)	Fill in all that apply. • If the higher taxon has yet to be created (in a later module, below) write “(new)” after its proposed name. • If no genus is specified, enter “unassigned” in the genus box.
Subfamily:	unassigned	
Family:	<i>Phenuiviridae</i> (NEW, see TP 2016.030M)	
Order:	<i>Bunyvirales</i> (NEW, see TP 2016.030M)	
Name of new species:	Representative isolate: (only 1 per species please)	GenBank sequence accession number(s)
<i>Gouleako goukovirus</i>	Gouléako virus (GOLV) A5/CI/2004	HQ541736–HQ541738
<i>Cumuto goukovirus</i>	Cumuto virus (CUMV) TR7094	KF543244–KF543246
<i>Yichang insect goukovirus</i>	Yíchāng insect virus (YIV) YCYC01	KM817763, KM817730, KM817703

<p>Reasons to justify the creation and assignment of the new species:</p> <ul style="list-style-type: none"> • Explain how the proposed species differ(s) from all existing species. <ul style="list-style-type: none"> ○ If species demarcation criteria (see module 3) have previously been defined for the genus, explain how the new species meet these criteria. ○ 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 11 <p>See genus justification.</p>
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MODULE 3: **NEW GENUS**

creating a new genus

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

Code	2016.022bM	(assigned by ICTV officers)
To create a new genus within:		
Subfamily:	unassigned	Fill in all that apply. • If the higher taxon has yet to be created (in a later module, below) write “(new)” after its proposed name. • If no family is specified, enter “unassigned” in the family box
Family:	<i>Phenuiviridae</i> (NEW, see TP 2016.030M)	
Order:	<i>Bunyavirales</i> (NEW, see TP 2016.030M)	

naming a new genus

Code	2016.022cM	(assigned by ICTV officers)
To name the new genus: <i>Goukovirus</i>		

Assigning the type species and other species to a new genus

Code	2016.022dM	(assigned by ICTV officers)
To designate the following as the type species of the new genus		
<i>Gouleako goukovirus</i>		Every genus must have a type species. This should be a well characterized species although not necessarily the first to be discovered
The new genus will also contain any other new species created and assigned to it (Module 2) and any that are being moved from elsewhere (Module 7b). Please enter here the TOTAL number of species (including the type species) that the genus will contain:		
3		

Reasons to justify the creation of a new genus:

Additional material in support of this proposal may be presented in the Appendix, Module 11

<p>Gouléako virus (GOLV) is most similar to the members of the accepted bunyaviral genus <i>Phlebovirus</i>. However, GOLV is significantly different from all known bunyaviruses (including phleboviruses) as listed below. A phylogenetic analyses is presented in the Appendix, Module 9.</p> <ul style="list-style-type: none"> • The GOLV genome is shorter than that of any known bunyavirus. • The GOLV genome termini are most similar to those of phleboviruses. <p>However, S and M segment termini are shorter with a length of only 5 instead of 8 nt. Genome termini are generally conserved within but invariably different between bunyavirus genera.</p> <ul style="list-style-type: none"> • GOLV has a different genome organization than phleboviruses. In contrast to phleboviruses, there is no evidence for encoded NSs and NSm proteins. • Only weak sequence similarity between GOLV and phleboviruses was identified. Amino acid pairwise identities between GOLV and the closest relatives, Uukuniemi virus (UUKV) and severe fever with thrombocytopenia syndrome virus (SFTSV), are 28% and 27% for the L protein, 21% and 24% for the glycoproteins, and 24% and 25% for the N protein, respectively. • GOLV does not serological cross-react with phleboviruses. According to the ICTV species demarcation criteria in the genus <i>Phlebovirus</i>, phleboviruses are defined by the
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serological relationships. Phleboviruses are antigenically unrelated to members of other genera but cross-react serologically among themselves. • GOLV seems to have a host range limited to insects and thus differs from phleboviruses that can in addition infect vertebrates. To date, GOLV has only been found in mosquitoes and does only replicate in mosquito but not in vertebrate cells.

Origin of the new genus name:

From Gouléako, the village from where the virus-positive mosquitoes originated.

Reasons to justify the choice of type species:

First virus of this group that was discovered.

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.

Until further goukoviruses are discovered, we propose to use the same species demarcation criteria for this genus as described for the proposed new genera “*Orthoferavirus*” and “*Herbevirus*” (see separate co-submitted proposals). Species demarcation criteria should be based on a ≈ 1 kb sequence fragment containing the core polymerase domain (premotif A to motif E) of the third conserved region of the L protein. These motifs can be aligned between all members of the proposed order *Bunyavirales* and would allow comparative species demarcation criteria for all genera of the entire family. Moreover, as the motifs are highly conserved between all bunyaviruses, amplification of this genome region from new viruses is facilitated. Species demarcation criteria of other viral families are also based on the replicative genes/domains and have been shown to be suitable criteria.

Species should be defined on the criterion that the ≈ 1 kb sequence fragment containing the core polymerase domain (premotif A to motif E) of the third conserved region of the L protein should be less than 90% identical on the amino acid level compared to that of any other described goukovirus.

This <90% aa identity threshold for the core polymerase domain is in agreement with the aa identity values for established bunyavirus species within the five established genera.

MODULE 11: **APPENDIX**: supporting material

additional material in support of this proposal

References:

Marklewitz M., Handrick S., Grasse W., Kurth A., Lukashev A., Drosten C., Ellerbrok H., Leendertz F.H., Pauli G., Junglen S. 2011. Gouleako virus isolated from West African mosquitoes constitutes a proposed novel genus in the family Bunyaviridae. *Journal of Virology* 85: 9227-9234.

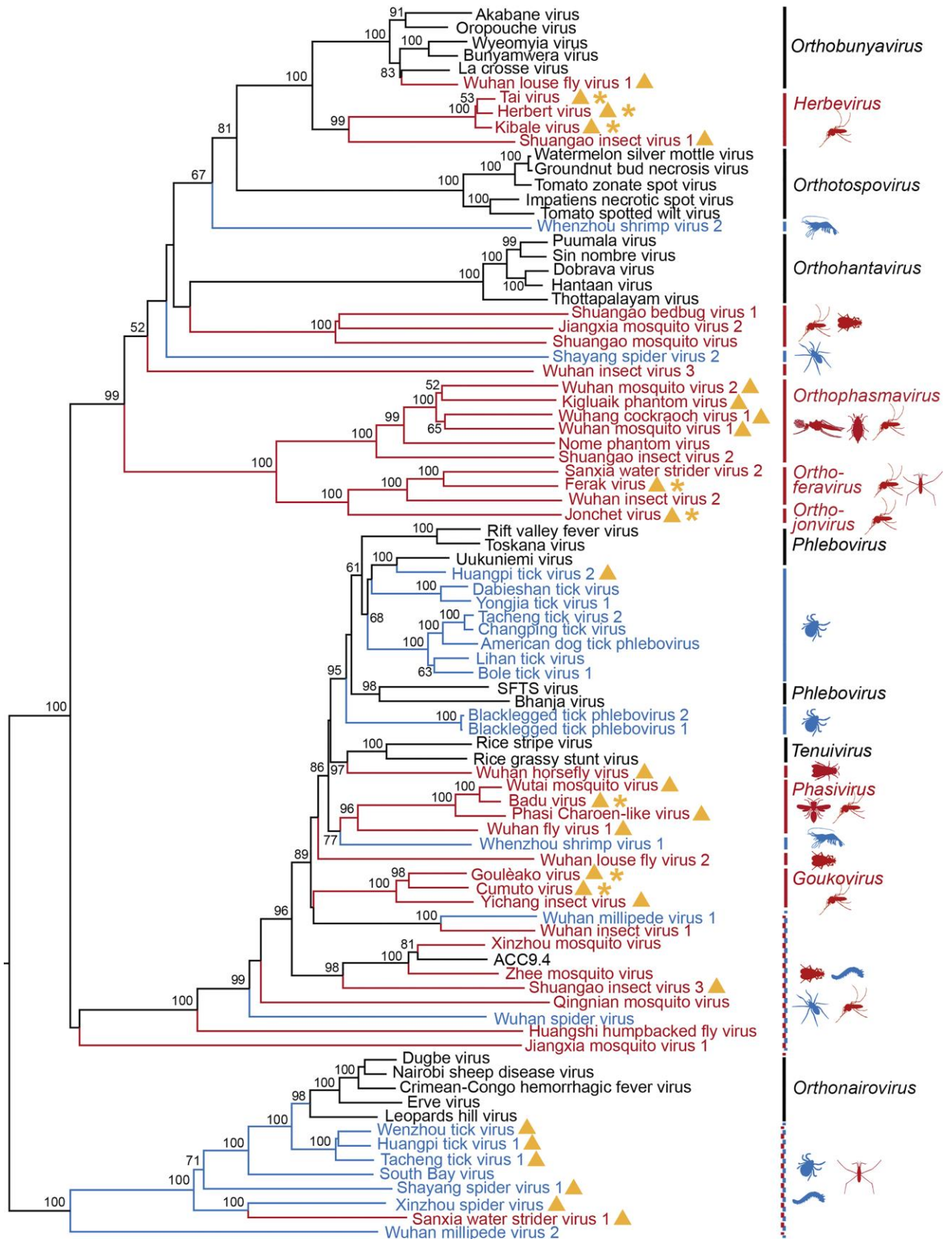
Marklewitz M., Zirkel F., Rwego I.B., Heidemann H., Trippner P., Kurth A., Kallies R., Briese T., Lipkin W.I., Drosten C., Gillespie T.R., Junglen S. 2013. Discovery of a Unique Novel Clade of Mosquito-Associated Bunyaviruses. *Journal of Virology* 87: 12850-12865.

Marklewitz M., Zirkel F., Kurth A., Drosten C., Junglen S. 2015. Evolutionary and phenotypic analysis of live virus isolates suggests arthropod origin of a pathogenic RNA virus family. *Proceedings of the National Academy of Sciences* 112: 7536-41.

Junglen S. (2016). Evolutionary origin of pathogenic arthropod-borne viruses — a case study in the family *Bunyaviridae*. *Current Opinion in Insect Science* 16: 81-86.

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.



Tick
 Shrimp
 Mosquito
 Horsefly
 Fly
 Louse fly / bed bug
 Spider
 Millipede
 Cockroach
 Water strider
 Phantom midge
 * Live virus isolate
 ▲ Entire coding sequence

Figure: Phylogenetic relationship of bunyaviruses. Phylogenetic analyses were based on RdRp proteins. Complete RdRp proteins were aligned using MAFFT (E-INS-I algorithm). Alignment columns were stripped to 10% gaps in Geneious. Maximum likelihood (ML) analyses were performed on a 508 amino acid alignment guided by the Blosum62 amino acid substitution matrix with 4 gamma categories and a gamma shape parameter of 1. Confidence testing was performed by 1000 bootstrap replicates. Only bootstrap values over 50 are shown.