

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:	2015.0200	a-aeB	(to be completed by ICTV officers)			
Short title: To amend the men in the subfamily <i>Tevenvirinae</i> (e.g. 6 new species in the genus A <b>Modules attached</b> (modules 1 and 10 are required)	1 0	genus <i>T4li</i> 1 ⊠ 6 □	ikevirus, a 2 🖂 7 🖂	and create 3 ⊠ 8 □	six (6) ne 4 9	w genera 5 □ 10 ⊠

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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 <u>http://www.ictvonline.org/subcommittees.asp</u> . If in doubt, contact the appropriate subcommittee chair (fungal, invertebrate, plant, prokaryote or vertebrate viruses)	Bacterial and Archaeal Virus Subcommittee
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### ICTV Study Group comments (if any) and response of the proposer:

Please note that we have chosen to refer to this genus as *T4virus* rather than *T4likevirus* since the Bacterial and Archaeal Virus Subcommittee of ICTV has voted overwhelmingly in favour of eliminating *"like"* from phage genus names.

Date first submitted to ICTV: Date of this revision (if different to above): May 2015

### **ICTV-EC** comments and response of the proposer:

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 201	5.020aB	(assigned by ICTV	office	rs)		
To create 10 new species within:						
Genus:T4likevirus (proposed name T4virus)Subfamily:TevenvirinaeFamily:MyoviridaeOrder:Caudovirales						
Name of new species: Representative isolate per species please)			nly 1	GenBank sequence accession number(s)		
Escherichia virus Ime09Escherichia phage ime0Escherichia virus E112Escherichia phage vB_IEscherichia virusEscherichia phage ECNECML134Escherichia virus RB3Escherichia virus C40Escherichia phage RB3VB_EcoM_ACG-C40		Escherichia phage		AP011113 JN202312 KJ668714.2 JX128259 KM606994 JN986846 HM035025		
Shigella virus Pss1Shigella phage Shil2Shigella virus Pss1Shigella phage pSs-1Yersinia virus PSTYersinia phage PSTYersinia virus D1Yersinia phage phiD1		Yersinia phage PST		KM501444 KF208315 HE956711		

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

- Explain how the proposed species differ(s) from all existing species.
  - 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 9

creating a new genus

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

Code 20	015.020bB	(assigned by ICTV officers)		
To create a ne	ew genus within:	Fill in all that apply.		
Subfamily	y: Tevenvirinae	If the higher taxon has yet to be created		
Family	y: Myoviridae	(in a later module, below) write "(new)" after its proposed name.		
Order	r: Caudovirales	<ul> <li>If no family is specified, enter</li> <li>"unassigned" in the family box</li> </ul>		

naming a new genus

Code	2015.020cB	(assigned by ICTV officers)
To name tl	ne new genus: <i>Rb49virus</i>	

Assigning the type species and other species to a new genus

Code	2015.020dB	(assigned by ICTV officers)				
To desig	nate the following as the type sp	ecies o	f the new genus			
<i>Escherichia phage RB49</i> (existing species) to be renamed <i>Escherichia virus RB49</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:						

### **Reasons to justify the creation of a new genus:**

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

This new genus will contain three species (*Escherichia virus RB49*, *Escherichia virus phi1 and Escherichia virus JSE*) that were previously classified as t4likeviruses and are now moved into the new genus. "Its [RB49] morphology is indistinguishable from the well-known T-even phage T4, but DNA hybridization indicated that it was phylogenetically distant from T4 and thus it was classified as a pseudo-T-even phage." (1). Phages of this group have 164.9 kb genomes with a mol%G+C of 40.5. They encode approximately 277 proteins and no tRNAs. Their DNA contains cytosine and not 5hydroxymethylcytosine. They only share 41% protein homologs with *Escherichia* phage T4. Our phylogenetic analysis (Fig. 1) and BLASTN analysis (Table 1) indicate further that this group of phages is separate from t4viruses.

#### **Origin of the new genus name:**

Named after the first virus of its type to be sequenced - Escherichia phage RB49

#### **Reasons to justify the choice of type species:**

The first virus of its type to be sequenced - Escherichia phage RB49

#### **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. Each of the proposed species differs from the others with more than 5% at the DNA level as confirmed with the BLASTN algorithm.

Please note that we have chosen to refer to this new genus as *Rb49virus* rather than *Rb49likevirus* since the Bacterial and Archaeal Virus Subcommittee of ICTV has voted overwhelmingly in favour of eliminating *"like"* and *"Phi"* from phage genus names.

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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	Code 2015.020eB (assigned by IC			ICTV office	rs)		
To crea	To create 3 new species within:						
Genus: <i>Rb69virus</i> (new)					<ul><li>Fill in all that apply.</li><li>If the higher taxon has yet to be</li></ul>		
Subfamily: <i>Tevenvirinae</i>					<ul> <li>created (in a later module, below) write</li> <li>"(new)" after its proposed name.</li> <li>If no genus is specified, enter</li> </ul>		
Fa	Family: <i>Myoviridae</i>						
(	Order:	Caudovirales	•		"unassigned" in the genus box.		
		<b>Representative isolate</b> per species please)	e: (only 1	GenBank sequence accession number(s)			
		Escherichia phage vB_EcoM_JS09		KF582788			
Escherichia virus HX01 Escherich		Escherichia phage HX Shigella phage Shf125		JX536493 KM407600			

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

• Explain how the proposed species differ(s) from all existing species.

- 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 9

creating a new genus

Ideally, a g	genus sho	uld be placed within a	higher taxon.
Code	201	5.020fB	(assigned by ICTV officers)
To crea	ite a new	genus within:	Fill in all that apply.
Su	ofamily:	Tevenvirinae	If the higher taxon has yet to be created     (in a later module halow) write "(result)"
	Family:	Myoviridae	(in a later module, below) write "(new)" after its proposed name.
	Order:	Caudovirales	<ul> <li>If no family is specified, enter</li> <li>"unassigned" in the family box</li> </ul>

naming a new genus

Code	2015.020gB	(assigned by ICTV officers)
To name tl	he new genus: <i>Rb69virus</i>	

Assigning the type species and other species to a new genus

Code	2015.020hB (assig		igned by ICTV officers)		
To designate the following as the type species of the new genus					
Escherichia virus RB69			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: 4

**Reasons to justify the creation of a new genus:** 

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

A long recognized member of a subgroup of T4-like phages (5) RB69 has been described as a "distant phylogenetic relative of T4" (6). Phages of this group have 168.7kb genomes with a mol%G+C of 37.6. They encode approximately 271 proteins and 2 tRNAs. They only share 74.7% protein homologs with *Escherichia* phage T4. Our phylogenetic analysis (Fig. 1) and BLASTN analysis (Table 1) indicate further that this group of phages is separate from *T4virus*.

Recently two *E.coli* O157 typing phages 3 (KP869101) & 6 (KP869104) have been characterized (11) and can be added to this genus.

#### **Origin of the new genus name:**

Named after the first virus of its type to be sequenced - *Escherichia* phage RB69

#### Reasons to justify the choice of type species:

The first virus of its type to be sequenced - *Escherichia* phage RB69

#### 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. Each of the proposed species differs from the others with more than 5% at the DNA level as confirmed with the BLASTN algorithm.

Please note that we have chosen to refer to this new genus as *Rb69virus* rather than *Rb69likevirus* since the Bacterial and Archaeal Virus Subcommittee of ICTV has voted overwhelmingly in favour of eliminating *"like"* and *"Phi"* from phage genus names.

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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	Code 2015.020iB (assigned by ICT			TV office	rs)		
To crea	To create 4 new species within:						
Genus: Js98virus (new)					<ul><li>Fill in all that apply.</li><li>If the higher taxon has yet to be</li></ul>		
Subfamily: <i>Tevenvirinae</i>				ated (in a later module, below) write			
Fa	Family: Myoviridae			<ul><li>"(new)" after its proposed name.</li><li>If no genus is specified, enter</li></ul>			
0	Order:	Caudovirales			"unassigned" in the genus box.		
Name of new species: Represer per specie		ntative isolate: es please)	(only 1	GenBank sequence accession number(s)			
Escherio	chia vi	rus IME08	Escheric	chia phage IME08		HM071924	
Escherio	<i>Escherichia virus Bp7</i> Escherichia		hia phage Bp7		HQ829472		
Escherichia virus JS10 Escherichia		hia phage JS10		EU863409			
Escherie	Escherichia virus VR5 Escherich		hia phage vb_EcoM-		KP007359		
			VR5				

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

• Explain how the proposed species differ(s) from all existing species.

- 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 9

creating a new genus

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

Code 20	015.020jB	(assigned by ICTV officers)		
To create a no	ew genus within:	Fill in all that apply.		
Subfamily	y: <b>Tevenvirinae</b>	If the higher taxon has yet to be created		
Family	y: Myoviridae	(in a later module, below) write " <b>(new)</b> " after its proposed name.		
Orde	r: Caudovirales	<ul> <li>If no family is specified, enter</li> <li>"unassigned" in the family box</li> </ul>		

naming a new genus

Code	2015.020kB	(assigned by ICTV officers)
To name tl	ne new genus: Js98virus	

Assigning the type species and other species to a new genus

Code	2015.020lB	(assigned by ICTV officers)		
To designate the following as the type species of the new genus				
Escherichia virus JS98		Every genus must have a type species. This should be a well characterized species although not necessarily the first to be discovered		
The new g	enus will also contain any other r	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: 5 5

**Reasons to justify the creation of a new genus:** 

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

Zuber et al. (5) stated "JS98 defines a new major subgroup of *E. coli* T4-like phages." Phages of this group possess 170.6 kb genomes (39.5 mol%G+C) and encode 262 proteins and usually 3 tRNAs. They only share 77.4% protein homologs with Escherichia phage T4. Our phylogenetic analysis (Fig. 1) and BLASTN analysis (Table 1) indicate further that this group of phages is separate from T4virus. A recent isolate, *Escherichia* virus RV5, is a member of this genus.

Origin of the new genus name:

Named after the first virus of its type to be sequenced - Escherichia phage JS98

Reasons to justify the choice of type species:

The first virus of its type to be sequenced - Escherichia phage JS98

#### 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.

Please note that we have chosen to refer to this new genus as *Js98virus* rather than *Js98likevirus* since the Bacterial and Archaeal Virus Subcommittee of ICTV has voted overwhelmingly in favour of eliminating *"like"* and *"Phi"* from phage genus names.

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 2015.020mB (assigned by 10			CTV office	rs)
To create 5 n	ew species wi	thin:		
Genus:Sp18virus (new)Subfamily:TevenvirinaeFamily:MyoviridaeOrder:Caudovirales			<ul> <li>Fill in all that apply.</li> <li>If the higher taxon has yet to be created (in a later module, below) write "(new)" after its proposed name.</li> <li>If no genus is specified, enter "unassigned" in the genus box.</li> </ul>	
-		Representative isolate per species please)	: (only 1	GenBank sequence accession number(s)
Shigella virus Escherichia vi		Shigella phage SP18 Escherichia phage vB EcoM VR7		GQ981382 HM563683
Escherichia virus VR20		Escherichia phage vB_EcoM_VR20		KP007360
Escherichia virus VR25 Escherichia virus VR26		Escherichia phage vB_EcoM_VR25 Escherichia phage		KP007361 KP007362
		vB_EcoM_VR26		

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

• Explain how the proposed species differ(s) from all existing species.

- 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 9

creating a new genus

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

Code 20.	15.020nB	(assigned by ICTV officers)		
To create a new	w genus within:	Fill in all that apply.		
Subfamily:	Tevenvirinae	If the higher taxon has yet to be created		
Family:	Myoviridae	(in a later module, below) write "(new)" after its proposed name.		
Order:	Caudovirales	<ul> <li>If no family is specified, enter</li> <li>"unassigned" in the family box</li> </ul>		

naming a new genus

Code	2015.020oB	(assigned by ICTV officers)
To name the new genus: Sp18virus		

Assigning the type species and other species to a new genus

Code	2015.020pB	(assigned by ICTV officers)			
To designa	To designate the following as the type species of the new genus				
Shigella virus SP18		Every genus must have a type species. This sho 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). <b>Please enter here the TOTAL number of species</b>					

(including the type species) that the genus will contain: 5 5

**Reasons to justify the creation of a new genus:** 

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

All VR phages fail to produce plaques at 37C, and that they are *E*.*coli* strain specific (VR7 and VR20 prefer BE strains while VR25 and VR26 form plaques on K12 strains only). Restriction enzymes that are known to recognize 5-hydroxymethylcytosine containing DNA (EcoRV, DraI, VspI, NdeI) cut the DNA suggesting to the authors that the DNA of phage VR7 has a similar modification to T4, yet the genomes failed to reveal alpha- or beta-glucosyltransferases or dCMP hydroxymethylase homologs (8). Phage SP18 was isolated from the Gap River in Korea (7) and lysed *Shigella sonnei*, but not *S. flexneri*, *S. boydii* or members of the genera *Escherichia* and *Salmonella*. Two common protein markers for these two phages are "hypothetical protein VR7\_gp288"-"hypothetical protein SP18\_gp282" and "hypothetical protein VR7\_gp147"-"hypothetical protein SP18\_gp148." Subsequent to our overall analysis three related phages were deposited by Kaliniene et al. *- Escherichia* phages vB\_EcoM\_VR20, vB\_EcoM\_VR25 and vB\_EcoM\_VR26.

These phages have genomes of approximately 170.0 kb with a GC content of 40.4%. They encode approximately 289 proteins and 2 tRNAs. Relative to coliphage T4 SP18 shares 71.8% proteins.

### **Origin of the new genus name:**

Named after the first virus of its type to be sequenced - Shigella phage S18

### Reasons to justify the choice of type species:

The first virus of its type to be sequenced - Shigella phage S18

### **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. Each of the proposed species differs from the others with more than 5% at the DNA level as confirmed with the BLASTN algorithm.

Please note that we have chosen to refer to this new genus as *Sp18virus* rather than *Sp18likevirus* since the Bacterial and Archaeal Virus Subcommittee of ICTV has voted overwhelmingly in favour of eliminating "like" and "Phi" from phage genus names.

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	201	015.020qB (assigned by IC		CTV office	ers)
To crea	ate 2 n	ew species wit	thin:		
Genus: S16virus (new)			w)	<ul><li>Fill in all that apply.</li><li>If the higher taxon has yet to be</li></ul>	
	amily:	Tevenvirinae	,		ated (in a later module, below) write
F	amily:	y: Myoviridae		<ul><li>"(new)" after its proposed name.</li><li>If no genus is specified, enter</li></ul>	
	Order:	Caudovirales		"unassigned" in the genus box.	
Name (	of new a	species:	Representative isolate per species please)	: (only 1	GenBank sequence accession number(s)
Salmonella virus S16			Salmonella phage STM Salmonella phage vB_S S16		JX181825 HQ331142

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

- Explain how the proposed species differ(s) from all existing species.
  - 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 9

We have chosen 95% DNA sequence identity as the criterion for demarcation of species in this new genus. Each of the proposed species differs from the others with more than 5% at the DNA level as confirmed with the BLASTN algorithm. Under these conditions, *Salmonella* phage STP4-a is considered a strain.

creating a new genus

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

 Code
 2015.020rB

 (assigned by ICTV officers)

To create a new	genus within:	Fill in all that apply.
Subfamily:	Tevenvirinae	If the higher taxon has yet to be creat
Family:	Myoviridae	<ul> <li>(in a later module, below) write "(new)"</li> <li>after its proposed name.</li> </ul>
Order:	Caudovirales	<ul> <li>If no family is specified, enter</li> <li>"unassigned" in the family box</li> </ul>

naming a new genus

Code	2015.020sB	(assigned by ICTV officers)

To name the new genus: *S16virus* 

Assigning the type species and other species to a new genus

Code 2015.020tB

(assigned by ICTV officers)

# To designate the following as the type species of the new genus

Salmonella virus S16

2

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: 2

Reasons to justify the creation of a new genus:

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

Virulent phage S16 is the second T-even-like phage specific for Salmonella. "It can infect a larger variety of *Salmonella* strains than Felix-O1" (9). Its physical dimensions are: head, 117 by 91; and, a contractile tail 120 nm in length. It can infect LPS 'Re' mutants and binds to OmpC. The "presence of DNA modification functions (dCMP hydroxymethylase,  $\beta$ -glucosyltransferase and  $\beta$ -glucosyl-HMC- $\alpha$ -glucosyl-transferase)" leads to resistance to most restriction endonucleases. There is no manuscript associated with *Salmonella* phage STML-198. The genomes of this genus are characterized by an average size of 159.2 kb (36.9 mol%G+C), encoding 262 proteins and 3 tRNAs. Using CoreGenes S16 shares 62.3% homologs with T4.

### **Origin of the new genus name:**

Named after the *Salmonella* phage S16

### **Reasons to justify the choice of type species:**

This is the only phage of the pair which has been fully described (9).

### 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. Each of the proposed species differs from the others with more than 5% at the DNA level as confirmed with the BLASTN algorithm.

Please note that we have chosen to refer to this new genus as *S16virus* rather than *S16likevirus* since the Bacterial and Archaeal Virus Subcommittee of ICTV has voted overwhelmingly in favour of eliminating "like" and "Phi" from phage genus names.

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	Code 2015.020uB (assigned by IC			TV office	rs)
To crea	ate 2 no	ew species wit	hin:		
Genus: Cc31virus (new)				<ul><li>Fill in all that apply.</li><li>If the higher taxon has yet to be</li></ul>	
	amily:			<ul> <li>created (in a later module, below) write</li> <li>"(new)" after its proposed name.</li> <li>If no genus is specified, enter</li> </ul>	
	amily:				
(	Order:	Caudovirales		"unassigned" in the genus box.	
Name of new species:		species:	<b>Representative isolate:</b> per species please)	(only 1	GenBank sequence accession number(s)
Enterobacter virus CC31 Enterobacter virus PG7			Enterobacter phage CC3 Enterobacter phage PG7		GU323318 KJ101592

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

- Explain how the proposed species differ(s) from all existing species.
  - 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 9

creating a new genus

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

Code	201	5.020vB	(assigned by ICTV officers)		
To create a	new	genus within:		Fill in all that apply.	
Subfam	nily:	Tevenvirinae		• If the higher taxon has yet to be created	
Fam	nily:	Myoviridae		(in a later module, below) write " <b>(new)</b> " after its proposed name.	
Or	der:	Caudovirales		<ul> <li>If no family is specified, enter</li> <li>"unassigned" in the family box</li> </ul>	

naming a new genus

Code	2015.020wB	(assigned by ICTV officers)
To name tl	ne new genus: <i>Cc31virus</i>	

Assigning the type species and other species to a new genus

Code	2015.020xB	(assigned by ICTV officers)				
To designate the following as the type species of the new genus						
Enterobacter virus CC31		Every genus must have a type species. This should be a well characterized species although not necessarily the first to be discovered				
		v 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: 2

**Reasons to justify the creation of a new genus:** 

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

The two members of the Cc31virus genus are lytic for *Enterobacter* species and are characterized by possessing genomes with the following average properties – size: 169.4 kb; mol%G+C: 39.9; number of proteins encoded: 287; number of tRNAs: 14. Percentage of CC31 proteins which find homologs in T4: 57.0.

### **Origin of the new genus name:**

Enterobacter phage CC31

### **Reasons to justify the choice of type species:**

The first virus of its type to be sequenced

### 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.

Please note that we have chosen to refer to this new genus as *Cc31virus* rather than *Cc31likevirus* since the Bacterial and Archaeal Virus Subcommittee of ICTV has voted overwhelmingly in favour of eliminating "like" and "Phi" from phage genus names.

### MODULE 7: **<u>REMOVE and MOVE</u>**

Use this module whenever an existing taxon needs to be removed:

- Either to abolish a taxon entirely (when only part (a) needs to be completed)
- Or to move a taxon and re-assign it e.g. when a species is moved from one genus to another (when BOTH parts (a) and (b) should be completed)

#### Part (a) taxon/taxa to be removed or moved

Code	2015.	020yB	(assigned by ICTV officers)			
To remove the following taxon (or taxa) from their present position:						
Aeromo	nas phage	25				
Aeromo	nas phage	31				
Aeromo	nas phage	44RR2.8t				
Enterob	acteria ph	age SV14				
Escheri	chia phage	e JS98				
Escherie	chia phage	e phi l				
Escherie	chia phage	e JSE				
	chia phage					
	chia phage					
	chia phage					
	chia phage					
	chia phage					
Pseudor	nonas pha	ge 42				
The pre	esent taxo	nomic position (	of these taxon/taxa:			
C	Genus: T4	Alikevirus				
Subfa	amily: <b>Te</b>	evenvirinae	Fill in all that apply			
Fa	amily: M	yoviridae	Fill in all that apply.			
(	Order: Co	audovirales				
			a not received to enother tower) write "wee"			

If the taxon/taxa are to be abolished (i.e. not reassigned to another taxon) write "yes" in the box on the right

#### **Reasons to justify the removal:**

Explain why the taxon (or taxa) should be removed

Based predominantly on the sequence of the major capsid protein (gp23) the T4-related phages were assigned to five groups: T-evens (T4, RB69), pseudoT-evens (AR1, RB49, RB42, RB43), schizoT-evens (nt-1, KVP20, 65, and Aeh1), JS98 and the exoT-even phages (cyanomyoviruses) [1-4]. Petrov et al. [5], re-examined the T4-related bacteriophages which they state contain genomes ranging from approximately 157-255 kb, and sharing a "core genome" of approximately 40 genes involved in nucleotide metabolism, DNA replication, repair, recombination and packaging, transcriptional regulation and morphogenesis. Their analysis indicates that this assemblage includes not only well-characterized coliphages but also viruses infecting *Acinetobacter* (e.g. 133), *Aeromonas* (44RR2.8t), *Campylobacter* (CP220), *Delftia* (φW-14), *Klebsiella* (KP15), *Prochlorococcus* (P-SSM2), *Salmonella* (ViI), *Shigella* (φSboM-AG3), *Synechococcus* (Syn9), and *Vibrio* (KVP40). The diversity displayed by these viruses is equivalent to that of the *Herpesvirales*. Previous ICTV proposals have identified a new genus *Viunalikevirus* and a subfamily the *Eucampyvirinae* (*Cp220likevirus*, *Cp8unalikevirus*) which, at some taxonomic level, fall within the "T4

superfamily."

According to the latest ICTV taxonomy report the subfamily *Tevenvirinae* contains only two genera – Schizot4likevirus (one species, Vibrio phage nt-1) and the T4likevirus of which the following phages are members: Aeromonas phages 25, 31 and 44RR2.8t, Escherichia phages T4, JS98, phi1, RB14, RB16, RB32, RB43, RB49, RB69 and SV14; and, Pseudomonas phage 42. The question here is not the total diversity of the T4-like phages but what viruses actually constitute the T4likevirus. In the past the Bacterial and Archaeal Virus Subcommittee has relied on percent homologous proteins to group phages, but with increasing deposits of complete phage genomes to public databases we are seeing that this results in "lumping" of species which can be clearly distinguished on the basis of numerous criteria, including DNA sequence identity - the gold standard to bacterial classification. Using the latter parameter as the primary taxonomic tool we have seen the suggestion made that the *Tunalikevirus* [7] and the *N4likevirus* (J. Whitman, submitted manuscript) are not monophyletic. With many more T4-like phages sequenced we have reassessed their This has been accomplished through BLASTN analysis in which we calculated taxonomy. the total sequence identity by multiplying the % coverage by the % identity; total proteome by TBLASTX comparisons and CoreGenes3.0 [8]; and a phylogenetic analysis of the large subunit terminase protein using phylogeny.fr [9]. It is clear that many of the phages listed as members of the T4likevirus are sufficiently different to warrant reclassification.

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10. Lavigne R, Darius P, Summer EJ, Seto D, Mahadevan P, Nilsson AS, Ackermann H-W, Kropinski AM: Classification of *Myoviridae* bacteriophages using protein sequence similarity. BMC Microbiology 2009, 9:224.

**Part (b)** re-assign to a higher taxon

	to a higher taxon				
	.020zB		ned by ICTV officers)		
<u> </u>	axon (or taxa) listed in Part (a) as				
1 0	RB49 (proposed name Escherichia	virus	Fill in all that apply.		
<i>RB49</i> )			<ul><li>If the higher taxon has yet to</li></ul>		
	phil (proposed name Escherichia v	irus	be created write "(new)"		
phil) Eacharichia naca l	SE (proposed nome Each grichig wir		after its proposed name and		
Genus:	SE (proposed name Escherichia vir <b>Rb49virus</b> (new)	us JSE)	complete relevant module to create it.		
Subfamily:	<i>Tevenvirinae</i>		If no genus is specified, enter		
Family:	Myoviridae		"unassigned" in the genus		
Order:			box.		
	.020aaB	(assigne	ed by ICTV officers)		
			a by icity oncers)		
0	axon (or taxa) listed in Part (a) as <i>RB69</i> (proposed name <i>Escherichia</i>		Fill in all that apply.		
RB69)	RB09 (proposed name Escherichid	virus	<ul> <li>If the higher taxon has yet to</li> </ul>		
Genus:	<i>Rb69virus</i> (new)		be created write "(new)"		
Subfamily:	Tevenvirinae		after its proposed name and complete relevant module to		
Family:	Myoviridae		create it.		
Order:	Caudovirales		If no genus is specified, enter		
			" <b>unassigned</b> " in the genus box.		
Code 2015	.020abB	(assigne	ed by ICTV officers)		
	axon (or taxa) listed in Part (a) as				
9	JS98 (proposed name Escherichia v		Fill in all that apply.		
JS98)			• If the higher taxon has yet to		
Genus:	Js98virus (new)		be created write " <b>(new)</b> " after its proposed name and complete relevant module to create it.		
Subfamily:	Tevenvirinae				
Family:	Myoviridae				
Order:	Caudovirales	If no genus is specified, enter			
			" <b>unassigned</b> " in the genus box.		
Code 2015	.020acB	(assigne	ed by ICTV officers)		
	axon (or taxa) listed in Part (a) as				
<u> </u>	<i>ige SV14</i> (proposed name <i>Enterobac</i>		La construction de la constructi		
virus SV14)	ige 5717 (proposed nume Lineroota				
/	RB16 (proposed name Escherichia	virus			
RB16)			Fill in all that apply.		
Escherichia phage	RB32 (proposed name Escherichia	virus	• If the higher taxon has yet to		
<i>RB32</i> )			be created write "(new)"		
	RB43 (proposed name Escherichia	virus	after its proposed name and complete relevant module to		
<i>RB43</i> )		1	create it.		
	ge 42 (proposed name Pseudomonas	phage	If no genus is specified, enter		
42)	Unaggiore of	]	" <b>unassigned</b> " in the genus box.		
Genus:	Unassigned Tevenvirinae		50A.		
Subfamily: Family:	Nyoviridae		-		
Order:	Caudovirales				
Uluel.	Canaorianes				

**Part (b)** re-assign to a higher taxon

Code 20	15.020adB	(assigned by ICTV officers)						
To re-assign	To re-assign the taxon (or taxa) listed in Part (a) as follows:							
Aeromonas p	Aeromonas phage 25 (proposed name Aeromonas							
virus 25)		Fill in all that apply.						
Aeromonas p	hage 31 (proposed name A	eromonas • If the higher taxon has yet to be						
virus 31)		created write "(new)" after its						
Genus	: Secunda5virus (new)	proposed name and complete relevant module to create it.						
Subfamily	:	If no genus is specified, enter						
Family	: Myoviridae	"unassigned" in the genus box.						
Order	: Caudovirales							

### Part (b) re-assign to a higher taxon

Code 20	)15.020aeB	(assigned by ICTV officers)			
0	the taxon (or taxa) listed bhage 44RR2.8t (proposed r virus 44RR2)				
Genus Subfamily		proposed name and complete relevant module to create it.			
Family Order	r: Myoviridae	If no genus is specified, enter " <b>unassigned</b> " in the genus box.			
•					

### **Reasons to justify the re-assignment:**

- If it is proposed to re-assign species to an existing genus, please explain how the proposed species differ(s) from all existing species.
  - 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.
- Provide accession numbers for genomic sequences

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

See above

### MODULE 10: APPENDIX: supporting material

additional material in support of this proposal

#### **References:**

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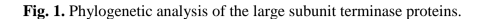
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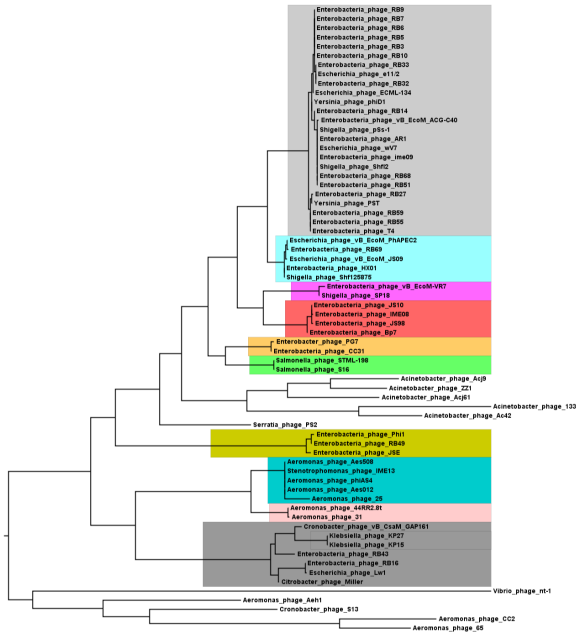
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#### Annex:

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.





0.05

## Table 1. BLASTN analyses

	T4	PST	Shfl2	e11/2	phiD1	ECML-134	RB3	RB14	AR1	ime09	RB32	ACG-C40	pSs-1
<b>T</b> 4	100	87	87	86	87	86	87	86	85	85	84	82	81
PST	87	100	89	84	89	91	86	93	88	89	91	83	86
Shfl2	86	88	100	86	91	89	89	90	91	94	89	84	91
e11/2	86	84	87	100	87	87	87	88	90	86	89	87	90
phiD1	86	88	92	86	100	86	89	90	90	91	90	85	89
ECML-134	86	91	90	86	86	100	90	92	89	86	92	89	89
RB3	86	86	90	87	90	91	100	93	88	88	91	88	88
RB14	84	91	90	87	89	91	91	100	90	89	92	88	90
AR1	84	87	92	89	91	89	87	91	100	87	89	86	93
ime09	84	89	94	85	91	86	97	90	86	100	89	82	92
RB32	82	90	89	87	89	91	89	92	88	88	100	88	89
vB_EcoM_ACG-C40	81	83	85	87	85	89	88	89	86	83	89	100	87
pSs-1	79	85	90	87	88	87	86	90	91	91	89	86	100
vB_EcoM_JS09	57	53	58	55	60	53	53	58	57	59	57	56	54
HX01	56	55	56	53	62	53	53	55	65	56	54	56	55
Shf125875	54	53	56	56	58	54	53	58	58	56	58	57	56
RB69	54	53	56	56	57	53	53	58	57	56	57	56	56
vB_EcoM_PhAPEC2	53	54	57	56	57	54	53	58	58	57	58	57	56
IME08	45	44	46	43	44	43	44	45	44	47	45	44	45
JS98	43	44	45	41	45	43	43	45	45	45	44	44	46
Bp7	43	43	43	49	47	42	42	43	43	43	43	43	45
JS10	43	43	44	40	44	43	42	43	44	43	43	43	45
SP18	42	42	52	42	54	52	41	52	53	53	52	41	54
vB_EcoM-VR7 S16	41 40	41 40	41 39	41 39	41 39	41 39	41 39	42 41	41 39	43 41	41 39	41 40	42
STML-198	40	40	39	39	39	39	39	41 42	39	41	39	40	41
PG7	38	39	40	30	40	40	39 41	39	39	42	39	39	41
CC31	30	38	39	38	39	38	39	39	39	40	39	39	41
JSE	14	14	15	14	13	14	14	13	13	13	14	13	13
RB49	14	14	14	14	13	14	14	14	13	14	14	14	12
Phi1	12	14	14	13	15	14	14	14	15	14	14	14	14
	JS09	HX01			RB69	PhAPEC2	IME08	JS98	Bp7	JS1		SP18	VR7
T4	58	5		54	54	54	45	43	4		42	42	42
PST	53		4	53	53	54	43	44	4		43	41	41
Shfl2	57		5	55	56	56	45	44	4		43	52	41
e11/2	56		3	56	56	56	43	41	5		40	42	41
phiD1	59	6		57	57	57	44	44	4		43	53	41
ECML-134	53		3	53	53	54	43	43	4	2	42		
RB3	53	5	3	53	53	50				_	42	51	41
RB14	57					53	43	43	4		42	51 41	41 41
AR1	57		4	58	58	53 58	43 44	43 44	4	3			
	_	5	4	58	58	58	44	44	4	3 3	42 43	41 52	41 41
ime09	57	5	4 5	58 57	58 57	58 58	44 43	44 45	4	3 3 4	42 43 44	41 52 52	41 41 41
ime09 RB32	57 58	5 6 5	4 5 6	58 57 56	58 57 55	58 58 57	44 43 46	44 45 44	4	3 3 4 3	42 43 44 43	41 52 52 52	41 41 41 42
RB32	57 58 56	5 6 5 5	4 5 6 3	58 57 56 57	58 57 55 56	58 58 57 57	44 43 46 44	44 45 44 43	4: 4: 4: 4: 4:	3 3 4 3 3	42 43 44 43 43	41 52 52 52 52 51	41 41 41 42 41
RB32 vB_EcoM_ACG-C4	57 58 56 0 55	5 6 5 5 5 5	4 5 6 3 5 5	58 57 56 57 57 57	58 57 55 56 56	58 58 57 57 57 57	44 43 46 44 43	44 45 44 43 44	4: 4: 4: 4: 4: 4: 4:	3 3 4 3 3 3 3	42 43 44 43 43 43 43	41 52 52 52 51 41	41 41 42 41 40
RB32 vB_EcoM_ACG-C4 pSs-1	57 58 56 0 55 53	5 6 5 5 5 5 5	4 5 6 3 5 3	58 57 56 57 57 57 55	58 57 55 56 56 55	58 58 57 57 57 57 55	44 43 46 44 43 43	44 45 44 43 44 46	4: 4: 4: 4: 4: 4: 4: 4: 4: 4: 4: 4: 4: 4	3 3 4 3 3 3 3 4	42 43 44 43 43 43 43 43	41 52 52 52 51 41 53	41 41 42 41 40 41
RB32 vB_EcoM_ACG-C4 pSs-1 vB_EcoM_JS09	57 58 56 0 55 53 100	5 6 5 5 5 5 5 9	4 5 6 3 5 3 4	58 57 56 57 57 57 55 91	58 57 55 56 56 55 91	58 58 57 57 57 57 55 92	44 43 46 44 43 43 43 48	44 45 44 43 44 46 48	4: 4: 4: 4: 4: 4: 4: 4: 4: 4: 4: 4: 4: 4	3 3 3 4 3 3 3 3 3 4 7 7	42 43 44 43 43 43 43 43 43 46	41 52 52 52 51 41 53 55	41 41 42 41 40 41 40 41 43
RB32 vB_EcoM_ACG-C4 pSs-1	57 58 56 0 55 53 100 94	5 6 5 5 5 5 5 9 9	4 5 6 3 5 5 3 4 00 0	58 57 56 57 57 55 91 90	58 57 55 56 56 55 91 90	58 58 57 57 57 57 55 92 91	44 43 46 44 43 43 43 48 51	44 45 44 43 44 46 48 50	43 44 43 44 44 44 44 44 44 44	3     3       4     3       3     3       3     4       7     4	42 43 44 43 43 43 43 43 43 46 49	41 52 52 51 41 53 55 54	41 41 42 41 40 41
RB32 vB_EcoM_ACG-C4 pSs-1 vB_EcoM_JS09	57 58 56 0 55 53 100	5 6 5 5 5 5 5 9 9	4 5 6 3 5 3 4	58 57 56 57 57 57 55 91	58 57 55 56 56 55 91	58 58 57 57 57 57 55 92	44 43 46 44 43 43 43 48	44 45 44 43 44 46 48 50 50	4: 4: 4: 4: 4: 4: 4: 4: 4: 4: 4: 4: 4: 4	3     3       4     3       3     3       3     4       7     4	42 43 44 43 43 43 43 43 43 46	41 52 52 51 41 53 55 54 50	41 41 42 41 40 41 40 41 43
RB32 vB_EcoM_ACG-C4 pSs-1 vB_EcoM_JS09 HX01	57 58 56 0 55 53 100 94	5 6 5 5 5 5 9 9 10 8	4 5 6 3 5 5 3 4 00 0	58 57 56 57 57 55 91 90	58 57 55 56 56 55 91 90	58 58 57 57 57 57 55 92 91	44 43 46 44 43 43 43 48 51	44 45 44 43 44 46 48 50	43 44 43 44 44 44 44 44 44 44	3     3       3     3       3     3       3     4       7     4       9     9	42 43 44 43 43 43 43 43 43 46 49	41 52 52 51 41 53 55 54	41 41 42 41 40 41 43 39
RB32 vB_EcoM_ACG-C4 pSs-1 vB_EcoM_JS09 HX01 Shf125875	57 58 56 55 53 100 94 91 90	5 6 5 5 5 5 5 9 9 11 8 8 8	4 5 5 5 3 5 3 5 4 6 9 9	58 57 56 57 57 57 55 91 90 100	58 57 55 56 56 55 91 90 94	58 58 57 57 57 55 92 91 94	44 43 46 44 43 43 43 48 51 47	44 45 44 43 44 46 48 50 50	4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4	3     3       4     3       3     3       3     4       7     4       9     8	42 43 44 43 43 43 43 43 43 46 49 47	41 52 52 51 41 53 55 54 50 51	41 41 42 41 40 41 43 39 41
RB32           vB_EcoM_ACG-C4           pSs-1           vB_EcoM_JS09           HX01           Shf125875           RB69	57 58 56 0 55 53 100 94 91 90 22 92	5 6 5 5 5 5 5 5 9 9 10 8 8 8 8 9 9	4 5 5 5 3 5 4 9 9 9 0 0	58       57       56       57       57       57       91       90       100       93       94	58 57 55 56 56 55 91 90 94 100 95	58 58 57 57 57 55 92 91 94 94 95 100	44 43 46 44 43 43 43 43 48 51 47 46 47	44 45 44 43 44 46 48 50 50 50 49 50	4: 44: 43: 44: 44: 44: 44: 44: 44: 44: 4	3	42 43 44 43 43 43 43 43 43 43 46 49 47 47 47	41 52 52 51 41 53 55 54 50 51 51	41 41 42 41 40 41 43 39 41 40 41
RB32           vB_EcoM_ACG-C4           pSs-1           vB_EcoM_JS09           HX01           Shf125875           RB69           vB_EcoM_PhAPEC           IME08	57 58 56 0 55 53 100 94 91 90 22 92 48	5 6 5 5 5 5 5 5 5 5 6 9 9 10 8 8 8 8 8 9 5 5	4 5 6 3 5 5 3 4 00 9 9 9 9 0 1	58           57           56           57           57           57           55           91           90           100           93           94           47	58 57 55 56 56 55 91 90 90 94 100 95 47	58 58 57 57 57 55 92 91 94 94 95 100 47	44 43 46 44 43 43 43 43 48 51 47 46 47 100	44 45 44 43 44 46 48 50 50 50 49 50 78	4: 4: 4: 4: 4: 4: 4: 4: 4: 4: 4: 4: 4: 4	3	42 43 44 43 43 43 43 43 43 43 46 49 47 47 47 78	41 52 52 51 41 53 55 54 50 51 51 62	41 41 42 41 40 41 43 39 41 40 41 57
RB32           vB_EcoM_ACG-C4           pSs-1           vB_EcoM_JS09           HX01           Shf125875           RB69           vB_EcoM_PhAPEC           IME08           JS98	57 58 56 0 55 53 100 94 91 90 22 92 48 48	5 6 5 5 5 5 5 5 5 5 5 9 9 9 11 8 8 8 8 8 9 9 5 5 5 5 5 5 5 5 5 5 5 5 5	4 5 5 6 3 3 5 5 3 4 9 9 9 9 0 0 1 0	58           57           56           57           57           55           91           90           100           93           94           47           49	58 57 55 56 55 91 90 94 100 95 47 50	58           58           57           57           55           92           91           94           95           100           47           50	44 43 46 44 43 43 43 43 48 51 47 46 47 40 77	44 45 44 43 44 46 48 50 50 50 49 50 78 100	44 44 44 44 44 44 44 44 44 44 44 44 44	3	42 43 44 43 43 43 43 43 43 43 46 49 47 47 47 78 93	41 52 52 51 41 53 55 54 50 51 51 62 59	41 41 42 41 40 41 43 39 41 40 41 57 62
RB32           vB_EcoM_ACG-C4           pSs-1           vB_EcoM_JS09           HX01           Shf125875           RB69           vB_EcoM_PhAPEC           IME08           JS98           Bp7	57 58 56 0 55 53 100 94 91 90 22 92 48 48 48 46	55 55 55 99 11 88 88 99 55 55 44	4 5 5 6 3 5 5 3 4 9 9 9 0 9 1 0 3 3	58           57           56           57           57           55           91           90           100           93           94           47           49           48	58 57 55 56 55 91 90 94 100 95 47 50 48	58           58           57           57           55           92           91           94           95           100           47           50           48	44 43 46 44 43 43 43 43 48 51 47 46 47 46 47 100 77 82	44 45 44 43 44 46 48 50 50 50 49 50 78 100 83	44 44 44 44 44 44 44 44 44 44 44 44 44	3	42 43 44 43 43 43 43 43 43 43 43 46 49 47 47 47 78 93 83	41 52 52 51 41 53 55 54 50 51 51 62 59 58	41 41 42 41 40 41 43 39 41 40 41 57 62 59
RB32           vB_EcoM_ACG-C4           pSs-1           vB_EcoM_JS09           HX01           Shf125875           RB69           vB_EcoM_PhAPEC           IME08           JS98           Bp7           JS10	57 58 56 0 55 53 100 94 91 90 22 92 48 48 48 48 46 46	55 55 55 99 11 88 88 99 55 55 55 44	4 5 5 6 3 3 5 5 3 4 9 9 9 9 0 0 1 1 0 3 3 9	58           57           56           57           57           55           91           90           100           93           94           47           48           47	58 57 55 56 55 91 90 94 100 95 47 50 48 47	58           58           57           57           55           92           91           94           95           100           47           50           48           47	44 43 46 44 43 43 43 43 48 51 47 46 47 40 77 82 78	44 45 44 43 44 46 48 50 50 50 49 50 78 100 83 94	44 44 44 44 44 44 44 44 44 44 44 44 44	3	42 43 44 43 43 43 43 43 43 43 46 49 47 47 47 47 83 83 83 100	41 52 52 51 41 53 55 54 50 51 51 62 59 58 59	41 41 42 41 40 41 43 39 41 40 41 57 62 59 60
RB32           vB_EcoM_ACG-C4           pSs-1           vB_EcoM_JS09           HX01           Shf125875           RB69           vB_EcoM_PhAPEC           IME08           JS98           Bp7           JS10           SP18	57 58 56 0 55 33 100 94 91 90 22 92 48 48 48 48 46 46 47 55	55 55 55 99 11 88 88 99 55 55 55 55 55 55 55 55 55 55 55 55	4 5 5 6 3 3 5 5 3 4 9 9 9 9 0 9 1 1 0 1 3 3 9 9 5 5	58           57           56           57           57           55           91           90           100           93           94           47           49           48           47           51	58           57           55           56           55           91           90           94           100           95           47           50           48           47           52	58           58           57           57           55           92           91           94           95           100           47           50           48           47           52	44 43 46 44 43 43 43 43 48 51 47 46 47 46 47 100 77 82 78 62	44 45 44 43 44 46 48 50 50 50 49 50 78 100 83 94 60	4: 4: 4: 4: 4: 4: 4: 4: 4: 4: 4: 4: 4: 4	3	42 43 44 43 43 43 43 43 43 43 43 46 49 47 47 47 47 83 83 83 100 59	41 52 52 51 41 53 55 54 50 51 51 62 59 58 59 100	41 41 42 41 40 41 43 39 41 40 41 57 62 59 60 94
RB32           vB_EcoM_ACG-C4           pSs-1           vB_EcoM_JS09           HX01           Shf125875           RB69           vB_EcoM_PhAPEC           IME08           JS98           Bp7           JS10	57 58 56 0 55 53 100 94 91 90 22 92 48 48 48 48 46 46 47 55 55 42	5 6 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5	4 5 5 6 3 3 5 5 3 4 9 9 0 0 1 0 3 9 9 5 5 9 9	58           57           56           57           57           55           91           90           100           93           94           47           49           48           47           51           41	58           57           55           56           55           91           90           94           100           95           47           50           48           47           52           41	58           58           57           57           55           92           91           94           95           100           47           50           48           47           52           41	44 43 46 44 43 43 43 48 51 47 46 47 46 47 100 77 82 78 62 55	44 45 44 43 44 46 48 50 50 50 49 50 78 100 83 94 60 62	4: 4: 4: 4: 4: 4: 4: 4: 4: 4: 4: 4: 4: 4	3	42 43 44 43 43 43 43 43 43 43 46 49 47 47 47 47 47 83 83 83 100 59 59	41 52 52 51 41 53 55 54 50 51 51 62 59 58 59 58 59 100 84	41 41 42 41 40 41 43 39 41 40 41 57 62 59 60 94 100
RB32           vB_EcoM_ACG-C4           pSs-1           vB_EcoM_JS09           HX01           Shf125875           RB69           vB_EcoM_PhAPEC           IME08           JS98           Bp7           JS10           SP18	57 58 56 0 55 53 100 94 91 90 22 92 48 48 48 46 46 47 55 55 42 36	5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5	4       5       6       3       5       3       4       00       9       0       1       0       3       9       5       9       5       9       6	58         57           56         57           57         55           91         90           100         93           94         47           49         48           47         51           51         36	58           57           55           56           55           91           90           94           100           95           47           50           48           47           52	58           58           57           57           55           92           91           94           95           100           47           50           48           47           52	44 43 46 44 43 43 43 48 51 47 47 46 47 100 77 82 78 62 55 33	44 45 44 43 44 46 48 50 50 50 49 50 78 100 83 94 60	4: 4: 4: 4: 4: 4: 4: 4: 4: 4: 4: 4: 4: 4	3	42 43 44 43 43 43 43 43 43 43 43 46 49 47 47 47 47 83 83 83 100 59	41 52 52 51 41 53 55 54 50 51 51 62 59 58 59 100	41 41 42 41 40 41 43 39 41 40 41 57 62 59 60 94
RB32           vB_EcoM_ACG-C4           pSs-1           vB_EcoM_JS09           HX01           Shf125875           RB69           vB_EcoM_PhAPEC           IME08           JS98           Bp7           JS10           SP18           vB_EcoM-VR7	57 58 56 0 55 53 100 94 91 90 22 92 48 48 48 48 46 46 47 55 55 42	5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5	4 5 5 6 3 3 5 5 3 4 9 9 0 0 1 0 3 9 9 5 5 9 9	58           57           56           57           57           55           91           90           100           93           94           47           49           48           47           51           41	58           57           55           56           55           91           90           94           100           95           47           50           48           47           52           41	58           58           57           57           55           92           91           94           95           100           47           50           48           47           52           41	44 43 46 44 43 43 43 48 51 47 46 47 46 47 100 77 82 78 62 55	44 45 44 43 44 46 48 50 50 50 49 50 78 100 83 94 60 62	4: 4: 4: 4: 4: 4: 4: 4: 4: 4: 4: 4: 4: 4	3	42 43 44 43 43 43 43 43 43 43 46 49 47 47 47 47 47 83 83 83 100 59 59	41 52 52 51 41 53 55 54 50 51 51 62 59 58 59 58 59 100 84	41 41 42 41 40 41 43 39 41 40 41 57 62 59 60 94 100
RB32           vB_EcoM_ACG-C4           pSs-1           vB_EcoM_JS09           HX01           Shf125875           RB69           vB_EcoM_PhAPEC           IME08           JS98           Bp7           JS10           SP18           vB_EcoM-VR7           S16	57 58 56 0 55 53 100 94 91 90 22 92 48 48 48 48 46 47 55 55 42 36 36	5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5	4       5       6       3       5       3       4       00       9       9       0       1       0       3       9       5       9       6       6	58           57           56           57           57           57           55           91           90           100           93           94           47           51           41           36           35	58           57           55           56           55           91           90           94           100           95           47           50           48           47           52           41           36           36	58           58           57           57           57           55           92           91           94           95           100           47           50           48           47           52           41           36           36	44 43 46 44 43 43 43 43 48 51 47 46 47 100 77 82 78 62 55 33 32	44 45 44 43 43 44 46 48 50 50 49 50 78 100 83 94 40 60 62 31 33	4: 4: 4: 4: 4: 4: 4: 4: 4: 4: 4: 4: 4: 4	3	42 43 44 43 43 43 43 43 43 43 46 49 47 47 47 47 83 93 83 100 59 59 33	41 52 52 51 41 53 55 54 50 51 51 62 59 58 59 100 84 31	41 41 42 41 40 41 43 39 41 40 41 57 62 59 60 94 100 32
RB32           vB_EcoM_ACG-C4           pSs-1           vB_EcoM_JS09           HX01           Shf125875           RB69           vB_EcoM_PhAPEC           IME08           JS98           Bp7           JS10           SP18           vB_EcoM-VR7           S16           STML-198           PG7	57 58 56 0 55 53 100 94 91 90 22 92 48 48 48 48 46 47 55 55 42 36 36 37	5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5	4       5       6       3       5       3       4       00       9       9       0       1       0       3       9       5       9       6       6       7	58           57           56           57           56           57           55           91           90           100           93           94           47           49           48           47           51           41           36           35           37	58           57           55           56           55           91           90           94           100           95           47           50           48           47           52           41           36           37	58           58           57           57           57           55           92           91           94           95           100           47           50           48           47           52           41           36           37	44 43 46 44 43 43 43 48 51 47 46 47 100 77 82 78 62 55 33 32 35	44 45 44 43 43 44 46 48 50 50 49 50 78 100 83 94 40 60 62 31 33 35	44 44 44 44 44 44 44 44 44 44 44 44 44	3	42 43 44 43 43 43 43 43 43 43 47 47 47 47 47 47 83 83 100 59 59 33 33 33 35	41 52 52 51 41 53 55 54 50 51 51 62 59 58 59 59 100 84 31 31 31	41 41 41 42 41 40 41 43 39 41 40 41 57 62 59 60 94 100 32 33 32
RB32           vB_EcoM_ACG-C4           pSs-1           vB_EcoM_JS09           HX01           Shf125875           RB69           vB_EcoM_PhAPEC           IME08           JS98           Bp7           JS10           SP18           vB_EcoM-VR7           S16           STML-198           PG7           CC31	57 58 56 0 55 53 100 94 91 90 22 92 48 48 48 48 46 46 47 55 42 36 36 37 37	5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5	4       5       6       3       5       3       4       00       9       9       0       1       0       3       9       5       9       6       6       7	58           57           56           57           56           57           55           91           90           100           93           94           47           49           48           47           51           41           36           35           37	58           57           55           56           55           91           90           94           100           95           47           50           48           47           52           41           36           37           37	58           58           57           57           57           55           92           91           94           95           100           47           50           48           47           52           41           36           37           37	44 43 46 44 43 43 43 43 48 51 47 46 47 100 77 82 78 62 55 33 32 35 34	44 45 44 43 43 44 46 48 50 50 50 49 50 78 100 83 94 60 62 31 33 35 35	44 44 44 44 44 44 44 44 44 44 44 44 44	3	42 43 44 43 43 43 43 43 43 43 46 49 47 47 47 47 47 83 83 100 59 59 33 33 33 35 34	41 52 52 51 41 53 55 54 50 51 51 62 59 58 59 59 100 84 31 31 31 32	41 41 41 42 41 40 41 43 39 41 40 41 57 62 59 60 94 100 32 33 32 31
RB32           vB_EcoM_ACG-C4           pSs-1           vB_EcoM_JS09           HX01           Shf125875           RB69           vB_EcoM_PhAPEC           IME08           JS98           Bp7           JS10           SP18           vB_EcoM-VR7           S16           STML-198           PG7           CC31           JSE	57 58 56 0 55 53 100 94 91 90 22 92 48 48 48 48 46 46 47 55 55 42 36 36 37 37	5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5	4       5       6       3       5       3       4       00       9       9       0       1       0       3       9       5       9       6       6       7       2	58           57           56           57           56           57           55           91           90           100           93           94           47           51           41           36           35           37           37           12	58           57           55           56           55           91           90           94           100           95           47           50           48           47           52           41           36           37           37           13	58           58           57           57           57           55           92           91           94           95           100           47           50           48           47           52           41           36           37           37           13	44 43 46 44 43 43 43 43 48 51 47 46 47 100 77 82 78 62 55 55 33 32 35 34 11	44 45 44 43 43 44 46 48 50 50 50 49 50 78 100 83 94 60 60 62 31 333 35 35	44 44 44 44 44 44 44 44 44 44 44 44 44	3	42 43 44 43 43 43 43 43 43 46 49 47 47 47 47 47 83 93 83 100 59 59 33 33 35 34 11	41 52 52 51 41 53 55 54 50 51 51 62 59 58 59 100 84 31 31 31 32 12	41 41 41 42 41 40 41 43 39 41 40 41 57 62 59 60 94 100 32 33 32 31 11
RB32           vB_EcoM_ACG-C4           pSs-1           vB_EcoM_JS09           HX01           Shf125875           RB69           vB_EcoM_PhAPEC           IME08           JS98           Bp7           JS10           SP18           vB_EcoM-VR7           S16           STML-198           PG7           CC31	57 58 56 0 55 53 100 94 91 90 22 92 48 48 48 48 46 46 47 55 42 36 36 37 37	5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5	4       5       6       3       5       3       4       00       9       9       0       1       0       3       9       5       9       6       6       7	58           57           56           57           56           57           55           91           90           100           93           94           47           49           48           47           51           41           36           35           37	58           57           55           56           55           91           90           94           100           95           47           50           48           47           52           41           36           37           37	58           58           57           57           57           55           92           91           94           95           100           47           50           48           47           52           41           36           37           37	44 43 46 44 43 43 43 43 48 51 47 46 47 100 77 82 78 62 55 33 32 35 34	44 45 44 43 43 44 46 48 50 50 50 49 50 78 100 83 94 60 62 31 33 35 35	44 44 44 44 44 44 44 44 44 44 44 44 44	3	42 43 44 43 43 43 43 43 43 43 46 49 47 47 47 47 47 83 83 100 59 59 33 33 33 35 34	41 52 52 51 41 53 55 54 50 51 51 62 59 58 59 59 100 84 31 31 31 32	41 41 41 42 41 40 41 43 39 41 40 41 57 62 59 60 94 100 32 33 32 31

	SP18	VR7	S16	STML-198	PG7	CC31	JSE	RB49	Phi1
T4	42	42	42	42	37	38	15	14	13
PST	41	41	42	42	38	38	14	14	14
Shfl2	52	41	41	41	39	39	15	14	14
e11/2	42	41	41	41	37	39	13	14	14
phiD1	53	41	41	41	39	39	13	14	15
ECML-134	51	41	41	41	38	38	14	14	14
RB3	41	41	42	41	39	40	15	14	14
RB14	52	41	42	44	38	39	13	14	14
AR1	52	41	41	41	38	39	13	13	15
ime09	52	42	42	44	39	40	13	14	14
RB32	51	41	41	41	38	39	14	14	14
vB_EcoM_ACG-C40	41	40	42	44	38	39	13	14	14
pSs-1	53	41	42	44	39	41	13	13	14
vB_EcoM_JS09	55	43	39	39	36	37	13	13	13
HX01	54	39	39	39	36	37	12	14	13
Shf125875	50	41	38	38	37	37	13	14	13
RB69	51	40	38	38	33	37	13	14	13
vB_EcoM_PhAPEC2	51	41	39	38	35	37	13	13	13
IME08	62	57	35	35	34	35	12	12	12
JS98	59	62	34	36	33	35	11	12	11
Bp7	58	59	35	35	33	34	12	11	11
JS10	59	60	35	36	33	35	11	12	11
SP18	100	94	33	34	31	33	12	12	12
vB_EcoM-VR7	84	100	33	36	31	31	11	12	11
S16	31	32	100	92	44	45	12	13	13
STML-198	31	33	91	100	43	45	12	13	13
PG7	31	32	48	47	100	87	13	14	14
CC31	32	31	47	48	84	100	13	14	14
JSE	12	11	12	13	13	13	100	94	91
RB49	12	11	13	13	14	14	92	100	94
Phi1	12	11	13	14	13	14	90	94	100

 Table 2.
 T4-related phage genera and species

Genus	Species	Host	Accession	Related strains
			No.	
T4virus	T4	Escherichia	AF158101.6	T4T;         T4         strain           wild;         T4         strain           GT7;         T4         strain           147;         RB55(KM607002)         ;           ;         RB59
				(KM607003)
	AR1	Escherichia	AP011113	RB51 (FJ839693); RB68 (KM607004); wV7 (HM997020); RB27 (KM607000)
	RB32	Escherichia	DQ904452	RB33(KM607001)
	RB14	Escherichia	FJ839692	
	ime09	Escherichia	JN202312	
	e11/2 (vB_EcoM_ 112)	Escherichia	KJ668714.2	
	ECML-134	Escherichia	JX128259	
	RB3	Escherichia	KM606994	RB5 (KM606995);

				RB6 (KM606996); RB7 (KM606997); RB9 (KM606998); RB10 (KM606999)
	vB_EcoM_ACG- C40	Escherichia	JN986846	(KW000999)
	Shfl2	Shigella	HM035025	
	pSs-1	Shigella	KM501444	
	PST	Yersinia	KF208315	
	phiD1	Yersinia	HE956711	
Rb69virus	RB69	Escherichia	AY303349	vB_EcoM_PhAPE C2 (KF562341)
	vB_EcoM_JS09	Escherichia	KF582788	O157 typing phages 3 (KP869101) & 6 (KP869104)
	HX01	Escherichia	JX536493	
	Shf125875	Shigella	KM407600	
Rb49virus	RB49	Escherichia	AY343333	GEC-3S (HE978309)
	Phi1	Escherichia	EF437941	
	JSE	Escherichia	EU863408	
Js98virus	JS98	Escherichia	EF469154	
	IME08	Escherichia	HM071924	
	Bp7	Escherichia	HQ829472	
	JS10	Escherichia	EU863409	
	vB_EcoM_VR5	Escherichia	KP007359	
S16virus	STML-198	Salmonella	JX181825	
	S16	Salmonella	HQ331142	STP4-a (KJ000058)
Sp18virus	SP18	Shigella	GQ981382	
*	vB_EcoM-VR7	Escherichia	HM563683	
	vB_EcoM-VR20	Escherichia	KP007360	
	vB EcoM-VR25	Escherichia	KP007361	
	vB_EcoM-VR26	Escherichia	KP007362	
Cc31virus	CC31	Enterobacter	GU323318	
	PG7	Enterobacter	KJ101592	

 Table 3. Average properties of each genus

Phage genus	Size (kb)	GC%	No. of Proteins	No. of tRNAs
T4virus	166.9	35.4	274	9

Rb69virus	168.7	37.6	271	2
Rb49virus	164.9	40.5	277	0
Js98virus	170.6	39.5	262	2
S16virus	159.2	36.9	262	3
Sp18virus	170.0	40.4	289	2
Cc31virus	169.4	39.9	287	14