

Bacteriophages and its applications: an overview

Sonika Sharma¹ · Soumya Chatterjee¹ · Sibnarayan Datta¹ · Rishika Prasad^{1,2} ·
Dharmendra Dubey¹ · Rajesh Kumar Prasad¹ · Mohan G Vairale¹

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Abstract Bacteriophages (or phages), the most abundant viral entity of the planet, are omni-present in all the ecosystems. On the basis of their unique characteristics and anti-bacterial property, phages are being freshly evaluated taxonomically. Phages replicate inside the host either by lytic or lysogenic mode after infecting and using the cellular machinery of a bacterium. Since their discovery by Twort and d’Herelle in the early 1900s, phage became an important agent for combating pathogenic bacteria in clinical treatments and its related research gained momentum. However, due to recent emergence of bacterial resistance on antibiotics, applications of phage (phage therapy) become an inevitable option of research. Phage particles become popular as a biotechnological tool and treatment of pathogenic bacteria in a range of applied areas. However, there are few concerns over the application of phage-based solutions. This review deals with the updated phage taxonomy (*ICTV 2015 Release and subsequent revision*) and phage biology and the recent development of its application in the areas of biotechnology, biosensor, therapeutic medicine, food preservation, aquaculture diseases, pollution remediation, and wastewater treatment and issues related with limitations of phage-based remedy.

Introduction

There are billions of viruses on the Earth. An estimation of their number reaches an astronomical figure of about 10^{31} viruses, which is much more than the number of stars in the universe (Weitz and Wilhelm 2013). Most of the viruses play a significant role in the global biogeochemical cycles and thrive by infecting microbes like bacteria, archaea, and microeukaryotes (Suttle 2007). Because of their astounding numbers, and intimate relationship with different microbes, they control both host populations and ecosystem functions (Weitz and Wilhelm 2012). Among the total viral population, bacteriophages, popularly known as phages (Greek “phagein” meaning “to eat”), are the group of viruses that infect and devour bacteria. This potential antibacterial property of phage is unique (Jamalludeen et al. 2009).

Phages are considered to be the most diverse and abundant entity on earth and are thought to exist in every ecosystem (Maranger and Bird 1995; Hendrix 2002; Hanlon 2007; Le Romancer et al. 2007) ranging from extremely hot environments like hot springs, the Sahara, to extremely cold environments like polar inland waters (Breitbart et al. 2004; Glud and Middleboe 2004; Prigent et al. 2005; Suttle 2005; Sävström et al. 2008; Lin et al. 2010). Sea water is among the hugely diverse and most dense natural environments for phages and other viruses; for example, the surface seawater has a concentration of approximately 10 million viruses per milliliter (Breitbart 2012). Similarly, forest floors and agricultural soils usually harbor a phage count of approximately 10^8 – 10^9 per gram of soil (dry mass; Williamson et al. 2003; Williamson et al. 2005). It is also reported that to carry out important ecological functions, phages express a variety of auxiliary metabolic genes (Breitbart 2012; Weitz and Wilhelm 2013).

A single phage particle can hunt for a specific bacterium species or a subset of the same species. After infecting a

Sonika Sharma and Soumya Chatterjee contributed equally.

✉ Soumya Chatterjee
drlsoumya@gmail.com

¹ Defence Research Laboratory, DRDO, Tezpur, Assam 784001, India

² School of Biomedical Engineering, Cornell University, Ithaca, NY 14850, USA

bacterium, phages replicate inside it. Once a bacterial cell is infected, phages multiply exponentially using the cellular machinery of the bacteria including protein-synthesizing and energy-generating systems. However, the method of propagation may either be lytic or lysogenic in nature. The lytic phages cause lysis of the host bacterial cells to release progeny viruses (Young et al. 2000). On the other hand, lysogenic or the temperate phages integrate their nucleic acid (genome) within the host bacterial cell and replicate along with the host, conferring new properties to the host bacteria (Brock et al. 1988). Phages are classified on the basis of certain criteria including its host specificity, morphology, nucleic acid type, mode of infection, morphogenesis, phylogeny, serology, sensitivity to physical and chemical agents, and its environment (Abeles et al. 1984). All bacterial genera, including the cyanobacteria, archaeobacterial, and mycoplasmas, are known to be vulnerable to a plethora of phages.

Brief history of bacteriophage

British bacteriologist Ernest Hanbury Hankin, in 1896, first reported antibacterial activity in the waters of the river Ganga and the river Yamuna in India (Hankin 1896: <http://icmr.nic.in/buapril02.pdf>). The then unidentified antibacterial agent was also noted as filterable and heat labile; it was found to limit the spread of cholera epidemic (Hankin 1896; Van Helvoort 1992; Sulakvelidze et al. 2001). However, how the implications of Hankin's findings related to the existence of bacteriophages remain dubious in the scientific society.

In 1915, Frederick William Twort FRS (October 22, 1877–March 20, 1950), an English bacteriologist, superintendent of the Brown Institute for Animals and professor of bacteriology at the University of London, while growing viruses in a laboratory condition, found zones of clearance in the form of “transmissible glassy transformation” with micrococcus bacteria. He also concluded that this agent multiplied itself in the process of killing the bacteria (Twort 1915, 1922, 1949).

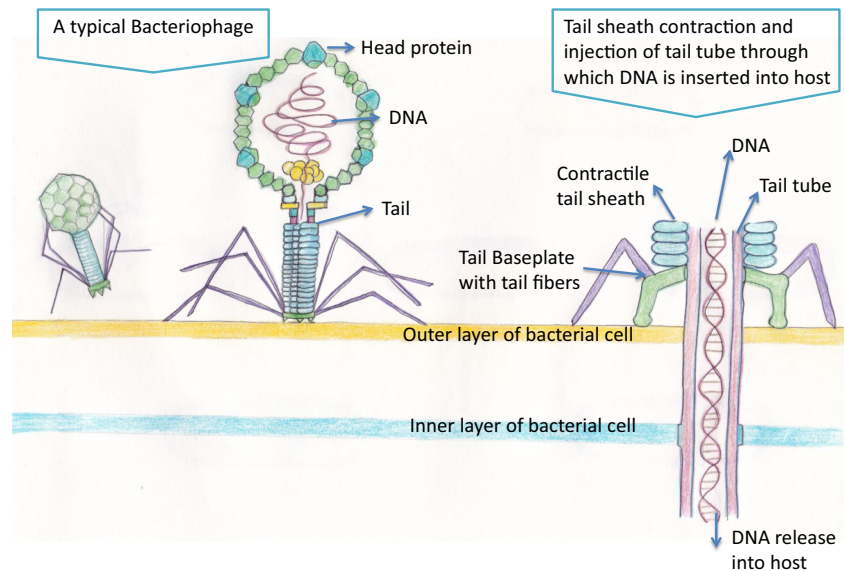
On the other hand, Félix d’Herelle (April 25, 1873–February 22, 1949), a French-Canadian microbiologist working at the Pasteur Institute in Paris, observed the bacteriophage phenomenon (in the year 1917) on severe hemorrhagic dysentery among French troops stationed at Maisons-Laffitte. He had observed the same phenomenon in 1910 when he was studying microbiology due to an epizootic locust infection in Mexico. d’Herelle prepared bacteria-free filtrates from fecal samples of patients and incubated the filtrate with the bacterium *Shigella*, isolated from those patients. His primary goal was to develop a vaccine against bacterial dysentery. For observing the growth of the bacteria, an aliquot of the filtrate-bacteria mixture was spread on agar medium and incubated. d’Herelle found the appearance of small, clear zones, which he primarily termed *taches*, then *taches vierges*, and later, *plaques*. This finding

was presented in September 1917 and subsequently published (d’Herelle 1917, 1930; Summers 1999; Sulakvelidze et al. 2001). d’Herelle also proposed that the phenomenon was caused by a bacteria parasitizing virus and named “bacteriophage”—*phages* implying to “eat” or “devour” bacteria. As per the recollection of d’Herelle, the name “bacteriophage” was decided together with his wife Marie, and the word came into existence on 18 October 1916—the day before their youngest daughter’s birthday (d’Herelle 1930; Summers 1999; Sulakvelidze et al. 2001). Thus, the credit of discovery of bacteriophages goes independently to two scientists: F.W. Twort and Félix d’Herelle. Primarily, the scientific society first called it “Twort-d’Herelle phenomenon,” and later, the “bacteriophage phenomenon” (d’Herelle 1949; Duckworth 1976; Sulakvelidze et al. 2001).

Structure of bacteriophage

Length of phages varies widely and usually ranges from 24 to 200 nm (Mayer 2016). T4 phages are among the largest phages that are approximately 200 nm in length and 80–100 nm wide. Phages having an icosahedral shape or a shape with 20 sides and filaments are also common (<http://www.microrao.com/micro notes/bacteriophage.pdf>). After the first commercial production of transmission electron microscope (called “Hypermicroscope”) by Siemens and Halske Company (Germany) in 1939, photomicrography of bacteriophages came into the existence. Ruska (1940) and Pfankuch and Kausche (1940), both working in the same company, reported separately in the same journal and on same date showing the first pictures of bacteriophages to the scientific world (Ruska 1940; Pfankuch and Kausche 1940; Ackermann, 2011a, b). On the other hand, in North America, Prebus and Hillier of University of Toronto constructed separately the first electron microscope in 1938, and subsequently, commercialized the same after years of research by Hiller at Radio Corporation of America (RCA, Princeton, NJ; Prebus and Hillier 1939; Haguénau et al. 2003; Ackermann 2011a, b). Luria and Anderson, in 1942, reported different unstained coli and staphylococcal phage particles, which were renamed later as T2 (T-even type; Luria and Anderson 1942; Ackermann 2011a, b). Despite continental difference, huge political unrest (because of World War II) and extreme limitations on scientific exchange, interestingly, both the group of scientists were aware of other progress in the development of this extremely important scientific instrument (Pfankuch and Kausche 1940; Ackermann 2011a, b). Later on, using the “negative contrast” technique in an electron microscope, Brenner and his team first reported a clearer image of bacteriophages in 1959 (Brenner and Horne 1959; Brenner et al. 1959). In subsequent years, Bradley and Kay (1960) and Bradley (1967) elaborated fine structures of phages. The basic features of a typical bacteriophage, in this case the T4 bacteriophage, include a “head” or capsid and a “tail” (Fig. 1).

Fig. 1 Schematic representation of a typical structure of a phage emphasizing tail region and contractile ability to insert its DNA into host bacterial cell by penetrating cell wall



The head or capsid structure, regardless of its size or shape, is a congregation of one or more protein subunits called protomers. The morphological subunit of a capsid is a capsomere which protects the viral nucleic acid (genome). The tail is a hollow tube through which the nucleic acid passes when the bacteriophage infects a bacterial host cell. Some phages do not possess a tail. The T4 phage has additional structures, namely the base plate and tail fibers attached to the tail that aid the phage in attaching itself to a bacterial cell (Leiman et al. 2003).

Taxonomy of phages

As mentioned earlier, there are billions and billions of phages. Deciphering taxonomic characterization is a challenging task, especially for such nano-sized phage particles. In 1967, phages were first classified by Bradley and were subsequently approved by the International Committee on Taxonomy of Viruses (ICTV) and total 111 phages were listed in classification (Wildy 1971). Bradley's classification projected six basic morphological types of tailed phages that further, categorized on the basis of morphotypes (contractile tails, long and noncontractile tails, and short tails), small isometric ssDNA viruses, filamentous phages, and small ssRNA phages. The regulating body for the viral taxonomic system (ICTV) characterizes phage particles taking into consideration numerous parameters like host range, physical characteristics (such as structure, capsid size, and shape), type of genomic material (single or double-stranded DNA or RNA), genome size, and resistance to organic solvents (Murphy et al. 1995). More than 96 % of phages are tailed and carries dsDNA as genetic material; however, they may vary in shape like cubic, filamentous, and pleomorphic (Ackermann 2011a, b). Polyphasic taxonomy was

revised and emphasis was given on genomic relationship (Thompson et al. 2015). Earlier in 2008, only 18 genera and 36 species were listed among three caudoviral families, Myoviridae, Podoviridae, and Siphoviridae. Subsequently, the order caudovirales expanded and an updated and detailed bacteriophage classification was presented in ICTV Release 2015 (<http://www.ictvonline.org/virusTaxonomy.asp>; Krupovic et al. 2016). Some alterations have also been suggested by committee, like replacement of word “phage” with “virus” in prokaryotic virus taxon names, omission of infix “like” from prokaryotic virus genus names, discouragement of use of “Phi” and other Greek letters in prokaryotic virus genera, exclusion of hyphens, and encouragement to use of host genus name in replacement of taxon names is being encouraged to avoid confusion related to phage action on specific bacterial host (Krupovic et al. 2016; Pietilä et al. 2016). The overview of the various bacteriophage families has been included in Table 1 compiled from (ICTV 2015 Release; EC 47, London, UK, July 2015 Email ratification 2016 (MSL #30) for ready reference to the readers.

Mechanism of proliferation

Specific receptors (like lipopolysaccharides, teichoic acids, proteins, and flagella) on the surface of the host bacteria are required for the phage to infect the bacteria. Due to this specificity of the receptor present on the bacterial cell surface, phages can infect specific hosts only. However, in solution, this interaction with the host is a random phenomenon for the phages. Bacterial type (Gram-negative and Gram-positive), growth conditions, and virulence also influence the phage to attach on the host's surface (Rakhuba et al. 2010). The outer membrane of Gram-negative bacteria has an external lipopolysaccharide (LPS) layer and embedded outer membrane

Table 1 The overview of the various phage families compiled from International Committee on Taxonomy of Viruses (ICTV 2015 Release; <http://www.ictvonline.org/virusTaxonomy.asp>)

Order	Family	Subfamily	Genus	Host bacteria	Species	Accession number (example species)	Isolate (example species)
<i>Caudovirales</i> (linear, dsDNA)	<i>Myoviridae</i> (icosahedral head with contractile tail)	<i>Eucampyvirinae</i>	<i>Cp220virus</i>	<i>Campylobacter</i>	<i>Campylobacter virus CP21</i>	HE815464	
					<i>Campylobacter virus CP220</i>	FN667788	
					<i>Campylobacter virus CP410</i>	FN667789	
					<i>Campylobacter virus IBB35</i>	HM246720	
					<i>Campylobacter virus CP81</i>	FR823450	
		<i>Peduvirinae</i>	<i>Cp8virus</i>	<i>Pasteurella</i> <i>Vibrio</i>	<i>Campylobacter virus CPX</i>	JN132397	
					<i>Campylobacter virus NCTC12673</i>	GU296433	
					<i>Aeromonas virus phiO18P</i>		
					<i>Haemophilus virus HP1</i>		
					<i>Haemophilus virus HP2</i>		
					<i>Vibrio virus K139</i>		
	<i>Podoviridae</i> (icosahedral head without contractile tail)	<i>P2virus</i>	<i>P2virus</i>	<i>Burkholderia</i>	<i>Vibrio virus Kappa</i>		
					<i>Burkholderia virus phi52237</i>		
					<i>Burkholderia virus phiE122</i>		
					<i>Burkholderia virus phiE202</i>		
					<i>Escherichia virus 186</i>		
					<i>Escherichia virus P2</i>		
					<i>Escherichia virus Wphi</i>		
					<i>Mannheimia virus MhaA1-PHL101</i>		
					<i>Pseudomonas virus phiCTX</i>		
					<i>Ralstonia virus RSA1</i>		
					<i>Salmonella virus Fels2</i>		
					<i>Salmonella virus PsP3</i>		
					<i>Salmonella virus SopEphi</i>		
	<i>Spounavirinae</i>	<i>Kayvirus</i>	<i>Kayvirus</i>	<i>Yersinia</i> <i>Staphylococcus</i>	<i>Yersinia virus L413C</i>		
					<i>Staphylococcus virus G1</i>		
					<i>Staphylococcus virus G15</i>	JQ686190	Staphylococcus phage G15
					<i>Staphylococcus virus JD7</i>	JX878671	Staphylococcus phage JD007
					<i>Staphylococcus virus K</i>		
					<i>Staphylococcus virus MCE2014</i>	KJ888149	Staphylococcus phage MCE-2014
					<i>Staphylococcus virus P108</i>		
					<i>Staphylococcus virus S253</i>	KM216423	Staphylococcus phage P108
					<i>Staphylococcus virus SA12</i>	AB853330	Staphylococcus phage S25-3
					<i>Staphylococcus virus SA12</i>	AB903967	Staphylococcus phage phiSA12
					<i>Listeria virus A511</i>		
					<i>Listeria virus P100</i>	DQ004855	Listeria phage P100
					<i>Staphylococcus virus Remus</i>	JX846613	Staphylococcus phage vB_SauM_Remus
	<i>SpoIVirus</i>	<i>SpoIVirus</i>	<i>SpoIVirus</i>	<i>Bacillus</i>	<i>Staphylococcus virus SA11</i>	JX194239	Staphylococcus phage SA11
					<i>Bacillus virus SPO1</i>		

Table 1 (continued)

Order	Family	Subfamily	Genus	Host bacteria	Species	Accession number (example species)	Isolate (example species)
		<i>Tevenvirinae</i>	<i>Twortvirus</i>	<i>Staphylococcus</i>	<i>Staphylococcus virus Twort</i>		
			<i>Unassigned</i>	<i>Enterococcus</i>	<i>Enterococcus virus phiEC24C</i>		
				<i>Lactobacillus</i>	<i>Lactobacillus phage Lb338-1</i>	FJ822135	
					<i>Lactobacillus virus LP65</i>		Enterobacter phage CC31
			<i>Cc31virus</i>	<i>Enterobacter</i>	<i>Enterobacter virus CC31</i>	GU323318	Enterobacter phage PG7
					<i>Enterobacter virus PG7</i>	KJ101592	Escherichia phage Bp7
			<i>Js98virus</i>	<i>Escherichia</i>	<i>Escherichia virus Bp7</i>	HQ829472	Escherichia phage IME08
					<i>Escherichia virus IME08</i>	HM071924	Escherichia phage JS10
					<i>Escherichia virus JS10</i>	EU863409	
					<i>Escherichia virus JS98</i>		Escherichia phage vb_EcoM-VR5
					<i>Escherichia virus VR5</i>	KP007359	
			<i>Rb49virus</i>		<i>Escherichia virus phi1</i>		
					<i>Escherichia virus RB49</i>		Escherichia phage HX01
			<i>Rb69virus</i>		<i>Escherichia virus HX01</i>	JX536493	Escherichia phage vB_EcoM_JS09
					<i>Escherichia virus JS09</i>	KF582788	
					<i>Escherichia virus RB69</i>		Shigella phage Shf125875
			<i>SI6virus</i>	<i>Shigella</i>	<i>Shigella virus UTAM</i>	KM407600	Salmonella phage vB_SenM-S16
				<i>Salmonella</i>	<i>Salmonella virus S16</i>	HQ331142	Salmonella phage STML-198
			<i>Schizot4virus</i>		<i>Salmonella virus STML198</i>	JX181825	
				<i>Vibrio</i>	<i>Vibrio virus KVP40</i>		Vibrio phage ValKK3
					<i>Vibrio virus ntl</i>		Escherichia phage vB_EcoM_VR7
			<i>Sp18virus</i>	<i>Escherichia</i>	<i>Vibrio virus ValKK3</i>	KP671755	Escherichia phage vB_EcoM_VR20
					<i>Escherichia virus VR7</i>	HM563683	Escherichia phage vB_EcoM_VR25
					<i>Escherichia virus VR20</i>	KP007360	Escherichia phage vB_EcoM_VR25
					<i>Escherichia virus VR25</i>	KP007361	Escherichia phage vB_EcoM_VR26
					<i>Escherichia virus VR26</i>	KP007362	Shigella phage SP18
			<i>T4virus</i>	<i>Shigella</i>	<i>Shigella virus SP18</i>	GQ981382	Escherichia phage AR1
				<i>Escherichia</i>	<i>Escherichia virus AR1</i>	AP011113	Escherichia phage vB_EcoM_ACG-C40
					<i>Escherichia virus C40</i>	JN986846	Escherichia phage vB_EcoM_112
					<i>Escherichia virus E112</i>	KJ668714.2	Escherichia phage ECML-134
					<i>Escherichia virus ECML134</i>	JX128259	Escherichia phage ime09
					<i>Escherichia virus Ime09</i>	JN202312	Escherichia phage RB3
					<i>Escherichia virus RB3</i>	KM606994	Escherichia phage RB3
					<i>Escherichia virus RB14</i>		

Order	Family	Subfamily	Genus	Host bacteria	Species	Accession number (example species)	Isolate (example species)				
	Unassigned			Shigella	Escherichia virus T4						
					Shigella virus Pss1	KM501444	Shigella phage pSs-1				
					Shigella virus Shf12	HM035025	Shigella phage Shf12				
					Yersinia virus D1	HE956711	Yersinia phage phiD1				
				Acinetobacter	Yersinia virus PST	KF208315	Yersinia phage PST				
					Acinetobacter virus I33						
				Aeromonas	Aeromonas virus 65						
					Aeromonas virus Aehl						
				Enterobacteria	Enterobacteria virus SV14						
					Escherichia virus RB16						
				Escherichia	Escherichia virus RB32						
					Escherichia virus RB43						
					Escherichia virus RB43						
					Pseudomonas virus 42						
				Cronobacter	Cronobacter virus CR3	JQ691612	Cronobacter phage CR3				
					Cronobacter virus CR8	KC954774	Cronobacter phage CR8				
					Cronobacter virus CR9	JQ691611	Cronobacter phage CR9				
					Cronobacter virus GAP31	JN882284	Cronobacter phage vB_CsaM_GAP31				
				Escherichia	Escherichia virus 4MG	KF550303	Escherichia phage 4MG				
					Salmonella virus SE1	GU070616	Salmonella phage SE1				
				Escherichia	Salmonella virus SSE121	JX181824	Salmonella phage SSE121				
					Escherichia virus FFH2	KJ190158	Escherichia phage FFH2				
					Escherichia virus FV3	JQ031132	Escherichia phage FV3				
					Escherichia virus JES2013	KC690136	Escherichia phage JES2013				
				Agatevirus	Escherichia virus V5	DQ832317	Escherichia phage V5				
					Bacillus virus Agate	JX238501	Bacillus phage phiAGATE				
				Ap22virus	Bacillus virus Bobb	KM051843	Bacillus phage Bobb				
					Bacillus virus Bp8pC	KJ010547	Bacillus phage Bp8p-C				
								Acinetobacter	Acinetobacter virus AB1	HM368260	Acinetobacter phage AB1
									Acinetobacter virus AB2	JX976549	Acinetobacter phage vB_AbaM-IME-AB2
Acinetobacter virus AbC62	KJ1817802	Acinetobacter phage YMC-13-01-C62									
Acinetobacter virus AP22	HE806280	Acinetobacter phage AP22									
B4virus	Bacillus virus B4	JN790865	Bacillus phage B4								
	Bacillus virus Bigbertha	KF669647	Bacillus phage BigBertha								
Bastillevirus	Bacillus virus Riley	KJ489402	Bacillus phage Riley								
	Bacillus virus Spock	KF669662	Bacillus phage Spock								
Bc431virus	Bacillus virus Troll	KF208639	Bacillus phage Troll								
	Bacillus virus Bastille	JF966203	Bacillus phage Bastille								
				Bc431virus	Bacillus virus CAM003	KJ489397	Bacillus phage CAM003				
					Bacillus virus Bc431	JX094431.1	Bacillus phage vB_BceM_Bc431v3				
					Bacillus virus Bcp1	KJ451625.1	Bacillus phage Bcp1				

Table 1 (continued)

Order	Family	Subfamily	Genus	Host bacteria	Species	Accession number (example species)	Isolate (example species)
	<i>Bcep78virus</i>		<i>Bcep78virus</i>	<i>Burkholderia</i>	<i>Bacillus virus BCP82</i>	KJ081346.1	<i>Bacillus</i> phage BCP8-2
					<i>Bacillus virus JBP901</i>	KJ676859.1	<i>Bacillus</i> phage JBP901
					<i>Burkholderia virus Bcep1</i>		
					<i>Burkholderia virus Bcep43</i>		
					<i>Burkholderia virus Bcep781</i>		
					<i>Burkholderia virus BcepNY3</i>		
					<i>Xanthomonas virus OP2</i>		
					<i>Burkholderia virus BcepMu</i>		
					<i>Burkholderia virus phiE255</i>		
					<i>Aeromonas virus 44RR2</i>	AY375531	<i>Aeromonas</i> phage 44RR2.8t
	<i>Biquartavirus</i> <i>Bx21virus</i> <i>Cd119virus</i>		<i>Biquartavirus</i> <i>Bx21virus</i> <i>Cd119virus</i>	<i>Aeromonas</i> <i>Mycobacterium</i> <i>Clostridium</i>	<i>Mycobacterium virus I3</i>		
					<i>Clostridium virus phiC2</i>		
					<i>Clostridium virus phiCD27</i>		
					<i>Clostridium virus phiCD119</i>		
					<i>Bacillus virus CP51</i>	KF554508.2	<i>Bacillus</i> phage CP-51
					<i>Bacillus virus JL</i>	KC595512.2	<i>Bacillus</i> phage JL
					<i>Bacillus virus Shanette</i>	KC595513.2	<i>Bacillus</i> phage Shanette
					<i>Escherichia virus CVM10</i>	GU903191	<i>Escherichia</i> phage vB_EcoM_ECO1230-10
					<i>Escherichia virus ep3</i>	KM360178	<i>Escherichia</i> phage vB_EcoM-ep3
					<i>Erwinia virus Ea214</i>		
	<i>Felixo1virus</i>		<i>Felixo1virus</i>	<i>Erwinia</i> <i>Escherichia</i>	<i>Escherichia virus AYO145A</i>	KR014248	<i>Escherichia</i> phage vB_EcoM_AYO145A
					<i>Escherichia virus EC6</i>	JX560968	<i>Escherichia</i> phage EC6
					<i>Escherichia virus JH2</i>	KF055347	<i>Escherichia</i> phage JH2
					<i>Escherichia virus VpaE1</i>	KM657822	<i>Escherichia</i> phage vB_EcoM-VpaE1
					<i>Escherichia virus wV8</i>		
					<i>Salmonella virus FelixO1</i>		
					<i>Salmonella virus HB2014</i>	KP010413	<i>Salmonella</i> phage HB-2014
					<i>Salmonella virus Mushroom</i>	KP143762	<i>Salmonella</i> phage Mushroom
					<i>Salmonella virus UAB87</i>	JN225449	<i>Enterobacteriophage</i> UAB_Phi87
					<i>Halomonas virus HAP1</i>		
	<i>Hapnavirus</i>		<i>Hapnavirus</i>	<i>Halomonas virus</i> <i>HAP1</i>	<i>Halomonas virus HAP1</i>		
					<i>Vibrio virus VP882</i>		
					<i>Pseudomonas virus Ab03</i>	LN610573	<i>Pseudomonas</i> phage vB_PaeM_PA01_Ab03
					<i>Pseudomonas virus KPP10</i>	AB472900.2	<i>Pseudomonas</i> phage KPP10
					<i>Pseudomonas virus PAKP3</i>	KC862299	<i>Pseudomonas</i> phage PAK_P3
					<i>Escherichia virus Mu</i>		
					<i>Halobacterium virus phiH</i>		
					<i>Bacillus virus Grass</i>	KF669652	<i>Bacillus</i> phage Grass
					<i>Bacillus virus NIT1</i>	AP013029	<i>Bacillus</i> phage phiNIT1

Table 1 (continued)

Order	Family	Subfamily	Genus	Host bacteria	Species	Accession number (example species)	Isolate (example species)
	<i>P1</i> virus		<i>Pakpuncavirus</i>	<i>Aeromonas</i> <i>Escherichia</i> <i>Pseudomonas</i>	<i>Bacillus virus SPG24</i>	AB930182	<i>Bacillus</i> phage SPG24
					<i>Aeromonas virus 43</i>		
					<i>Escherichia virus P1</i>	HE983845	<i>Pseudomonas</i> phage vB_PaeM_C2-10_Ab1
					<i>Pseudomonas virus CAb1</i>		<i>Pseudomonas</i> phage vB_PaeM_C2-10_Ab02
					<i>Pseudomonas virus CAb02</i>	LN610572	<i>Pseudomonas</i> phage JG004
					<i>Pseudomonas virus JG004</i>	GU988610.2	<i>Pseudomonas</i> phage PAK_P1
					<i>Pseudomonas virus PAKP1</i>	KC862297	<i>Pseudomonas</i> phage PAK_P4
					<i>Pseudomonas virus PAKP4</i>	KC862300	<i>Pseudomonas</i> phage PaP1
					<i>Pseudomonas virus PaP1</i>	HQ832595	<i>Pseudomonas</i> phage PaP1
					<i>Burkholderia virus BcepF1</i>		
	<i>Phunavirus</i>			<i>Burkholderia</i> <i>Pseudomonas</i>	<i>Pseudomonas virus l41</i>		<i>Pseudomonas</i> phage vB_PaeM_C1-14_Ab28
					<i>Pseudomonas virus Ab28</i>	LN610589	<i>Pseudomonas</i> phage DL60
					<i>Pseudomonas virus DL60</i>	KR054030	<i>Pseudomonas</i> phage DL60
					<i>Pseudomonas virus DL68</i>	KR054033	<i>Pseudomonas</i> phage DL68
					<i>Pseudomonas virus F8</i>		
					<i>Pseudomonas virus JG024</i>	GU815091	<i>Pseudomonas</i> phage JG024
					<i>Pseudomonas virus KPP12</i>	AB560486	<i>Pseudomonas</i> phage KPP12
					<i>Pseudomonas virus LBL3</i>		
					<i>Pseudomonas virus LMA2</i>		
					<i>Pseudomonas virus PBI</i>		
	<i>Phikzvirus</i>				<i>Pseudomonas virus SN</i>		
					<i>Pseudomonas virus EL</i>		
					<i>Pseudomonas virus Lin68</i>		
					<i>Pseudomonas virus phiKZ</i>		
					<i>Rhizobium virus RHEph4</i>	JX483876	<i>Rhizobium</i> phage RHEph04
					<i>Aeromonas virus 25</i>	DQ529280	<i>Aeromonas</i> phage 25
					<i>Aeromonas virus 31</i>		
					<i>Aeromonas virus Aes12</i>	JN377895	<i>Aeromonas</i> phage Aes012
					<i>Aeromonas virus Aes508</i>	JN377894	<i>Aeromonas</i> phage Aes508
					<i>Aeromonas virus AS4</i>	HM452125	<i>Aeromonas</i> phage phiAS4
	<i>Rheph4</i> virus <i>Secunda5</i> virus			<i>Rhizobium</i> <i>Aeromonas</i>	<i>Stenotrophomonas virus</i>	JX306041	<i>Stenotrophomonas</i> phage IME13
					<i>IME13</i>		
					<i>Yersinia virus RJRT</i>	HE956709	<i>Yersinia</i> phage phiRI-RT
					<i>Yersinia virus TGI</i>	KP202158	<i>Yersinia</i> phage vB_YenM_TG1
					<i>Bacillus virus G</i>		
					<i>Bacillus virus PBS1</i>		
					<i>Microcystis virus Ma-LMM01</i>		
					<i>Vibrio virus MAR</i>	JX556417	<i>Vibrio</i> phage vB_VpaM_MAR
					<i>Vibrio virus VHML</i>	AY133112	<i>Vibrio</i> phage VHML
					<i>Vibrio virus VP585</i>	FN297812	<i>Vibrio</i> phage VP585
	<i>VlmL</i> virus <i>Vil</i> virus			<i>Dickeya</i>	<i>Dickeya virus Limestone</i>		

Table 1 (continued)

Order	Family	Subfamily	Genus	Host bacteria	Species	Accession number (example species)	Isolate (example species)
Podoviridae (Icosahedral head with short tail)	Autographivirinae	Wphvirus	Kp34virus	<i>Escherichia virus</i>	<i>Escherichia virus CBA120</i>	JX128257	<i>Escherichia</i> phage ECML-4
					<i>Escherichia virus ECML4</i>		
				<i>Salmonella</i>	<i>Escherichia virus Phaxl</i>		
					<i>Salmonella virus Det7</i>	KP797973	<i>Salmonella</i> phage Det7
					<i>Salmonella virus Marshall</i>	KF669653	<i>Salmonella</i> phage Marshall
					<i>Salmonella virus Maynard</i>	KF669654	<i>Salmonella</i> phage Maynard
					<i>Salmonella virus SFP10</i>		
					<i>Salmonella virus SH19</i>		
					<i>Salmonella virus SJ2</i>	KJ174317	<i>Salmonella</i> phage vB_SalM_SJ2
					<i>Salmonella virus SJ3</i>	KJ174318	<i>Salmonella</i> phage vB_SalM_SJ3
					<i>Salmonella virus STM131</i>	JX181828	<i>Salmonella</i> phage STM13-1
					<i>Salmonella virus Vi1</i>		
				<i>Shigella</i>	<i>Shigella virus AG3</i>	HM144387	<i>Bacillus</i> phage W.Ph.
				<i>Bacillus</i>	<i>Bacillus virus WPh</i>	KF765493.2	<i>Klebsiella</i> phage F19
				<i>Klebsiella</i>	<i>Klebsiella virus F19</i>	AB716666	<i>Klebsiella</i> phage NTUH-K2044-K1-1
					<i>Klebsiella virus K244</i>		
					<i>Klebsiella virus KP34</i>	GQ413938.2	<i>Klebsiella</i> phage KP34
					<i>Klebsiella virus SU503</i>	KP708985	<i>Klebsiella</i> phage vB_KpnP_SU503
					<i>Klebsiella virus SU552A</i>	KP708986	<i>Klebsiella</i> phage vB_KpnP_SU552A
Picovirinae	P68virus	Unassigned	Phi29virus	<i>Pantoea</i>	<i>Pantoea virus Limelight</i>		
					<i>Pantoea virus Limezero</i>		
				<i>Pseudomonas</i>	<i>Pseudomonas virus LKA1</i>		
				<i>Erwinia virus</i>	<i>Pseudomonas virus phiKMV</i>		
				<i>Era103</i>	<i>Erwinia virus Era103</i>		
				<i>Escherichia</i>	<i>Escherichia virus K5</i>		
					<i>Escherichia virus K1-5</i>		
					<i>Escherichia virus K1E</i>		
				<i>Salmonella</i>	<i>Salmonella virus SP6</i>		
				<i>Escherichia</i>	<i>Escherichia virus T7</i>		
				<i>Kluyvera</i>	<i>Kluyvera virus Kvp1</i>		
				<i>Pseudomonas</i>	<i>Pseudomonas virus gh1</i>		
				<i>Prochlorococcus</i>	<i>Prochlorococcus virus PSSP7</i>		
					<i>Synechococcus virus P60</i>		
					<i>Synechococcus virus Syn5</i>		
				<i>Staphylococcus</i>	<i>Staphylococcus virus 44AHJD</i>		
					<i>Streptococcus virus C1</i>		
				<i>Bacillus</i>	<i>Bacillus virus B103</i>		

Table 1 (continued)

Order	Family	Subfamily	Genus	Host bacteria	Species	Accession number (example species)	Isolate (example species)
	Unassigned			Kurthia Actinomyces Mycoplasma Streptococcus Burkholderia	<i>Bacillus virus GA1</i>		
					<i>Bacillus virus phi29</i>		
					<i>Kurthia virus 6</i>		
					<i>Actinomyces virus Av1</i>		
					<i>Mycoplasma virus P1</i>		
					<i>Streptococcus virus Cpl</i>		
					<i>Burkholderia virus Bcep22</i>	AY349011	
					<i>Burkholderia virus Bcep102</i>	FJ937737	
					<i>Burkholderia virus</i>	JX104231	
					<i>Bcep102</i>		
	Bcep22likevirus			Bordetella Burkholderia	<i>Burkholderia virus DC1</i>	JN662425	
					<i>Bordetella virus BPP1</i>		
					<i>Burkholderia virus</i>		
					<i>BcepC6B</i>		
					<i>Cellulophaga virus Cba41</i>	KC821632	Cellulophaga phage phi4:1
					<i>Cellulophaga virus Cba172</i>	KC821609	Cellulophaga phage phi17:2
					<i>Escherichia virus phiV10</i>		
					<i>Salmonella virus Epsilon15</i>	AY625898	
					<i>Pseudomonas virus F116</i>	KC262634	
					<i>Pseudomonas virus APEC5</i>	KF192075	Escherichia phage vB_EcoP_PhAPEC5
	G7cvirus			Escherichia	<i>Escherichia virus APEC7</i>	KF562340	Escherichia phage vB_EcoP_PhAPEC7
					<i>Escherichia virus Bp4</i>	KJ135004.2	Escherichia phage Bp4
					<i>Escherichia virus EC1UPM</i>	KC206276.2	Escherichia phage EC1-UPM
					<i>Escherichia virus ECBP1</i>	JX415535	Escherichia phage ECBP1
					<i>Escherichia virus G7C</i>	HQ259105	Escherichia phage vB_EcoP_G7C
					<i>Escherichia virus IME11</i>	JX880034	Escherichia phage IME11
					<i>Shigella virus Sb1</i>	KF620435	Shigella phage pSb-1
					<i>Pseudomonas virus Ab09</i>	HG962375	Pseudomonas phage vB_PaeP_C2-10_Ab09
					<i>Pseudomonas virus LIT1</i>	FN422399	Pseudomonas phage LIT1
					<i>Pseudomonas virus PA26</i>	JX194238	Pseudomonas phage PA26
	Lit1virus			Shigella Pseudomonas	<i>Pseudomonas virus Ab22</i>	LN610578	Pseudomonas phage vB_PaeP_C2-10_Ab22
					<i>Pseudomonas virus CHU</i>	KP233880	Pseudomonas phage PhiCHU
					<i>Pseudomonas virus LUZ24</i>		
					<i>Pseudomonas virus PAA2</i>	KF856712	Pseudomonas phage phiIBB-PAA2
					<i>Pseudomonas virus PaP3</i>		
					<i>Pseudomonas virus PaP4</i>	KC294142	Pseudomonas phage PaP4
					<i>Pseudomonas virus TL</i>	HG518155	Pseudomonas phage TL
					<i>Escherichia virus N4</i>		
					<i>Escherichia</i>		

Table 1 (continued)

Order	Family	Subfamily	Genus	Host bacteria	Species	Accession number (example species)	Isolate (example species)
Siphoviridae (Icosahedral head with Long and noncontractile Tail)	Guernseyvirinae	Jerseyvirus	Nonanavirus	Salmonella	<i>Salmonella virus 9NA</i>	KJ802832	Salmonella phage 9NA
					<i>Salmonella virus SP069</i>	KC139649	Salmonella phage SP069
					<i>Salmonella virus HK620</i>		
					<i>Salmonella virus P22</i>		
			P22virus	Salmonella	<i>Salmonella virus ST64T</i>		
					<i>Shigella virus Sf6</i>		
					<i>Bacillus virus Page</i>	KF669655	Bacillus phage Page
					<i>Bacillus virus Palmer</i>	KP411017	Bacillus phage Palmer
					<i>Bacillus virus Pascal</i>	KM236247	Bacillus phage Pascal
			Pagevirus	Shigella Bacillus	<i>Bacillus virus Pony</i>	KF669660	Bacillus phage Pony
					<i>Bacillus virus Pookie</i>	KM236248	Bacillus phage Pookie
					<i>Escherichia virus ECB2</i>	KX415536	Escherichia phage ECBP2
					<i>Escherichia virus NJ01</i>	JX867715	Escherichia phage NJ01
			Phieco32virus	Escherichia	<i>Escherichia virus phiEco32</i>		
					<i>Escherichia virus Septimal1</i>	KF981730	Escherichia phage KBNP1711
					<i>Escherichia virus SU10</i>	KM044272	Escherichia phage vB_EcoP_SU10
			Unassigned	Hamiltonella Lactococcus Phormidium	<i>Hamiltonella virus APSE1</i>		
					<i>Lactococcus virus KSY1</i>		
					<i>Phormidium virus WMP3</i>		
					<i>Phormidium virus WMP4</i>		
			Vp5virus	Pseudomonas Roseobacter Vibrio	<i>Pseudomonas virus 1J9X</i>		
					<i>Roseobacter virus SIO1</i>		
					<i>Vibrio virus VpV262</i>		
					<i>Vibrio virus VC8</i>	JF712866	Vibrio phage phiVC8
Siphoviridae (Icosahedral head with Long and noncontractile Tail)	Guernseyvirinae	Jerseyvirus	Vp5virus	Vibrio	<i>Vibrio virus VP2</i>	AY505112	Vibriophage VP2
					<i>Vibrio virus VP5</i>	AY510084	Vibriophage VP5
					<i>Salmonella phage L13</i>	KC832325	Salmonella phage L13
					<i>Salmonella phage LSPA1</i>	KM272358	Salmonella phage LSPA1
			Guernseyvirinae	Salmonella	<i>Salmonella virus AG11</i>	JX297445	Salmonella phage vB_SenS_AG11
					<i>Salmonella virus Ent1</i>	HE775250	
					<i>Salmonella virus Jersey</i>	KF148055	
			K1Igvirus	Escherichia	<i>Salmonella virus SE2</i>	JQ007353	
					<i>Salmonella virus SETP3</i>	EF177456	
					<i>Salmonella virus SETP7</i>	KF562865	Salmonella phage SETP7
					<i>Salmonella virus SETP13</i>	KF562864	Salmonella phage SETP13
			Sp31virus	Salmonella	<i>Salmonella virus SP101</i>	KC139511	Salmonella phage FSL SP-101
					<i>Salmonella virus SS3e</i>	AY730274	
					<i>Salmonella virus wks3</i>	JX202565	
					<i>Escherichia virus K1G</i>	GU196277	Escherichia phage K1G
			Sp31virus	Salmonella	<i>Escherichia virus K1H</i>	GU196278	Escherichia phage K1H
					<i>Escherichia virus K1ind1</i>	GU196279	Escherichia phage K1ind1
					<i>Escherichia virus K1ind2</i>	GU196280	Escherichia phage K1ind2
					<i>Salmonella virus SP31</i>	KC139518	Salmonella phage FSL SP-031

Table 1 (continued)

Order	Family	Subfamily	Genus	Host bacteria	Species	Accession number (example species)	Isolate (example species)
	Tunavirinae	Kp36virus	Enterobacter	Klebsiella	Enterobacter virus F20	KP658157	Klebsiella phage 1513
					Klebsiella virus 1513	JF501022	Klebsiella phage KP36
					Klebsiella virus KP36	KF771237	Escherichia phage vB_EcoS_AHP42
					Escherichia virus AHP42		Escherichia phage vB_EcoS_AHS24
		Rogue1virus	Escherichia	Escherichia virus AHS24	KF771238	Escherichia phage vB_EcoS_AHS24	
				Escherichia virus AKS96	KF771239	Escherichia phage vB_EcoS_AKS96	
				Escherichia virus E41c	KJ668713	Escherichia phage e4/1c	
				Escherichia virus Eb49			
		Rtpvirus	Escherichia	Escherichia virus Jk06			
				Escherichia virus KP26	KC579452	Escherichia phage phiKP26	
				Escherichia virus Rogue1	KC333879	Escherichia phage Rogue1	
				Escherichia virus ACGM12	JN986845	Escherichia phage vB_Eco_ACG-M12	
	Tl1virus	Escherichia	Escherichia virus Rtp	AM156909	Escherichia phage Rtp		
			Escherichia virus ADB2	JX912252	Escherichia phage ADB-2		
			Escherichia virus T1				
			Shigella virus PSJ2	KP085586	Shigella phage pSF-2		
	Tlsvirus	Citrobacter	Shigella virus Shf11				
			Citrobacter virus Stevie	KM236241	Citrobacter phage Stevie		
			Escherichia virus TLS	AY308796	Escherichia phage Tls		
			Salmonella virus SP126	KC139513	Salmonella phage FSL SP-126		
	Unassigned	Andromedavirus	Bacillus	Cronobacter virus Exp2949-1			
				Bacillus virus Andromeda	KC330684		
				Bacillus virus Blastoid	KF669648		
				Bacillus virus Curly	KC330679		
		Barnyardvirus	Mycobacterium	Bacillus virus Eoghan	KC330680		
				Bacillus virus Fimi	KC330683		
				Bacillus virus Glittering	KF669651		
				Bacillus virus Riggi	KF669659		
				Bacillus virus Taylor	KC330682		
				Mycobacterium virus Barnyard	AY129339		
				Mycobacterium virus Konstantine	FJ174691		
				Mycobacterium virus Patience	JN412589		
			Mycobacterium virus Predator	EU770222			
			Mycobacterium virus Bignuz	JN412591			
			Mycobacterium virus Jebeks	JN572061			
			Staphylococcus virus I3	AF424783			

Table 1 (continued)

Order	Family	Subfamily	Genus	Host bacteria	Species	Accession number (example species)	Isolate (example species)
	<i>Bronvirus</i>			<i>Mycobacterium</i>	<i>Staphylococcus virus 77</i>	AY508486	Cellulophaga phage phi12:1 Cellulophaga phage phi17:1 Cellulophaga phage phi18:1 Cellulophaga phage phiST
					<i>Staphylococcus virus 108PVL</i>	AB243556	
					<i>Mycobacterium virus Bron</i>	HM152763	
					<i>Mycobacterium virus Faith1</i>	JF744988	
					<i>Mycobacterium virus Joedirt</i>	JF704108	
					<i>Mycobacterium virus Rumpelstiltskin</i>	JN680858	
					<i>Lactococcus virus bLL67</i>		
					<i>Lactococcus virus c2</i>		
					<i>Lactobacillus virus c5</i>	EU340421	
					<i>Lactobacillus virus LLKu</i>	AY739900	
	<i>C2virus</i>			<i>Lactococcus</i>	<i>Cellulophaga virus Cba121</i>	KC821613	Cellulophaga phage phi12:1 Cellulophaga phage phi17:1 Cellulophaga phage phi18:1 Cellulophaga phage phiST
					<i>Cellulophaga virus Cba171</i>	KC821617	
					<i>Cellulophaga virus Cba181</i>	KC821619	
					<i>Cellulophaga virus ST</i>	HQ634192	
					<i>Bacillus virus 250</i>	GU229986	
					<i>Bacillus virus IEBH</i>	EU874396	
					<i>Mycobacterium virus Charlie</i>	JN256079	
					<i>Mycobacterium virus Redi</i>	JN624851	
					<i>Mycobacterium virus Ardmore</i>	GU060500	
					<i>Mycobacterium virus Avani</i>	JQ809702	
	<i>Che8virus</i>			<i>Mycobacterium</i>	<i>Mycobacterium virus Boomer</i>	EU816590	Cellulophaga phage phi12:1 Cellulophaga phage phi17:1 Cellulophaga phage phi18:1 Cellulophaga phage phiST
					<i>Mycobacterium virus Che8</i>	AY129330	
					<i>Mycobacterium virus Che9d</i>	AY129336	
					<i>Mycobacterium virus Deadp</i>	JN698996	
					<i>Mycobacterium virus Dlane</i>	JF937093	
					<i>Mycobacterium virus Dorothy</i>	JX411620	
					<i>Mycobacterium virus Dopproduct</i>	JN859129	
					<i>Mycobacterium virus Drago</i>	JN542517	
					<i>Mycobacterium virus Fruitloop</i>	FJ174690	
					<i>Mycobacterium virus Gumbie</i>	JN398368	
	<i>Iblubesi</i>			<i>Mycobacterium</i>	<i>Mycobacterium virus Iblubesi</i>	JF937098	Cellulophaga phage phi12:1 Cellulophaga phage phi17:1 Cellulophaga phage phi18:1 Cellulophaga phage phiST
					<i>Mycobacterium virus Llij</i>	DQ398045	
					<i>Mycobacterium virus Mozy</i>	JF937102	
					<i>Mycobacterium virus Mutaformal3</i>	JN020142	

Order	Family	Subfamily	Genus	Host bacteria	Species	Accession number (example species)	Isolate (example species)

	<i>Mycobacterium virus Pucc40</i>	FJ174692
	<i>Mycobacterium virus PMC</i>	DQ398050
	<i>Mycobacterium virus Ramsey</i>	FJ174693
	<i>Mycobacterium virus Rockyhorror</i>	JF704117
	<i>Mycobacterium virus SG4</i>	JN699012
	<i>Mycobacterium virus Shauna1</i>	JN020141
	<i>Mycobacterium virus Shilam</i>	JN020143
	<i>Mycobacterium virus Spartacus</i>	JQ300538
	<i>Mycobacterium virus Taj</i>	JX121091
	<i>Mycobacterium virus Tweety</i>	EF536069
	<i>Mycobacterium virus Wee</i>	HQ728524
	<i>Mycobacterium virus Yoshi</i>	JF704115
	<i>Mycobacterium virus Babstiella</i>	JN699001
	<i>Mycobacterium virus Brujita</i>	FJ168659
	<i>Mycobacterium virus Che9c</i>	AY129333
	<i>Salmonella virus Chi</i>	JX094499
	<i>Salmonella virus FSLSP030</i>	KC139519
	<i>Salmonella virus FSLSP088</i>	KC139512
	<i>Salmonella virus iEP55</i>	KC677662
	<i>Salmonella virus SPN19</i>	JN871591
	<i>Mycobacterium virus 244</i>	DQ398041
	<i>Mycobacterium virus Bask21</i>	JF937091
	<i>Mycobacterium virus CJW1</i>	AY129331
	<i>Mycobacterium virus Eureka</i>	JN412590
	<i>Mycobacterium virus Kosya</i>	EU816591
	<i>Mycobacterium virus Porky</i>	EU816588
	<i>Mycobacterium virus Pumpkin</i>	GQ303265
	<i>Mycobacterium virus Sirduracell</i>	JF937106
	<i>Mycobacterium virus Toto</i>	JN006061
	<i>Mycobacterium virus Cornudog</i>	AY129335
	<i>Mycobacterium virus Firecracker</i>	JN698993
	<i>Pseudomonas virus D3112</i>	AY394005
	<i>Pseudomonas virus DMS3</i>	DO631426
<i>Che9c</i>	<i>Mycobacterium</i>	
<i>Chivirus</i>	<i>Salmonella</i>	
<i>Cjw1</i>	<i>Mycobacterium</i>	
<i>Corndog</i>	<i>Mycobacterium</i>	
<i>D3112</i>	<i>Pseudomonas</i>	

Table 1 (continued)

Order	Family	Subfamily	Genus	Host bacteria	Species	Accession number (example species)	Isolate (example species)
					<i>Pseudomonas virus</i> <i>FHA0480</i>	JN808773	
					<i>Pseudomonas virus LPB1</i>	HE584812	
					<i>Pseudomonas virus MP22</i>	DQ873690	
					<i>Pseudomonas virus MP29</i>	EU272036	
					<i>Pseudomonas virus MP38</i>	EU272037	
					<i>Pseudomonas virus</i> <i>PAIKOR</i>	HM624080	
			<i>D3virus</i>	<i>Pseudomonas</i>	<i>Pseudomonas virus D3</i>	AF165214	
			<i>E125virus</i>	<i>Burkholderia</i>	<i>Pseudomonas virus PMG1</i>	HQ711985	
					<i>Burkholderia virus phi6442</i>	CP000625	
					<i>Burkholderia virus</i> <i>phi1026b</i>	AY453853	
			<i>Ff47virus</i>	<i>Mycobacterium</i>	<i>Burkholderia virus phiE125</i>	AF447491	
			<i>Hk578virus</i>	<i>Escherichia</i>	<i>Mycobacterium virus Ff47</i>	JX901189	
					<i>Mycobacterium virus Muddy</i>	KF024728	
					<i>Escherichia virus HK578</i>	JQ086375	
					<i>Escherichia virus JLI1</i>	JX865427	
					<i>Escherichia virus SSL2009a</i>	FJ750948	
				<i>Shigella</i>	<i>Shigella virus EP23</i>	JN984867	
				<i>Sodalis</i>	<i>Sodalis virus SO1</i>	GQ502199	
			<i>L5virus</i>	<i>Mycobacterium</i>	<i>Mycobacterium virus Alma</i>	JN699005	
					<i>Mycobacterium virus Arturo</i>	JX307702	
					<i>Mycobacterium virus Astro</i>	JX015524	
					<i>Mycobacterium virus</i> <i>Backyardigan</i>	JF704093	
					<i>Mycobacterium virus</i> <i>BBPrebs31</i>	JF957057	
					<i>Mycobacterium virus</i> <i>Benedict</i>	JN083852	
					<i>Mycobacterium virus</i> <i>Bethlehem</i>	AY500153	
					<i>Mycobacterium virus</i> <i>Billknuckles</i>	JN699000	
					<i>Mycobacterium virus Bruns</i>	JN698998	
					<i>Mycobacterium virus Bxb1</i>	AF271693	
					<i>Mycobacterium virus Bxz2</i>	AY129332	
					<i>Mycobacterium virus Che12</i>	DQ398043	
					<i>Mycobacterium virus Cuco</i>	JN408459	
					<i>Mycobacterium virus D29</i>		
					<i>Mycobacterium virus Doom</i>	JN153085	
					<i>Mycobacterium virus Erich</i>	JN049605	
					<i>Mycobacterium virus</i> <i>Euphoria</i>	JN153086	
						JF704107	
							<i>Mycobacterium virus FF47</i>
							<i>Mycobacterium virus Muddy</i>

Table 1 (continued)

Order	Family	Subfamily	Genus	Host bacteria	Species	Accession number (example species)	Isolate (example species)
					<i>Mycobacterium virus George</i>	JF704097	
					<i>Mycobacterium virus Gladiator</i>	JX307704	
					<i>Mycobacterium virus Goose</i>	JF937094	
					<i>Mycobacterium virus Hammer</i>	JF957058	
					<i>Mycobacterium virus Heldan</i>	EU744251	
					<i>Mycobacterium virus Jasper</i>	JF937099	
					<i>Mycobacterium virus JC27</i>	JN699019	
					<i>Mycobacterium virus Jeffabunny</i>	JF704098	
					<i>Mycobacterium virus JHC117</i>	EU744248	
					<i>Mycobacterium virus KBG</i>	JF937110	
					<i>Mycobacterium virus Kssjeb</i>	JN699016	
					<i>Mycobacterium virus Kugel</i>	JF937100	
					<i>Mycobacterium virus L5</i>	JN699015	
					<i>Mycobacterium virus Lesedi</i>		
					<i>Mycobacterium virus LHTSCC</i>	EU744249	
					<i>Mycobacterium virus lockley</i>	JX307705	
					<i>Mycobacterium virus Marcell</i>	JF704101	
					<i>Mycobacterium virus Microwolf</i>	JN020140	
					<i>Mycobacterium virus Mrgordo</i>	JF937103	
					<i>Mycobacterium virus Museum</i>	JQ698665	
					<i>Mycobacterium virus Nepal</i>	JF704110	
					<i>Mycobacterium virus Packman</i>	GQ303263	
					<i>Mycobacterium virus Peaches</i>	JN572689	
					<i>Mycobacterium virus Perseus</i>	EU744250	
					<i>Mycobacterium virus Pukovnik</i>	JX411619	
					<i>Mycobacterium virus Rebeuca</i>	GU339467	
					<i>Mycobacterium virus Redrock</i>	JN398369	

Table 1 (continued)

Order	Family	Subfamily	Genus	Host bacteria	Species	Accession number (example species)	Isolate (example species)
					<i>Mycobacterium virus Ridgecb</i>	JF704111	
					<i>Mycobacterium virus Rockstar</i>	JN831654	
					<i>Mycobacterium virus Saintus</i>	GU247132	
					<i>Mycobacterium virus Skipole</i>	EU826470	
					<i>Mycobacterium virus Solon</i>	JF937108	
					<i>Mycobacterium virus Switzer</i>	JF946695	
					<i>Mycobacterium virus SWU1</i>	JN400277	
					<i>Mycobacterium virus Ta17a</i>	JQ684677	
					<i>Mycobacterium virus Tiger</i>	JF957060	
					<i>Mycobacterium virus Timshel</i>	JN408461	
					<i>Mycobacterium virus Trixie</i>	JN408460	
					<i>Mycobacterium virus Turbido</i>	JQ512844	
					<i>Mycobacterium virus Twister</i>	AY500152	
					<i>Mycobacterium virus U2</i>	JN687951	
					<i>Mycobacterium virus Violet</i>	HM755814	
					<i>Mycobacterium virus Wonder</i>	JN116827	
				<i>Rhodococcus</i>	<i>Rhodococcus virus RER2</i>	JN116826	
					<i>Rhodococcus virus RGL3</i>		
				<i>Escherichia</i>	<i>Escherichia virus HK022</i>		
					<i>Escherichia virus HK97</i>		
					<i>Escherichia virus Lambda</i>		
				<i>Mycobacterium</i>	<i>Mycobacterium virus Halo</i>	DQ398042	
					<i>Mycobacterium virus Liefie</i>	JN412593	
					<i>Escherichia virus N15</i>		
				<i>Escherichia</i>	<i>Escherichia virus 9 g</i>	KJ419279	Enterobacteria phage 9 g
					<i>Escherichia virus JenK1</i>	KP719134	Enterobacteria phage JenK1
				<i>Escherichia</i>	<i>Escherichia virus JenP1</i>	KP719132	Enterobacteria phage JenP1
					<i>Escherichia virus JenP2</i>	KP719133	Enterobacteria phage JenP2
				<i>Mycobacterium</i>	<i>Mycobacterium virus Baka</i>	JF937090	
					<i>Mycobacterium virus Courthouse</i>	JN698997	
					<i>Mycobacterium virus Littlee</i>	JF937101	
				<i>Mycobacterium virus</i>	<i>Mycobacterium virus</i>	AY129338	
					<i>Omega</i>		
				<i>Mycobacterium virus</i>	<i>Mycobacterium virus</i>	JF957059	
					<i>Optimus</i>		

Table 1 (continued)

Order	Family	Subfamily	Genus	Host bacteria	Species	Accession number (example species)	Isolate (example species)
					<i>Mycobacterium virus Thibault</i>	JN201525	
	<i>P23virus</i>			<i>Thermus</i>	<i>Thermus virus P23-45</i>	EU100883	
					<i>Thermus virus P74-26</i>	EU100884	
	<i>P70virus</i>			<i>Listeria</i>	<i>Listeria virus LP26</i>	KJ094020	Listeria phage LP-026
					<i>Listeria virus LP37</i>	JX126920.2	Listeria phage LP-037
					<i>Listeria virus LP110</i>	JX126919	Listeria phage LP-110
					<i>Listeria virus LP114</i>	KJ094021	Listeria phage LP-114
					<i>Listeria virus P70</i>	JX442241	Listeria phage P70
	<i>PhiIvirus</i>			<i>Mycobacterium</i>	<i>Mycobacterium virus PBI1</i>	DQ398047	
					<i>Mycobacterium virus</i>	JN699007	
					<i>Acadian</i>		
					<i>Mycobacterium virus</i>	JN699003	
					<i>Athens</i>		
					<i>Mycobacterium virus</i>	JF704094	
					<i>Chrismnich</i>		
					<i>Mycobacterium virus</i>	DQ398044	
					<i>Cooper</i>		
					<i>Mycobacterium virus Gadjet</i>	JN698992	
					<i>Mycobacterium virus Nigel</i>	EU770221	
					<i>Mycobacterium virus Oline</i>	JN192463	
					<i>Mycobacterium virus Pgl</i>	AF547430	
					<i>Mycobacterium virus</i>	DQ398049	
					<i>Pipefish</i>		
					<i>Mycobacterium virus</i>	AY129334	
					<i>Rosebush</i>		
					<i>Mycobacterium virus</i>	JN699011	
					<i>Stinger</i>		
					<i>Mycobacterium virus</i>	JF704104	
					<i>Zemanar</i>		
	<i>Phic31virus</i>			<i>Streptomyces</i>	<i>Streptomyces virus phiBT1</i>	AJ550940	
					<i>Streptomyces virus phiC31</i>		
					<i>Streptomyces virus TGI</i>	JX182372	
					<i>Caulobacter virus Karma</i>	JX100811	
	<i>Phicbkvirus</i>			<i>Caulobacter</i>	<i>Caulobacter virus Magneto</i>	JX100812	
					<i>Caulobacter virus phiCbK</i>	JX100813	
					<i>Caulobacter virus Rogue</i>	JX100814	
					<i>Caulobacter virus Swift</i>	JX100809	
					<i>Staphylococcus virus 11</i>	AF424781	
	<i>Phietavirus</i>			<i>Staphylococcus</i>	<i>Staphylococcus virus 29</i>	AY954964	
					<i>Staphylococcus virus 37</i>	AY954958	
					<i>Staphylococcus virus 53</i>	AY954952	
					<i>Staphylococcus virus 55</i>	AY954963	
					<i>Staphylococcus virus 69</i>	AY954951	
					<i>Staphylococcus virus 71</i>	AY954962	

Table 1 (continued)

Order	Family	Subfamily	Genus	Host bacteria	Species	Accession number (example species)	Isolate (example species)
					<i>Staphylococcus virus 80</i>	DQ908929	
					<i>Staphylococcus virus 85</i>	AY954953	
					<i>Staphylococcus virus 88</i>	AY954966	
					<i>Staphylococcus virus 92</i>	AY954967	
					<i>Staphylococcus virus 96</i>	AY954960	
					<i>Staphylococcus virus 187</i>	AY954950	
					<i>Staphylococcus virus 52a</i>	AY954965	
					<i>Staphylococcus virus 80alpha</i>	DQ517338	
					<i>Staphylococcus virus CNPH82</i>	DQ831957	
					<i>Staphylococcus virus EW</i>	AY954959	
					<i>Staphylococcus virus IPLA5</i>	JN192400	
					<i>Staphylococcus virus IPLA7</i>	JN192401	
					<i>Staphylococcus virus IPLA88</i>	EU861004	
					<i>Staphylococcus virus PH15</i>	DQ834250	
					<i>Staphylococcus virus phiETA</i>	AP001553	
					<i>Staphylococcus virus phiETA2</i>	AP008953	
					<i>Staphylococcus virus phiETA3</i>	AP008954	
					<i>Staphylococcus virus phiMR11</i>	AB370268	
					<i>Staphylococcus virus phiMR25</i>	AB370205	
					<i>Staphylococcus virus phiNM1</i>	DQ530359	
					<i>Staphylococcus virus phiNM2</i>	DQ530360	
					<i>Staphylococcus virus phiNM4</i>	DQ530362	
					<i>Staphylococcus virus SAP26</i>	GU477322	
					<i>Staphylococcus virus X2</i>	AY954968	
			<i>Phifelvirus</i>	<i>Enterococcus</i>	<i>Enterococcus virus FL1</i>	GQ478081	
					<i>Enterococcus virus FL2</i>	GQ478084	
					<i>Enterococcus virus FL3</i>	GQ478086	
			<i>Phijlivirus</i>	<i>Lactobacillus</i>	<i>Lactobacillus virus ATCC8014</i>	JX486087	
					<i>Lactobacillus virus phiJL1</i>	AY236756	
				<i>Pediococcus</i>	<i>Pediococcus virus ePI</i>	JN051154	
			<i>Psavirus</i>	<i>Listeria</i>	<i>Listeria virus LP302</i>	JX120799.2	Listeria phage LP-030-2
			<i>Psimunavirus</i>	<i>Methanobacterium</i>	<i>Listeria virus PSA</i>	AJ312240.2	Listeria phage PSA

Table 1 (continued)

Order	Family	Subfamily	Genus	Host bacteria	Species	Accession number (example species)	Isolate (example species)
					<i>Methanobacterium virus psiM1</i>		
	<i>Reyvirus</i>			<i>Mycobacterium</i>	<i>Mycobacterium virus Bongo</i>	JN699628	
	<i>Sapovirus</i>			<i>Enterococcus</i>	<i>Mycobacterium virus Rey</i>	JF937105	
					<i>Enterococcus virus BC611</i>	AB712291	
					<i>Enterococcus virus IMEEF1</i>	KF192053	
					<i>Enterococcus virus SAP6</i>	JF731128	
					<i>Enterococcus virus VD13</i>	KJ094032	
	<i>Septina3virus</i>			<i>Streptococcus</i>	<i>Streptococcus virus SPQ51</i>	HE962497	
				<i>Burkholderia</i>	<i>Burkholderia virus KLI1</i>	JF939047	Burkholderia phage vB_BceS_KLI1
				<i>Pseudomonas</i>	<i>Pseudomonas virus 73</i>	NC_007806	Pseudomonas phage 73
					<i>Pseudomonas virus Ab26</i>	HG962376	Pseudomonas phage vB_PaeS_SCH_Ab26
					<i>Pseudomonas virus Kakheti25</i>	JQ307387	Pseudomonas phage vB_Pae-Kakheti25
	<i>Seuravirus</i>			<i>Escherichia</i>	<i>Escherichia virus Cajan</i>	KP064094	Escherichia phage CAjan
	<i>Sextaevirus</i>			<i>Staphylococcus</i>	<i>Escherichia virus Seurat</i>	KM236243	Escherichia phage Seurat
					<i>Staphylococcus virus SEP9</i>	KF929199	Staphylococcus phage vB_SepS_SEP9
					<i>Staphylococcus virus</i>	KJ804259	Staphylococcus phage 6ec
					<i>Sextaev</i>		
	<i>Sfi11virus</i>			<i>Streptococcus</i>	<i>Streptococcus virus 858</i>	EF529515	
					<i>Streptococcus virus 2972</i>	AY699705	
					<i>Streptococcus virus ALQ132</i>	FJ226752	
					<i>Streptococcus virus OI205</i>	U88974	
					<i>Streptococcus virus Sfi11</i>	AF158600	
					<i>Streptococcus virus 7201</i>	AF145054	
	<i>Sfi21dt1virus</i>			<i>Streptococcus</i>	<i>Streptococcus virus DT1</i>	AF085222	
					<i>Streptococcus virus phiAbc2</i>	FJ236310	
					<i>Streptococcus virus Sfi19</i>	AF115102	
					<i>Streptococcus virus Sfi21</i>	AF115103	
	<i>Sitaravirus</i>			<i>Paenibacillus</i>	<i>Paenibacillus virus Diva</i>	KP296791	Paenibacillus phage Diva
					<i>Paenibacillus virus Hb10c2</i>	KP202972	Paenibacillus phage HB10c2
					<i>Paenibacillus virus Rani</i>	KP296793	Paenibacillus phage Rani
					<i>Paenibacillus virus Shelly</i>	KP296795	Paenibacillus phage Shelly
					<i>Paenibacillus virus Sitara</i>	KP296796	Paenibacillus phage Sitara
	<i>Sk1virus</i>			<i>Lactococcus</i>	<i>Lactococcus virus 712</i>	DQ227763	
					<i>Lactococcus virus ASCC191</i>	JQ740813	
					<i>Lactococcus virus ASCC273</i>	JQ740788	
					<i>Lactococcus virus ASCC281</i>	JQ740787	
					<i>Lactococcus virus ASCC465</i>	JQ740804	
					<i>Lactococcus virus ASCC532</i>	JQ740789	
					<i>Lactococcus virus Bibb29</i>	EU221285	
					<i>Lactococcus virus bLI170</i>	AF009630	

Table 1 (continued)

Order	Family	Subfamily	Genus	Host bacteria	Species	Accession number (example species)	Isolate (example species)
					<i>Lactococcus virus CB13</i>	FJ848882	Bacillus phage Slash Bacillus phage Stahl Bacillus phage Staley Bacillus phage Stills Vibrio phage vB_VpaS_ MAR10 Vibrio phage SSP002
					<i>Lactococcus virus CB14</i>	FJ848883	
					<i>Lactococcus virus CB19</i>	FJ848884	
					<i>Lactococcus virus CB20</i>	FJ848885	
					<i>Lactococcus virus JJ50</i>	DQ227764	
					<i>Lactococcus virus P2</i>	GQ979703	
					<i>Lactococcus virus P008</i>	DQ054536	
					<i>Lactococcus virus skl</i>	AF011378	
					<i>Lactococcus virus SI4</i>	FJ848881	
					<i>Bacillus virus Slash</i>	KF669661	
				<i>Bacillus</i>	<i>Bacillus virus Stahl</i>	KP696447	Bacillus phage Slash Bacillus phage Stahl Bacillus phage Staley Bacillus phage Stills Vibrio phage vB_VpaS_ MAR10 Vibrio phage SSP002
					<i>Bacillus virus Staley</i>	KF669663	
					<i>Bacillus virus Stills</i>	KP696448	
					<i>Bacillus virus SPbeta</i>		
					<i>Vibrio virus MAR10</i>	JX556418	
					<i>Vibrio virus SSP002</i>	JQ692107	
					<i>Escherichia virus AKFV33</i>		
					<i>Escherichia virus BF23</i>		
					<i>Escherichia virus DT57C</i>	KM979354	
					<i>Escherichia virus EPS7</i>		
				<i>Escherichia</i>	<i>Escherichia virus FFH1</i>	KJ190157	Escherichia phage DT57C Escherichia phage vB_EcoS_ FFH_1 Salmonella phage Shivani Salmonella phage Stitch Macncheese Mycobacterium virus Pixie Mycobacterium virus TM4 Bacillus virus BMBtp2 Bacillus virus TP21 Staphylococcus virus 47 Staphylococcus virus 3a Staphylococcus virus 42e
					<i>Escherichia virus H8</i>		
					<i>Escherichia virus T5</i>		
					<i>Salmonella virus Shivani</i>	KP143763	
					<i>Salmonella virus SPC35</i>		
					<i>Salmonella virus Stitch</i>	KM236244	
					<i>Mycobacterium virus Anaya</i>	JF704106	
					<i>Mycobacterium virus Angelica</i>	HM152764	
					<i>Mycobacterium virus Crimd</i>	HM152767	
					<i>Mycobacterium virus Fionn</i>	JN831653	
				<i>Mycobacterium</i>	<i>Mycobacterium virus Jaws</i>	JN185608	Escherichia phage DT57C Escherichia phage vB_EcoS_ FFH_1 Salmonella phage Shivani Salmonella phage Stitch Macncheese Mycobacterium virus Pixie Mycobacterium virus TM4 Bacillus virus BMBtp2 Bacillus virus TP21 Staphylococcus virus 47 Staphylococcus virus 3a Staphylococcus virus 42e
					<i>Mycobacterium virus Larva</i>	JN243855	
					<i>Mycobacterium virus Macncheese</i>	JX042579	
					<i>Mycobacterium virus Pixie</i>	JF937104	
					<i>Mycobacterium virus TM4</i>	AF068845	
					<i>Bacillus virus BMBtp2</i>	JX887877	
					<i>Bacillus virus TP21</i>	EU887664	
					<i>Staphylococcus virus 47</i>	AY954957	
					<i>Staphylococcus virus 3a</i>	AY954956	
					<i>Staphylococcus virus 42e</i>	AY954955	
				<i>Bacillus</i>	<i>Staphylococcus virus 42e</i>	EU861005	Escherichia phage DT57C Escherichia phage vB_EcoS_ FFH_1 Salmonella phage Shivani Salmonella phage Stitch Macncheese Mycobacterium virus Pixie Mycobacterium virus TM4 Bacillus virus BMBtp2 Bacillus virus TP21 Staphylococcus virus 47 Staphylococcus virus 3a Staphylococcus virus 42e

Table 1 (continued)

Order	Family	Subfamily	Genus	Host bacteria	Species	Accession number (example species)	Isolate (example species)
Unassigned	Corticoviridae (Circular dsDNA, Lipid present in capsid) Cystoviridae (dsRNA, icosahedral. Enveloped) Inoviridae (Circular ssDNA, Filamentous)				<i>Staphylococcus virus IPLA35</i>		
					<i>Staphylococcus virus phi12</i>	AF424782	
					<i>Staphylococcus virus phiSLT</i>	AB045978	
			<i>Whetavirus</i> <i>Xp10virus</i>	<i>Bacillus</i> <i>Xanthomonas</i>	<i>Bacillus virus Wbeta</i>	DQ289555	
					<i>Xanthomonas virus CP1</i>	AB720063	
					<i>Xanthomonas virus OP1</i>	AP008979	
					<i>Xanthomonas virus phi17</i>	EU717894	
					<i>Xanthomonas virus Xop411</i>	DQ777876	
			<i>Yuavirus</i>	<i>Alphaproteobacteria</i>	<i>Xanthomonas virus Xp10</i>	AY299121	
					<i>Alphaproteobacteria virus phiJ1001</i>		
			<i>Corticovirus</i>	<i>Pseudomonas</i>	<i>Pseudomonas virus M6</i>		
					<i>Pseudomonas virus Yua</i>		
			<i>Cystovirus</i>	<i>Pseudomonas</i>	<i>Pseudomonas virus PM2</i>		
					<i>Pseudomonas virus phi6</i>		
			<i>Inovirus</i>	<i>Escherichia</i>	<i>Escherichia virus AE2</i>		
					<i>Escherichia virus DeltaA</i>		
					<i>Escherichia virus Ec9</i>		
					<i>Escherichia virus f1</i>		
					<i>Escherichia virus fd</i>		
					<i>Escherichia virus HR</i>		
					<i>Escherichia virus I22</i>		
					<i>Escherichia virus If1</i>		
					<i>Escherichia virus M13</i>		
					<i>Escherichia virus PR64FS</i>		
					<i>Escherichia virus SF</i>		
					<i>Escherichia virus tf1</i>		
					<i>Escherichia virus X</i>		
					<i>Escherichia virus X2</i>		
					<i>Escherichia virus ZJ2</i>		
					<i>Pseudomonas virus Pj1</i>		
					<i>Pseudomonas virus Pj2</i>		
					<i>Pseudomonas virus Pj3</i>		
				<i>Salmonella</i>	<i>Salmonella virus C2</i>		
					<i>Salmonella virus IKE</i>		
					<i>Vibrio virus 493</i>		
					<i>Vibrio virus CTXphi</i>		
					<i>Vibrio virus fs1</i>		
				<i>Vibrio</i>	<i>Vibrio virus fs2</i>		
					<i>Vibrio virus v6</i>		

Table 1 (continued)

Order	Family	Subfamily	Genus	Host bacteria	Species	Accession number (example species)	Isolate (example species)
				<i>Xanthomonas</i>	<i>Vibrio virus Vj12</i>		
					<i>Vibrio virus Vj33</i>		
					<i>Vibrio virus VSK</i>		
					<i>Xanthomonas virus Cf16</i>		
					<i>Xanthomonas virus Cf1c</i>		
					<i>Xanthomonas virus Cf1t</i>		
					<i>Xanthomonas virus Cf1v</i>		
					<i>Xanthomonas virus Lf</i>		
					<i>Xanthomonas virus Xf</i>		
					<i>Xanthomonas virus Xfo</i>		
			<i>Plectrovirus</i>	<i>Acholeplasma Spiroplasma</i>	<i>Xanthomonas virus Xfv</i>		
					<i>Acholeplasma virus MV-L51</i>		
					<i>Spiroplasma virus laa</i>		
					<i>Spiroplasma virus C74</i>		
					<i>Spiroplasma virus KC3</i>		
					<i>Spiroplasma virus R8A2B</i>		
					<i>Spiroplasma virus S102</i>		
					<i>Spiroplasma virus T78</i>		
					<i>Escherichia virus F1</i>		
					<i>Escherichia virus Qbeta</i>		
	<i>Leviviridae</i> (ssRNA+, icosahedral)	<i>Bullavirinae</i>	<i>Allolevivirus</i>	<i>Escherichia</i>	<i>Escherichia virus BZ13</i>		
					<i>Escherichia virus MS2</i>		
					<i>Escherichia virus alpha3</i>		
					<i>Escherichia virus ID21</i>	DQ079870	Escherichia phage ID21
					<i>Escherichia virus ID32</i>	DQ079871	Escherichia phage ID32
					<i>Escherichia virus ID62</i>	DQ079876	Escherichia phage ID62
					<i>Escherichia virus NC28</i>	DQ079875	Escherichia phage NC28
					<i>Escherichia virus NC29</i>	DQ079879	Escherichia phage NC29
					<i>Escherichia virus NC35</i>	DQ079872	Escherichia phage NC35
					<i>Escherichia virus phiK</i>		
			<i>G4microvirus</i>	virus G4	<i>Escherichia virus Stl</i>		
					<i>Escherichia virus WA45</i>	DQ079874	Escherichia phage WA45
					<i>Escherichia virus G4</i>		
					<i>Escherichia virus ID52</i>	DQ079877	Escherichia phage ID52
					<i>Escherichia virus Talmos</i>	DQ079869	Escherichia phage ID2 Moscow/ID/2001
					<i>Escherichia virus phiX174</i>		
					<i>Bdellovibrio virus MAC1</i>		
					<i>Bdellovibrio virus MH2K</i>		
					<i>Chlamydia virus Chp1</i>		
					<i>Chlamydia virus Chp2</i>		
	<i>Gokushovirinae</i>		<i>Phix174microvirus</i> <i>Bdellovibrio</i>	<i>Escherichia Bdellovibrio</i>	<i>Chlamydia virus CPAR39</i>		
					<i>Chlamydia virus CPG1</i>		
					<i>Spiroplasma virus SpV4</i>		
					<i>Acholeplasma virus L2</i>		
			<i>Spiromicrovirus</i> <i>Plasmavirus</i>	<i>Spiroplasma Acholeplasma</i>			

Table 1 (continued)

Order	Family	Subfamily	Genus	Host bacteria	Species	Accession number (example species)	Isolate (example species)
Plasmaviridae (Circular dsDNA, Lipid Envelop dsDNA)	Pleolipoviridae (Pleomorphic membrane vesicle enclosing single and double stranded DNA genomes)		Alphapleolipovirus	Haloarcula	Haloarcula virus HHPV-1	GU321093	Haloarcula hispanica pleomorphic virus 1
					Haloarcula virus HHPV-2	KF056323	Haloarcula hispanica pleomorphic virus 2
Halorubrum virus HRPV-1					FJ685651	Halorubrum pleomorphic virus 1	
Halorubrum virus HRPV-2					JN882264	Halorubrum pleomorphic virus 2	
Halorubrum virus HRPV-6					JN882266	Halorubrum pleomorphic virus 6	
Halogeometricum virus HGPV-1					JN882267	Halogeometricum pleomorphic virus 1	
Sphaerolipoviridae (Icosahedral, tailless haloarchaeal)			Betapleolipovirus	Halogeometricum	Halorubrum virus HRPV-3	JN882265	Halorubrum pleomorphic virus 3
					Haloarcula virus His2	AF191797	His2 virus
					Haloarcula hispanica icosahedral virus 2	JN968479	
					Haloarcula hispanica virus PHI	KC252997	
					Haloarcula hispanica virus SH1	AY950802	
					Natrinema virus SNJ1	AY048850	
Tectiviridae (Icosahedral Non-Enveloped Linear dsDNA)			Betasphaerolipovirus Gammaphaerolipovirus Tectivirus Unassigned	Natrinema Thermus Bacillus Salmonella Thermus Alicyclobacillus	Thermus virus IN93	AB063393	
					Thermus virus P23-77	GQ403789	
					Bacillus virus AP50		
					Bacillus virus Bam35		
					Salmonella virus PRD1		
					Thermus virus P37-14		
				Alicyclobacillus virus NS11			

proteins (OMPs) for transport and diffusion of nutrients. These act as phage receptors, and in some infection strategies, they are essential for adsorption of phage particles as well (Bugla-Ploskonska et al. 2007; Rakhuba et al. 2010). In comparison, teichoic acids (peptidoglycan interspersed with acid polysaccharides) present in the Gram-positive bacteria cell wall act as receptors for their corresponding phages (Brown et al. 2013; León and Bastías 2015).

The penetration processes in bacterial cell also vary in different phage groups. In general, myoviridae phage inserts its genetic material into the bacterial cell by using a syringe-like movement of its tail (Fig. 2). After receptor recognition, in a reversible binding mode, the phage particle attaches its base plate with the bacterial surface utilizing the flexing activity of tail fibers. The phage takes sufficient time to make its surface binding irreversible. Thereafter, with the help of ATP, the contraction of its tail takes place, along with insertion of its genetic material. On the contrary, podoviridae phage, which is devoid of the tail part of the myoviridae phage, inserts its genetic material after enzymatically degrading a portion of the bacterial cell membrane using its small, tooth-like tail fibers (Rakhuba et al. 2010; Brown et al. 2013; León and Bastías 2015).

Phages undergo two possible life cycles: lytic and lysogenic cycles.

Lytic cycle

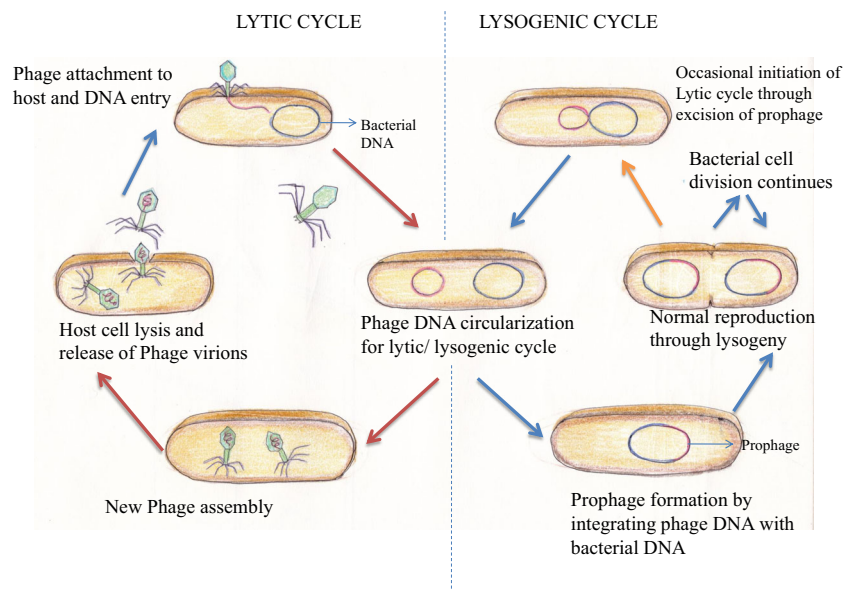
Phages, which are in their lytic phase, are called virulent phages. During the lytic cycle, lysis (death) of host bacterium occurs as the phage multiplies in great numbers within the host, following which the phage disintegrates and ruptures the host cell for release of new phage particles (Fig. 2).

The lytic cycle initiates with the attachment of the phage on the bacteria with the aid of a complex of proteins (Karlsson et al. 2003). Once the attachment of the viral particle is complete, the genetic material of the bacteriophage is inserted into the bacterial host cell. Upon penetration, the bacterial metabolic machinery is utilized by the phage to create multiple copies of its own genetic material (DNA or RNA). DNA viruses directly transcribe themselves into mRNA (messenger RNA) molecules that are then used to direct the host cell's ribosomes. In case of RNA viruses (retroviruses), a unique enzyme—reverse transcriptase—transcribes the viral RNA into DNA, and thereafter, follows the path of DNA virus for transcription of the viral RNA. Towards the later stages of this translation, the newly translated proteins are assembled to form the capsid and the tail of the phages that break out of the host cell, bursting its cellular membrane. Each newly formed particle continues to infect new host cells and subsequently proliferates. In few cases, instead of the phage genome, host chromosome gets packed in to the capsid during phage replication and represents an example of horizontal gene transfer within the bacterial population via transduction (Madigan and Martinko 2006). A direct application of phages exhibiting only the lytic cycle is that they can be conveniently employed for tackling the problem of antibiotic-resistant pathogenic bacteria.

Lysogenic cycle

In comparison to the lytic cycle, the lysogenic phase is exhibited particularly by temperate phages (Campbell and Reece 2005), and results in the integration of the viral genetic material with the bacterial genome (called prophage), ensuring continued replication of the viral genetic

Fig. 2 Schematic representation of lytic and lysogenic cycle of phage



material without any fatal consequences to the infected host (Inal 2003). However, due to incorporation of viral genetic material into the host, change in the phenotype of the infected bacteria is a common phenomenon. This conversion may induce the pathogenicity of bacteria, which is evident for many common bacterial strains (Brussow et al. 2004; Keen 2012). Prevention of such lysogenic conversion may also be done using hydrogen peroxide by production of a reactive oxygen species, glutathione, and overexpression of transcriptional repressors (Wagner et al. 2001; Liu et al. 2005; Ptashne 2006; Keen 2012).

Application of phages

Biotechnological relevance

Recently, phages have become an important instrument to biotechnologists. It is being studied for various purposes like drug designing, synthesis of novel protein, delivery of protein and DNA vaccines, detection of pathogenic bacteria, and screening of protein libraries, peptides, or antibodies (Hart et al. 1994; Sperinde et al. 2001; Clark and March 2004; Donnelly et al. 2015; Gao et al. 2015). Further, phages are being used as a substitute to antibiotics for many antibiotic-resistant bacterial strains, biocontrol agents in agriculture, aquaculture, and oil and petroleum industry (Haq et al. 2012). Researchers are also working upon synthesizing a novel peptide by exploiting phage display techniques, gene inclusion, and replication using host machinery (Smith 1985; Sidhu 2000; Haq et al. 2012; Wang et al. 2016). Phage display was first described by Smith (1985) to identify polypeptides with precise bait-binding activity; the technique has evolved with several versatile applications (Paschke 2006; Li and Caberoy 2010). The displayed molecule encoded by the phage genetic information will be expressed on its surface along with the coat protein. Researchers reported using phages like M13, Lambda, and T7 for phage display techniques (Benhar 2001; Willats 2002; Wang et al. 2016). A range of ligands can be prepared by fusing the coat protein gene with different gene-encoding peptides. Such ligands must have capabilities like detecting viable bacteria like *Escherichia coli* (Wang et al. 2016) and targeting specific pathogen proteins, recognizing specific receptors, and blocking interaction between ligand and receptor (Watson and Eveland 1965; Kodikara et al. 1991; Funatsu et al. 2002; Haq et al. 2012). Furthermore, bacteriophages are being modified as tunable nanocontainers for the packaging and delivering of peptides (Kelly et al. 2015). In an interesting study, Dickerson et al. (2005) demonstrated the use of engineered filamentous bacteriophage in cocaine sequestration for blocking the behavioral effects of cocaine in rodent model. Furthermore, phage technology for detecting affinity of antibody to a particular antigen

(pathogenic) or phage amplification for detecting pathogenic bacteria, like *Mycobacterium tuberculosis*, *E. coli*, *Pseudomonas*, etc., improvised research in this field (Stewart et al. 1989; Barry et al. 1996; Donnelly et al. 2015). Exploring phage sensitivity and specificity to design phage-based biosensors can be an improved alternative to antibody-based immunoassay techniques (Peltomaa et al. 2015).

Medicine and clinical application

About a century ago, in 1917, the bacteriophage era was started when Félix d'Herelle published a paper demonstrating “un bactériophage obligatoire” (d'Herelle 1917, 1949; Abedon et al. 2011). Bruynoghe and Maisin (1921) initiated bacteriophage therapy by treating patients having staphylococcal infections. “Phage therapy” has become a potential therapeutic option (Duckworth and Gulig 2002; Wright et al. 2009a, b; Sarker et al. 2012). Immense research has been initiated to employ phage therapy as an alternative (Matsuzaki et al. 2003; Quintin et al. 2005; Chatterjee et al. 2015; Cui 2015; Zschach et al. 2015; Abedon 2016; Maszewska et al. 2016). Phage based products were first developed in the commercial laboratory of d'Herelle in Paris. Treatment of various pathogens and antibiotic-resistance bacteria (like *Salmonella* spp. *Clostridium difficile*, and diarrheagenic *E. coli*) has been tried using phage (Mai et al. 2010; Frampton et al. 2012). Phage-based products were first developed in the commercial laboratory of d'Herelle in Paris. The company (later became the successful French company L'Oréal) produced five phage preparations (Bacté-coli-phage, Bacté-rhinophage, Bacté-intesti-phage, Bacté-pyophage, and Bacté-staphyphage) (Summers 1999; Sulakvelidze et al. 2001). In 1940s, Eli Lilly and Company (Indianapolis, IN, USA) developed seven phage products for human uses. These products were either in the form of a lysate in broth cultures (e.g., Colo-lysate, Ento-lysate, Neiso-lysate, and Staphylo-lysate) or in the form of a water soluble jelly (e.g., Colo-jel, Ento-jel, and Staphylo-jel) against targeted bacteria like, staphylococci, streptococci, and *Escherichia coli*. They were used for treating abscesses, septic wounds and vaginitis, mastoid infections, and respiratory tract infections (Sulakvelidze et al. 2001).

However, due to exponential growth of antibiotic-based drug companies and pharmaceutical giant companies, phage products became less popular (Alisky et al. 1998; Sulakvelidze and Kutter 2005). In favor of therapeutic uses of phages, recent reports on the application of phage cocktails for open septic wounds and burn injuries have contributed encouraging results. Phage cocktail containing 82 phages against *Pseudomonas aeruginosa* and 8 phages against *Staphylococcus aureus* was successfully applied on eight patients (Merabishvili et al. 2009; Wright et al., 2009a, b; Nilsson 2014). Recently, Pherecydes Pharma, a biotechnology company in France, have announced the phase I/II single-blind multicenter clinical trials of its

bacteriophage based product ‘Phagoburn’ (<http://www.pherecydes-pharma.com/phagoburn-clinical-study.html>; Reardon 2014; Ravat et al. 2015; Servick 2016). Phagoburn contains bacteriophage cocktail targeting pathogenic strain of *Escherichia coli* and *Pseudomonas aeruginosa* and its infection in serious burn patients. The Phagoburn is a collaborative European project (www.phagoburn.eu) coordinated by French Ministry of Defense and being conducted in eleven major burns units in France, Switzerland and Belgium along with eight civilian hospitals. Phage therapy is important alternative to overcome critical limitations of antibiotic therapy due to emergence of bacterial resistance, like, in the case of *Clostridium difficile* infection (CDI). CDI is responsible for inducing the dysbiosis, which has extremely high recurrence rates. As per reports, treatment of CDI using current antibiotics has become more and more ineffective (Sangster et al. 2014). Further, prolonged use of antibiotics can also harm beneficial gut flora causing discomfort to the patients. Phage therapy against *C. difficile* involves specifically targeting the causative agent, sparing the other bacterial organisms of the human gut (Sangster et al. 2014, 2015). Similarly, treatment of nosocomial infections (common hospital-acquired infections) caused by *Pseudomonas aeruginosa* is a huge therapeutic challenge currently. High rate of morbidity and mortality is connected with the infection, along with a greater possibility of drug resistance in the bacteria during the course of therapy. With such limited scope for developing new drugs, scientists have also found considerable success in alternative treatment options including phage-based approaches (Viertel et al. 2014; Chatterjee et al. 2015). Babalova et al. (1968) used *Shigella* phages to treat bacterial dysentery and successfully controlled dysentery in patients. Till date, inadequate references are available on human clinical trials conducted and published; the reported cases are phase I studies and mostly from European Medicines Agency (EMA) or United States Food and Drug Administration (FDA) jurisdictions (Bruttin and Brüssow 2005; Merabishvili et al. 2009; Rhoads et al. 2009). In 2009, phase I clinical trial of bacteriophage cocktail (Biophage-PA) targeting three pathogenic strains of *S. aureus*, *P. aeruginosa*, and *E. coli* targeting venous ulcers was approved US FDA (Rhoads et al. 2009). Similarly, a randomized, double-blind, placebo-controlled phase I/II clinical trial approved by UK Medicines and Healthcare products Regulatory Agency (MHRA) and the Central Office for Research Ethics Committees (COREC) ethical review process was performed to treat chronic otitis against multidrug resistant bacteria *P. aeruginosa* and found lower bacterial count and significant improvement with a single input dose of 600,000 bacteriophages (Wright et al. 2009a, b). Bacteriophage application trials have shown that PFU count of 10^2 – 10^3 plaque forming

unit (PFU) is adequate to counter 10^6 – 10^9 CFU per milliliter proliferation threshold of bacteria in vivo (Marza et al. 2006; Wright et al. 2009a, b; Parracho et al. 2012). However, the reported trials with bacteriophages have been performed with 10^5 and 10^9 PFU of individual bacteriophages (Bruttin and Brüssow 2005; Merabishvili et al. 2009; Rhoads et al. 2009; Wright et al. 2009a, b). Development of *phage bioderm* is another important area of application in clinical sector. It is a therapeutic autodegrading, non-toxic, biopolymer complex containing phages that help in healing of wounds and burns, osteomyelitis, and periodontal diseases (Mathur et al. 2003). Studies reported 20–95 % lysis of clinical isolates of *Serratia marcescens* using specific phage strains and phage type (Alisky et al. 1998). However, there are certain limitations on the application of phage bioderm, like reduction in phage activity due to development of antibodies against phages, induction of toxin genes, and fast discharge of bacterial endotoxins due to the lytic effect of phage (Mathur et al. 2003).

Biosensor development

IUPAC defines a biosensor as “a device that uses specific biochemical reactions mediated by isolated enzymes, immune-systems, tissues, organelles or whole cells to detect chemical compounds usually by electrical, thermal or optical signals” (Nagel et al. 1992; Hwang 2014). A biosensor is typically composed of bio-based recognition and transducer components, and electronic systems (signal amplification, processing, and display). Potential applications of biosensors are diverse and numerous, from defense security to pharmaceutical science to environmental monitoring and assessment (Kirsch et al. 2013). Biosensors have a number of advantages like sensitivity, specificity, speed and accuracy of detection, easy sample preparation, cost-effectiveness, etc. (Singh et al. 2013; Hwang 2014; Rackus et al. 2015). Like that of many other biomaterials, unique biological, geometrical, and mechanical characters of bacteriophages can be exploited for bacterial identification, pathogen detection, and biocontrol (Hwang 2014).

It is simpler to develop biosensor surfaces by surface adsorption of phages, though it may give inconsistent results due to unstable immobilization densities (Balasubramanian et al. 2007; Lakshmanan et al. 2007). However, chemically anchoring (cysteamine-modified and glutaraldehyde-activated gold substrate) phages on a detection platform display a consistent improvement in the phage density and detection (Cademartiri et al. 2010; Singh et al. 2013). For chemically functionalizing phage-based biosensors, selection (biopanning) purity of the phage suspension is an important criterion which should be devoid of various other bio-contaminating agents like other carbohydrates, proteins, and lipids (Naidoo et al. 2012). However, genetically modified or engineered phages are more appropriate to develop bio-probes than intact wild-type phages. Being biologically active, wild-type phages upon infection lyse

the host bacterium that may lead to reduction of signal on a biosensor platform (Singh et al. 2010, 2013).

Recently, utilizing unique ability of phages to display peptides or proteins on their surface, called “phage display,” is becoming a powerful tool to screen diversity of targets like proteins, carbohydrates, small molecules, or an entire cell. For the phage display technology, lambda, f1, M13, fd, T4, and T7 phages have most widely been used (Smith 1985; Atias et al. 2008; Singh et al. 2013). This emerging technology can revolutionize diagnostics by creating molecules that are otherwise unavailable via conventional approaches. Researchers have successfully expressed cellular proteins and peptides, antibody (or its fragments), and antigen molecule on the surface of phages for developing pathogen detection biosensors, molecular imaging, and gene delivery (Petrenko 2008; Pande et al. 2010; Singh et al. 2013; Hwang 2014; Chuang et al. 2015).

Agriculture and food safety

Phage application in agriculture and food material is an upcoming area of development. *Salmonella*, *Campylobacter*, *Listeria*, and *E. coli* are common bacterial contaminants in food borne-related infections (DuPont 2007). Bacterial infection in crops is a grave problem that reduces the yields (Allen et al. 2011). High resistivity to multiple antibiotics was reported by a number of researchers in the agricultural field (Price et al. 2012; Zhu et al. 2013; Micallef et al. 2013; Popowska et al. 2012). To evade infection, a number of reports of phage application on various crops (like tomato, citrus, and onion) are available (Huff et al. 2005; Lang et al. 2007; Balogh et al. 2008; Jones et al. 2014). Phage products like AgriPhage™, Omnilytics are being used at a large scale for protecting crops from a number of bacterial diseases. Similarly, other phage products like LISTEX (for *Listeria* sp.; <http://www.listex.eu/product/>), Listshield™, and Intralytix were approved by the FDA (USA), for treating food products before they are being marketed (Meaden and Koskella 2013). Phages have been exploited to treat colibacillosis and prevent food-borne pathogens (Huff et al. 2005; Sharma et al. 2009). Phage therapy is applicable in restricting bacterial contamination on a variety of food materials like chicken flesh, meat, fruits, and vegetables (Flaherty et al. 2000; Goode et al. 2003; Allwood et al. 2004; Toro et al. 2005; Higgins et al. 2005; Fiorentin et al. 2005; Ravensdale et al. 2007; Sharma et al. 2009; Guenther et al. 2009; Marcó et al. 2012). Further, antimicrobial packaging against the activity of *Listeria monocytogenes* for cantaloupes and ready to eat meat and *E. coli* for alfalfa seeds and sprouts are available in the market (Lone et al. 2016).

Aquaculture industries

Aquaculture industries often undergo economic losses chiefly due to uncontrolled microbial diseases that threaten their

development and sustainability (Almeida et al. 2009, Silva et al. 2014). Fish-based products are becoming more popular throughout the world as a cheap source of protein. For the last three decades, considerable growth of aquaculture and related industries has taken place. On the other hand, pathogenic bacteria like *Flavobacterium psychrophilum*, *Photobacterium damsela*, *Vibrio anguillarum*, *Vibrio vulnificus*, *Aeromonas hydrophila*, and *Aeromonas salmonicida* have become more prominent, causing heavy loss and/or diseases in humans due to consumption of contaminated aquaculture products (Subasinghe et al. 2001; Nakai and Park 2002; Flegel 2006; Saksida et al. 2006; FAO Report 2015). Mostly in marine and estuarine fisheries and occasionally in freshwater fisheries, vibriosis is a common disease that causes significant mortality in fish (up to 100 % mortality in larvae). This is also responsible for disease outbreaks in fisheries units. Bacterial genera *Vibrio* (with its various species like *V. vulnificus*, *V. anguillarum*, *V. parahaemolyticus*, *V. alginolyticus*, and *V. salmonicida*) and *Photobacterium damsela* are the causative agents of vibriosis (Noorlis et al. 2011; Higuera et al. 2013; Martínez-Díaz and Hipólito-Morales 2013; Silva et al. 2014). As antibiotic treatments have exhibited reduced effectiveness against multidrug-resistant bacteria, treatments with phages have proven efficient in case of *V. anguillarum* infection in fish larvae (Silva et al. 2014). However, the success of aquaculture phage therapy depends mainly on two factors: number of phages produced by each host cell after host lysis and on latent period for fresh infection to a new host by new phage particles. Ideally, for phage therapy in aquatic condition, maintaining abundant phage number and short period for fresh infection is required, which is a challenge for the days to come (Abedon et al. 2001; Crothers-Stomps et al. 2010; Silva et al. 2014). Table 2 has summarized the phage applicability and trials performed in recent era of antibiotic resistance bacteria.

Wastewater phages and their application in effluent water decontamination

Common wastewater treatment involves technology based on chemical precipitation, ion exchange, sedimentation, or coagulation-flocculation (Otte and Jacob 2006). These techniques are energy and maintenance intensive and require high-infrastructure facilities. Researchers across the world are trying to harness phage-based techniques in wastewater treatments to improve quality of effluent and sludge either released into the environment or reused (Fu and Wang 2011). This treatment has the potential to manage the problems of environmental wastewater processes like: reduction in pathogenic bacteria, foaming in activated sludge plants, digestibility, and water ability of sludge, and reduction in competition between functionally important bacterial populations and nuisance bacteria (Withey et al. 2005; Mulani et al. 2015). This is a highly

Table 2 Representing information on phage therapy trials and its applications in various fields

Target organism	Disease	Phage used	Phage dose/MOI	Comments/result	Reference
1 <i>Xanthomonas axonopodis</i> pv. <i>Vignaeradiatae</i>	Bacterial leaf spot of mung bean	XMP ⁻¹	6×10^9 PFU/mL	Phage application efficiently provide protection against bacterial infection in mung bean seed	Borah et al. (2000)
2 <i>Pseudomonas plecoglossicida</i>	Bacterial haemorrhagic ascites	Phage PPp-W4, PPpW-3	10^7 PFU/g pellets	Oral administration of phage-impregnated feed successfully reduced infection of <i>Pseudomonas plecoglossicida</i>	Park et al. (2000)
3 <i>Salmonella enteritidis</i>	Contamination of food	<i>Salmonella enteritidis</i> specific phages	10^7 PFU/mL	Phage application on fresh cut honeydew melon reduced the bacterial count	Leverentz et al. (2001)
4 <i>Streptomyces scabiei</i>	Potato scab	AS1	10^9 and 10^{12} PFU/mL	Prior to planting, phage treatment of mother tuber reduced scab lesion coverage	McKenna et al. (2001)
5 Vancomycin-resistant <i>Enterococcus faecium</i> (VRE)	Individuals with compromised immune systems are particularly prone to develop VRE infections	Phage ENB6 and C33	3×10^8 PFU/mL	45 min exposure to phage therapy was sufficient to rescue 100 % diseased mice	Biswas et al. (2002)
6 <i>E. coli</i> , <i>Proteus</i> , <i>Pseudomonas</i> , and <i>Staphylococcus</i>	Ulcers and wounds	Pyophage	10^6 PFU/cm ²	PhagoBioDerm effectively healed wounds and eliminated causative organism of ulcers in 22 patients	Markoishvili et al. (2002)
7 <i>Xanthomonas campestris</i> pv. <i>Vesicatoria</i>	Bacterial spot of tomato	Agriphage	10^{10} PFU/mL	PCF and Casecrete formulation of Agriphage increased phage longevity and decreased the pathogen number in field trials	Balogh et al. (2003)
8 <i>Salmonella enterica</i> serovar Enteritidis and <i>Campylobacter jejuni</i>	Food poisoning	<i>Salmonella enteritidis</i> phage type 4 strain P125589, phage 29C, phage P22, HTint, <i>Campylobacter jejuni</i> phage 12673	MOI:100 to 1000	Bacterial count reduced by 2 log10 within 48 h after phages application (MOI of 100 to 1000)	Goode et al. (2003)
9 <i>Streptococcus pneumoniae</i> 541, serotype 6B	Pneumococcal bacteraemia and death	Phage-coded lysins (enzymatic): Pal amidase and/or Cpl-1 lysozyme	1100 U of Pal amidase or Cpl-1 lysozyme	Phage-coded enzymes successfully protected mice from Bacteraemia after 1 h of phage injection	Jado et al. (2003)
10 <i>Pseudomonas plecoglossicida</i>	Bacterial haemorrhagic ascites	Phage PPp-W4, PPpW-3	10^7 PFU/fish	Mortalities of fish reduced and inhibited bacterial infection	Park and Nakai (2003)
11 <i>Pseudomonas aeruginosa</i>	Septicemia	PF3R, PF3, and Pt1	10^6 – 10^9 PFU/mL	PF3R-treated mice showed higher survival rate with reduced inflammatory response	Hagens et al. (2004)
12 <i>Salmonella enteritidis</i>	Food poisoning	Phages CNPSA 1, CNPSA3 and CNPSA4	10^9 PFU/mL	Significant reduction in <i>Salmonella</i> count after treating with bacteriophage and found effective lytic action on days 3, 6 and 9 of post-treatment.	Fiorentin et al. (2005)
13 Nonmotile, serotype O2 isolate of <i>Escherichia coli</i>	Colibacillosis	<i>Escherichia coli</i> phages SPR02 and DAF6	DAF6, 10^9 PFU/mL; SPR02, 10^8 PFU/mL	Phage therapy successfully recovered <i>E. coli</i> infection in birds after 24 or 48 h	Huff et al. (2005)

Table 2 (continued)

Target organism	Disease	Phage used	Phage dose/MOI	Comments/result	Reference
14 <i>Vibrio harveyi</i>	Mass mortalities of shrimp larvae	<i>Vibrio harveyi</i> specific phage	10 ⁹ PFU/mL	Bacteriophage treatment enhanced larval survival by (80 %) as compared control (25 %)	Vinod et al. (2006)
15 <i>Staphylococcus aureus</i> strain 2698	Wound and soft-tissue infection	LS2a	10 ⁹ PFU/mL	Treatment of wound infection by <i>Staphylococcus aureus</i> phages prevent formation of abscess in rabbit model	Wills et al. (2005)
16 <i>Aeromonas salmonicida</i> HER 1107	Furunculosis	Bacteriophage HER 110	10 ⁹ PFU/mL	In brook trout, bacteriophage application delayed furunculosis onset by 7 days	Imbeault et al. (2006)
17 <i>Aeromonas salmonicida</i> subsp. <i>Salmonicida</i>	Furunculosis of Atlantic salmon	<i>Aeromonas salmonicida</i> phages O, R, and B	10 ⁵ , 10 ⁷ , and 10 ⁸ PFU/fish	Slow down the mortality rate, but at the end 100 % mortality was reported	Verner-Jeffreys et al. (2007)
18 <i>Vibrio harveyi</i>	Mortality of Penaeus monodon larvae	Bacteriophage specific to <i>Vibrio harveyi</i>	10 ⁶ PFU/mL	85 % increase in survival at larval stages of shrimp	Karunasa gar et al. (2007)
19 <i>Xanthomonas axonopodis</i> Pv. Allii	Xanthomonas leaf blight of onion	Agriphage	10 ⁸ , 10 ⁷ , 10 ⁶ , or 10 ⁵ PFU/mL	Biweekly or weekly applications of bacteriophages under green house condition reduced leaf blight disease by 26 to 50 % and proved equally better as weekly applications of copper hydroxide plus mancozeb	Lang et al. (2007)
20 <i>Pseudomonas aeruginosa</i> strain D4	Septicemia in immunocompromised hosts	KPP10	10 ¹⁰ PFU/mL	Oral administration KPP10 significantly reduced gut-derived sepsis caused by <i>P. aeruginosa</i>	Watanabe et al. (2007)
21 <i>Xanthomonas axonopodis</i> subspecies <i>citri</i> and <i>citrumelo</i>	Citrus canker and Citrus bacterial spot	CP2, Φ Xac2005-1, cc Φ 7, cc Φ 13, Φ Xacm2004-4, Xacm2004-16, Φ X44, and Φ XaacA1	10 ⁶ , 10 ⁸ , 10 ⁹ , and 10 ¹⁰ PFU/mL	Treatment of sensitive Valencia oranges with bacteriophage provided significant disease reduction in plants	Balogh et al. (2008)
22 <i>Pseudomonas aeruginosa</i> and <i>Staphylococcus aureus</i>	Burn wound patients	Bacteriophage cocktail PNM, 14/1 and ISP	10 ⁹ PFU/mL	Bacteriophage cocktail found effective in treating burn wound and is being evaluated in a pilot clinical study	Merabishvili et al. 2009
23 <i>Escherichia coli</i> O157:H7	Watery diarrhea, hemorrhagic colitis, hemolytic uremic syndrome, and in some cases death	Bacteriophages e11/2 and e4/1c	10 ¹¹ PFU	cleared by Medical Ethical Committee	Rivas et al. (2010)
24 <i>Ralstonia solanacearum</i>	Bacterial wilt	RSA1, RSBI, and RSL1	10 ⁹ PFU/mL	Bacteriophages e11/2 and e4/1c reduced the number of <i>Escherichia coli</i> O157:H7 in Rumen model	Fujiwara et al. (2011)
25 <i>Klebsiella pneumoniae</i> B5055	Burn wound infection	Phage Kpn5	MOI:200	phage ϕ RSL1 prevents bacterial wilt	Kumari et al. (2011)
26	Columnaris		10 ⁵ –10 ⁹ PFU/mL	Kpn5 phages have therapeutic importance in treating burn wound infection in mice	Laanto et al. (2011)

Table 2 (continued)

Target organism	Disease	Phage used	Phage dose/MOI	Comments/result	Reference
<i>Flavobacterium columnare</i>		Flavobacterium phages (a) FCV-1, (b) FCL-2, (c) Fly-3, (d) FK0-2, and (e) FKy-1		Flavobacterium phages were found more host specific to <i>F. columnare</i> and can be useful in treating Columnaris in fishes	
27 <i>Dickeya solani</i>	Rotting and blackleg of potato	T4-related Bacteriophage (vB_DsoM_LIMEstone1 and vB_DsoM_LIMEstone)	10 ¹⁰ PFU/L	Phage application on infected tubers resulted in a higher yield potato tubers	Adriaenssens et al. (2012)
28 <i>Pseudomonas aeruginosa</i>	Infections of MDR <i>Pseudomonas aeruginosa</i>	Phage PS5	~9 × 10 ⁸ PFU (First dose) ~3 × 10 ⁸ PFU (Second dose)	In murine model treatment with Phage PS5 resolved the infection of MDR <i>Pseudomonas aeruginosa</i>	Golkar et al. (2013)
29 <i>Vibrio anguillarum</i>	Vibriosis	KVP40 and (Phages H1, H2, H4, H5, H7, H8, H20, and 2E-1, S4-7, and S4-18)	MOI: 0.1, 1, and 10	Lytic phages were identified against <i>V. anguillarum</i> hosts to treat Vibriosis	Tan et al. (2014)
30 <i>Escherichia coli</i> (Antibiotic-resistant clinical strain)	Wounds	TPR7	10 ¹⁰ PFU/mL	Phage administration in animal cured from infection after 48 h and results were equally effective as found in the group treated with gentamicin.	Rahmani et al. (2015)
31 <i>E. coli</i> strain DPC6051	<i>E. coli</i> infections	APCEc01, APCEc02, and APCEc03	MOI 10 ⁻³ and 10 ⁵	Phage cocktail reduced biofilm formation and effective in preventing the appearance of phage-resistant mutants	Dalmaso et al. (2016)
32 <i>Pseudomonas aeruginosa</i>	<i>Pseudomonas</i> infections	Phage KTN4	MOI 100	KTN4 phage injection in <i>Galleria</i> larvae model inoculated with <i>P. aeruginosa</i> strains successfully reduced colony count (4–7 log) of <i>P. aeruginosa</i>	Danis-Wlodarczyk et al. (2016)
33 <i>Escherichia coli</i> (ETEC) K88	ETEC infection in post-weaning pigs	Phages against enterotoxigenic <i>Escherichia coli</i> (ETEC) K88	10 ⁷ PFU/kg	Treatment with phages was found effective for reduction of acute ETEC K88 infection in post-weaning pigs	Lee et al. (2016)
34 <i>Escherichia coli</i> and <i>Klebsiella pneumoniae</i> strains	Urinary tract infections	Pyo, Intesti, Ses, Enko bacteriophages, 29 <i>E. coli</i> , and 10 <i>K. pneumoniae</i> bacteriophages	10 ⁷ PFU	Lytic action of commercial bacteriophage cocktails on <i>E. coli</i> and <i>K. pneumoniae</i> strains isolated UTIs patients	Sybesma et al. (2016)

potential area of research and development especially in a country like India; there is potential here for application of appropriate, affordable, sustainable, and eco-friendly approaches. Phage-mediated bactericidal effects in any wastewater treatment process have many controlling factors that lead to treatment performance. Wastewater is rich in microbial diversity that contains a wide variety of microbes and bacteriophages, among which, somatic coliphages are the most abundant. These coliphages are virulent, in nature, generally following lytic cycle for their reproduction. These phages belong to the families Myoviridae, Siphoviridae, Podoviridae, and Microviridae that mainly target the member of *Enterobacteriaceae* family especially *E. coli* (Hayes 1968). Coliphages find favorable environment in wastewater. However, their abundance depends upon factors like optimal condition and density of the host bacteria, pH, temperature, cations like calcium and magnesium, organic matter concentration, etc. (Grabow 1990; Grabow et al. 1980, 1998). For instance, fecal coliform bacteria closely related to *E. coli*, especially *Klebsiella* species, are heterotrophic and favor certain water environments for its multiplication and in turn support the multiplication of somatic coliphages (Grabow et al. 2000). It has been reported that phage titer against *Salmonella typhi* in both laboratory media and sewage was found to be 10^9 – 10^{10} per milliliter (Goyal et al. 1980). Successful phage replication requires a minimum of 10^4 host bacteria per milliliter. According to Goyal et al. (1987), phages under nutrient-limited conditions were found concentrated on the solid surface instead of flowing water. Coliphages also inhabit inside the gut of humans and other warm blooded animals, multiply within it, and are excreted along with fecal matter. Different pieces of information are available on phages in sewage. A number of studies have reported density of somatic coliphages to be 10^6 – 10^8 per milliliter (Bell 1976; Ignazitto et al., 1980; Havelaar and Hogeboom 1984; Havelaar et al. 1984; Tartera et al. 1989; Grabow et al. 1993; Grabow et al., 2000).

Interestingly, some members of coliphages showed similarity with human viruses like enteroviruses such as polio viruses (Family Picornaviridae) and F-RNA coliphages (Family Leviviridae) (Kamiko and Ohgaki 1993). Both viral particles consist of an icosahedral capsids (about 25 nm diameter) and a single strand (ss)-RNA genome. Due to these similarities, coliphages are considered to be valuable models or surrogates for human enteric viruses (Grabow 2001), which are also excreted along with fecal matter (Grabow et al. 1995). Although coliphages are shed regularly along with fecal matter, human enteric viruses are excreted generally during infection. Greater number of adenoviruses than enteroviruses have been constantly present in raw sewage around the world (Irving and Smith 1981; Hurst et al. 1988; Krikelis et al. 1985a, b), and according to Hurst et al. 1988, 80 % of infectious adenoviruses shed in the feces and present in raw sewage may be enteric adenoviruses. Recently, human fecal

contamination was indicated using potential source indicators such as bacteriophages against Bacteroides (GA-17, GB-124, and ARABA 84) as well as sorbitol-fermenting Bifidobacteria (Wicki et al. 2015). Furthermore, according to Haramoto et al. (2015), coliphages belonging to the genogroup, GI F-RNA, may be used as a suitable indicator of virus reduction during wastewater treatment.

Impediments of phage application: interplay between phage and bacterial resistance

There is a tug of war between phage and bacterial infection and immunity. The selective pressure among these two entities is unique. Bacteria develop resistance to phage; subsequently, phages improve their antiviral mechanism and vice versa. This interplay between phage and bacteria is an interesting phenomenon; elevated mutation rates is a normal event for bacteria like *Pseudomonas fluorescens* when they are subjected to coevolve in the invitro systems (Buckling and Rainey 2002). Again, Pal et al. (2007) showed that in laboratory conditions, coevolution of *P. fluorescens* with its bacteriophage help in faster bacterial mutation rates. They have also found that 25 % of the *P. fluorescens* populations (approx. 200 bacterial generations) coevolving with the lytic phages had evolved 10 to 100-fold increase in mutation rates (Pal et al. 2007). For lytic bacteriophages, reciprocal selection for bacterial resistance and phage infectivity is essential. And this bacteria-phage interface generates more beneficial resistance mutations to the bacteria (Gomez and Buckling 2013). However, natural populations of the bacteria-phage coevolving in their natural environment (as for example, soil) may not follow the same mutation pattern as, in nature, several other factors may influence the selection rates for mutation. Innumerable microbial species including natural microbial and viral community (NMC) as a whole and other physico-chemical factors possibly will influence the rate of mutation (Gomez and Buckling 2013).

Bacterial resistance towards phage, therefore, may be due to several mechanisms. These include modification of phage attachment or adsorption sites (receptors) and CRISPR sequence-based adaptive immunity (Hyman and Abedon 2010). Alteration of phage adsorption sites/receptors is the most common phenomenon by which bacteria evade phage infection and become resistant to phage. However, phages can also adapt themselves to recognize these new modified receptors. Synthesis of exopolysaccharide (EPS) or masking proteins (like protein A of *S. aureus*) to mask the phage receptor is another strategy of bacteria. Again, phages conquer the barrier by cleaving the EPS layer using polysaccharide lyase or a polysaccharide hydrolase (Labrie et al. 2010, Örmälä and Jalasvuori 2013).

Further, self/non-self discrimination in prokaryotes is ubiquitous and provided by restriction-modification (RM) systems that defend hosts from exogenous DNA (Pleška et al. 2016). Bacterial system has the ability to recognize and modify the phage DNA. In many instances, the phage DNA is cleaved by the restriction endonucleases on entering into the bacterial cell through RM system and protects bacterial cell from foreign phage DNA attack (Pleška et al. 2016).

It has also been reported that phages adopt anti-restriction strategy to avoid recognition by endonuclease enzyme. For example, T4 phage escapes of restriction endonuclease attack as it contains hydroxymethylcytosine (HMC) instead of cytosine. However, some bacteria, again, modified their system to specially recognize hydroxymethylcytosine (HMC) and shatter phage DNA (Bickle and Kruger 1993; Borgaro and Zhu 2013). Yet again, anti-restriction strategy in *Staphylococcus* phage *K* is interesting where no 5' GATC-3' cleavage site is present; hence, DNA is protected from restriction (Kruger et al. 1988; Tock and Dryden 2005; Bryson et al. 2015).

CRISPER-Cas mechanism or Clustered regularly interspaced short palindromic repeats (CRISPRs) and the CRISPR-associated (cas) genes present a novel example of acquired resistance against viruses in prokaryotes. This mechanism is widely present in genome of bacteria and archaeal population and CRISPER-Cas loci were first described in 1987 in *E. coli* (Ishino et al. 1987). CRISPER is composed of 21–48 bp direct repeats separated by (26–72 bp) non-repetitive spacers, and at 5' edge, this loci is flanked by a number of *cas* genes (4–20 in number) in bacterial strains. The variation in *cas* gene and spacer make this system more unique and robust (Marraffini 2015). Main function of CRISPER-Cas is to provide immunity against foreign DNA including phage genomic DNA or plasmid DNA (Makarova et al. 2006; Bolotin et al. 2005; Haft et al. 2005; Mojica et al. 2005; Pourcel et al. 2005; Marraffini and Sontheimer 2008). Further, as for example of antiphage activity of CRISPER-Cas mechanism in *Streptococcus thermophilus*, exposure to virulent phage gave rise to the phage-resistant mutants due to insertion of additional 30 bp spacer similar to protospacer of infecting phage (Barrangou et al. 2007). The event of acquiring immunity against phage can be explained in following steps like adaptation or spacer attainment, transcription of acquired spacer (small CRISPER RNAs (crRNAs), on recurrent phage attack this crRNAs form a complex with Cas protein), and immunity against phage (crRNAs-Cas complex direct nuclease to trace and chop the invading phage DNA; Marraffini 2015).

Conclusive remarks

Phages are omni-present, ubiquitous, and important part of nature. Being antibacterial and natural predators of bacteria, the role of phage can be extensive as nano-cleaner of

ecosystem. Detailed classification of bacteriophages may be beneficial to mankind in the screening of therapeutic phages against a number of diseases and economically important phages help in industries and its preservation. High-throughput sequencing like metagenomics study of environmental samples can provide information of novel bacteriophages or new bacteriophage species. Various biological, environmental, medical, and pharmaceutical applications related to the use of phage are becoming more and more attractive. Further, employability of phage for sustainable wastewater treatment is a challenging approach. Labeled phages are used in situ bacterial detection using phage typing of clinical bacterial strains. Although discovered a century ago, their use in various fields are still restrictive. Comprehensive studies on basic molecular biology, host range increment, and genetics processes of these tiny antibacterial agents are required which will help them consider novel beneficial agent for bactericidal activities, even against multidrug-resistant bacteria.

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References

- Abedon ST (2016) Bacteriophage exploitation of bacterial biofilms: phage preference for less mature targets? FEMS Microbiol Lett. doi:10.1093/femsle/fnv246
- Abedon ST, Herschler TD, Stopar D (2001) Bacteriophage latent-period evolution as a response to resource availability. Appl Environ Microbiol 13:4233–4241
- Abedon ST, Sarah JK, Bob GB, Kutter EM (2011) Phage treatment of human infections. Bacteriophage. 1(2):66–85
- Abeles AL, Snyder KM, Chatteraj DK (1984) P1 plasmid replication: replicon structure. J Mol Biol 17:307–324
- Ackermann HW (2011a) Bacteriophage taxonomy Microbiology Australia, 90–94. Downloaded from <http://journals.cambridge.org/11%20ackermann244.pdf>
- Adriaenssens EM, Vaerenbergh JV, Vandenheuvel D (2012) T4-related bacteriophage LIMESTONE isolates for the control of soft rot on potato caused by *Dickeya solani*. PLoS One 7:e33227
- Alisky J, Iczkowski K, Rapoport A, Troitsky N (1998) Bacteriophages show promise as antimicrobial agents. J Infect 36:5–15
- Allen HK, Looft T, Bayles DO, Humphrey S, Levine UY, Alt D, Stanton TB (2011) Antibiotics in feed induce prophages in swine fecal microbiomes. MBio 2:00260–00211
- Allwood PB, Malik YS, Maherchandani S, Vought K, Johnson LA, Braymen C, Hedberg CW, Goyal SM (2004) Occurrence of *Escherichia coli*, noroviruses, and f-specific coliphages in fresh market-ready produce. J Food Protect 67:2387–2390

- Almeida A, Cunha A, Gomes NCM, Alves E, Costa L et al (2009) Phage therapy and photodynamic therapy: low environmental impact approaches to inactivate microorganisms in fish farming plants. *Mar Drugs* 7:268–313
- Atias D, Lobel L, Virta M, Marks RS (2008) Phage-displayed epitopes as bioreceptors for biosensors part two. Biological and molecular recognition systems. Wiley, Handbook of Biosensors and Biochips. doi:10.1002/9780470061565.hbb012
- Babalova EG, Katsitadze KT, Sakvarelidze LA, Imnaishvili NS, Sharashidze TG, Badashvili VA, Kiknadze GP, Meipariani AN, Gendzekhadze ND, Machavariani EV, Gogoberidze KL, Gozalov EI, Dekanosidze NG (1968) Preventive value of dried dysentery bacteriophages. *Zh Mikrobiol Epidemiol Immunobiol* 45:143–145
- Balasubramanian S, Sorokulova IB, Vodyanov VJ, Simonian AL (2007) Lytic phage as a specific and selective probe for detection of *Staphylococcus aureus*—a surface plasmon resonance spectroscopic study. *Biosens. Bioelectron.* 22:948–955
- Balogh B, Canteros BI, Stall RE, Jones JB (2008) Control of citrus canker and citrus bacterial spot with bacteriophages. *Plant Dis* 92:1048–1052
- Balogh B, Jones JB, Momolet MT (2003) Improved efficacy of newly formulated bacteriophages for management of bacterial spot on tomato. *Plant Dis* 87:949–954
- Barrangou R, Fremaux C, Deveau H, Richards M, Boyaval P, Moineau S, Romero DA, Horvath P (2007) CRISPR provides acquired resistance against viruses in prokaryotes. *Science* 315:1709–1712
- Barry MA, Dower WJ, Johnston SA (1996) Toward cell-targeting gene therapy vectors: selection of cell-binding peptides from random peptide presenting phage libraries. *Nat Med* 2:299–305
- Bell RG (1976) The limitation of the ratio of fecal coliform to total coliphage as a water pollution index. *Water Res* 10:745–748
- Benhar I (2001) Biotechnological applications of phage and cell display. *Biotechnol Adv* 19:1–33
- Bickle TA, Kruger DH (1993) Biology of DNA restriction. *Microbiol Rev* 57:434–450
- Biswas B, Adhya S, Washart P, Paul B, Trostel AN, Powell B, Carlton R, Merrill CR (2002) Bacteriophage therapy rescues mice bacteremic from a clinical isolate of vancomycin-resistant *Enterococcus faecium*. *Infect Immun* 70:204–210
- Bolotin A, Quinquis B, Sorokin A, Ehrlich SD (2005) Clustered regularly interspaced short palindrome repeats (CRISPRs) have spacers of extrachromosomal origin. *Microbiology* 151:2551–2561
- Borgaro JG, Zhu Z (2013) Characterization of the 5-hydroxymethylcytosine-specific DNA restriction endonucleases. *Nucleic Acids Res* 41:4198–4206
- Bradley DE (1967) Ultrastructure of bacteriophages and bacteriocins. *Bacteriol Rev* 31:230–314
- Bradley DE, Kay D (1960) The fine structure of Bacteriophages. *J Gen Microbiol* 23:553–563
- Breitbart M (2012) Marine viruses: truth or dare. *Annu Rev Mar Sci* 4:425–448
- Breitbart M, Wegley L, Leeds S, Schoenfeld T, Rohwer F (2004) Phage community dynamics in hot springs. *Appl Environ Microbiol* 70:1633–1640
- Brenner S, Horne RW (1959) A negative staining method for high-resolution electron microscopy of viruses. *Biochim Biophys Acta* 34:103–110. doi:10.1016/0006-3002(59)90237-9
- Brenner S, Streisinger G, Horne RW, Champe SP, Barnett L, Benzer S, Rees MW (1959) Structural components of bacteriophage. *J Mol Biol* 1:281
- Brock TD, Madigan, Michael T (1988) Biology of microorganisms, 5th edn. Prentice-Hall, USA, pp. 1988–1835
- Brown S, Maria JPS, Walker S (2013) Wall teichoic acids of gram-positive bacteria. In: S. Gottesman (ed) Annual Review of Microbiology, 67 (Palo Alto: Annual Reviews), pp 313–336
- Brussow H, Canchaya C, Hardt W (2004) Phages and the evolution of bacterial pathogens: from genomic rearrangements to lysogenic conversion. *Microbiol Mol Biol Rev* 68:560–602
- Bruttin A, Brüssow H (2005) Human volunteers receiving *Escherichia coli* phage T4 orally: a safety test of phage therapy. *Antimicrob Agents Chemother* 49:2874–2878
- Bruynoghe R, Maisin J (1921) Essais de thérapie au moyen du bacteriophage. *J Soc Biol* 85:1120–1121
- Bryson AL, Hwang Y, Sherrill-Mix S, Wu GD, Lewis JD, Black L, Clark TA, Bushman FD (2015) Covalent modification of bacteriophage T4 DNA inhibits CRISPR-Cas9. *MBio* 6:e00648. doi:10.1128/mBio.00648-15
- Buckling A, Rainey PB (2002) Antagonistic coevolution between a bacterium and a bacteriophage. *P Roy Soc B Biol Sci* 269:931–936
- Bugla-Ploskonska G, Futoma-Koloch B, Doroszkiewicz W (2007) Role of outer membrane proteins of gram-negative bacteria in interaction with human organism. *Postepy Mikrobiol* 46:139–152
- Cademartiri R, Anany H, Gross I, Bhayani R, Griffiths M, Brook MA (2010) Immobilization of bacteriophages on modified silica particles. *Biomaterials* 31:1904–1910
- Campbell NA, Reece JB (2005) Biology. Pearson, Benjamin Cummings, San Francisco, pp. 338–339
- Chatterjee M, Anju CP, Biswas L, Anil Kumar V, Gopi Mohan C, Biswas R (2015) Antibiotic resistance in *Pseudomonas aeruginosa* and alternative therapeutic options. *Int J Med Microbiol* doi:10.1016/j.ijmm.2015.11.004
- Chuang CH, Wu TF, Chen CH, Chang KC, Ju JW, Huang YW, Van Nhan V (2015) Lab on a chip for multiplexed immunoassays to detect bladder cancer using multifunctional dielectrophoretic manipulations. *Lab Chip* 15:3056–3064
- Clark JR, March JB (2004) Bacterial viruses as human vaccines? *Expert Rev Vaccines* 3:463–476
- Crothers-Stomps C, Høj L, Bourne DG, Hall MR, Owens L (2010) Isolation of lytic bacteriophages against *Vibrio harveyi*. *J Appl Microbiol* 108:1744–1750
- Cui Z (2015) Advances in the treatment of wound bacterial infection with phage. *Zhonghua Shao Shang ZaZhi* 31:389–391
- D’Herelle F (1917a) Sur un microbe invisible antagoniste des bacilles dysentériques. *C R Academy of Sciences (Paris)* 165:373–375
- D’Herelle F (1930) The bacteriophage and its clinical applications. Charles C Thomas, Springfield, Ill
- Dalmaso M, Strain R, Neve H, Franz CMAP, Cousin FJ, Ross RP (2016) Three new *Escherichia coli* phages from the human gut show promising potential for phage therapy. *PLoS One* 11(6):e0156773. doi:10.1371/journal.pone.0156773
- d’Herelle F (1917b) On an invisible microbe antagonistic to dysentery bacilli. *Comptes Rendus Academie des Sciences* 165:373–375
- d’Herelle F (1949) The bacteriophage. *Science News* 14:44–59
- Dickerson TJ, Kaufmann GF, Janda KD (2005) Bacteriophage-mediated protein delivery into the central nervous system and its application in immune pharmacotherapy. *Expert Opin Biol Ther* 5:773–781
- Donnelly A, Yata T, Bentayebi K, Suwan K, Hajitou A (2015) Bacteriophage mediates efficient Gene transfer in combination with conventional Transfection reagents. *Viruses* 7:6476–6489
- Duckworth DH (1976) Who discovered Bacteriophage? *Bacteriol Rev* 40:793–802
- Duckworth DH, Gulig PA (2002) Bacteriophages: potential treatment for bacterial infections. *BioDrugs* 16:57–62
- DuPont HL (2007) The growing threat of foodborne bacterial enteropathogens of animal origin. *Clin Infect Dis* 45:1353–1361
- FAO Report (2015) Downloaded from: <http://www.fao.org/3/a-i4910e.pdf>. Accessed 09 Jan 2016
- Fiorentin L, Vieira ND, Bavioni JW, Aves ES (2005) Use of lytic bacteriophages to reduce salmonella enteritidis in experimentally contaminated chicken cuts. *Braz J Poult Sci* 7:255–260

- Flaherty JE, Jones JB, Harbaugh BK, Somodi GC, Jackson LE (2000) Control of bacterial spot on tomato in the greenhouse and field with h-mutant bacteriophages. *Hort Sci* 35:882–884
- Flegel TW (2006) Detection of major penaeid shrimp viruses in Asia, a historical perspective with emphasis on Thailand. *Aquaculture* 258: 1–33
- Frampton RA, Pitman AR, Fineran PC (2012) Advances in Bacteriophage-mediated control of plant pathogens. *Int J Microbiol Res*. doi:10.1155/2012/326452 Article ID 326452, 11 pages
- Fu F, Wang Q (2011) Removal of heavy metal ions from wastewaters: a review. *J. Environ Manag* 92:407–418. doi:10.1016/j.jenvman.2010.11.011
- Fujiwara A, Fujisawa M, Hamasaki R, Kawasaki T, Fujie M, Yamada T (2011) Biocontrol of *Ralstonia solanacearum* by treatment with lytic bacteriophages. *Appl Env Microbiol* 77:4155–4162
- Funatsu T, Taniyama T, Tajima T, Tadakuma H, Namiki H (2002) Rapid and sensitive detection method of a bacterium by using a GFP reporter phage. *Microbiol Immun* 46:365–369
- Gao C, Hong M, Geng J, Zhou H, Dong J (2015) Characterization of PI (breast cancer cell special peptide) in MDA-MB-231 breast cancer cells and its potential therapeutic applications. *Int J Oncol* 47:1371–1378
- Glud RN, Middleboe M (2004) Virus and bacteria dynamics of coastal sediment: implication for benthic carbon cycling. *Limnol Oceanogr* 49:2073–2081
- Golkar Z, Bagasra O, Jamil N (2013) Experimental phage therapy on multiple drug resistant *Pseudomonas Aeruginosa* infection in mice. *J Antivir Antiretrovir*:S10–005. doi:10.4172/jaa.S10-005
- Gomez P, Buckling A (2013) Coevolution with phages does not influence the evolution of bacterial mutation rates in soil. *ISME J* 7:2242–2244
- Goode D, Allen VM, Barrow PA (2003) Reduction of experimental salmonella and campylobacter contamination of chicken skin by application of lytic bacteriophages. *Appl Environ Microbiol* 69:5032–5036
- Goyal SM, Gerba CP, Bitton G (1987) Phage ecology. Wiley, New York, p. 321
- Goyal SM, Zerda KS, Gerba CP (1980) Concentration of coliphages from large volumes of water and wastewater. *Appl Environ Microbiol* 39: 85–91
- Grabow WOK (2001) Bacteriophages: update on application as models for viruses in water. *Water SA* 27:251–268
- Grabow WOK (1990) Microbiology of drinking water treatment: reclaimed wastewater. In: McFeters GA (ed) *Drinking water microbiology-progress and recent developments*. Springer Verlag, New York, pp. 185–203
- Grabow WOK, Burger JS, Nupen EM (1980) Evaluation of acid-fast bacteria, *Candida albicans*, enteric viruses and conventional indicators for monitoring wastewater reclamation systems. *Prog Water Technol* 12:803–817
- Grabow WOK, Holtzhausen CS and De Villiers JC (1993) Research on Bacteriophages as Indicators of Water Quality. WRC Report No 321/1/93 Water Res Comm Pretoria 147
- Grabow WOK, Neubrech TE, Holtzhausen CS, Jofre J (1995) *Bacteroides fragilis* And *Escherichia coli* bacteriophages: excretion by humans and animals. *Water Sci Technol* 31:223–230
- Grabow WOK, Taylor MB, Clay CG and De Villiers JC (2000) Molecular detection of viruses in drinking water: Implications for safety and disinfection. Proc. 2nd Conf. of the Int. Life Sciences Inst.: The Safety of Water Disinfection: Balancing Chem. and Microb. Risks. Radisson Deauville Resort, Miami Beach, Florida, USA, pp 15–17
- Grabow WOK, Vrey A, Uys M and De Villiers JC (1998) Evaluation of the application of bacteriophages as indicators of water quality. WRC Report No 540/1/98. Water Res. Commission, Pretoria, p 55
- Guenther S, Huwyler D, Richard S, Loessner MJ (2009) Virulent bacteriophage for efficient biocontrol of listeria *Listeria monocytogenes* in ready-to-eat foods. *Appl Environ Microbiol* 75:93–100
- Haft DH, Selengut J, Mongodin EF, Nelson KE (2005) A guild of 45 CRISPR-associated (Cas) protein families and multiple CRISPR/Cas subtypes exist in prokaryotic genomes. *PLoS Comput Biol* 1: 474–483
- Hagens S, Habe A, Ahsen UV, Gabain A, Blasi U (2004) Therapy of experimental pseudomonas infections with a non replicating genetically modified phage. *Antimicrob. Agents Chemother* 48:3817–3822. doi:10.1128/AAC0066-4804/04/\$08.000
- Haguenau F, Hawkes PW, Hutchison JL, Satiat-Jeuñemaître B, Simon GT, Williams DB (2003) Key events in the history of the electron microscope. *Microsc Microanal* 9:96–138. doi:10.1017/S1431927603030113
- Hankin EHL (1896) Action bactericide des Eaux de la Jumna et du Gange sur le vibron du cholera. *Annales de l'Institut Pasteur* 10:511
- Hanlon GW (2007) Bacteriophages: an appraisal of their role in the treatment of bacterial infections. *Int J Antimicrob Agents* 30:118–128
- Ackermann HW (2011b) The first phage electron micrographs. *Bacteriophage*. 1(4):225–227
- Haq IU, Chaudhry WN, Akhtar MN, Andleeb S, Qadri I (2012) Bacteriophages and their implications on future biotechnology: a review. *Virol J* 9:1–8
- Haramoto E, Fujino S, Otagiri M (2015) Distinct behaviors of infectious F-specific RNA coliphage genogroups at a wastewater treatment plant. *Sci Total Environ* 1:32–38
- Hart SL, Knight AM, Harbottle RP, Mistry A, Hunger HD, Cutler DF, Williamson R, Coutelle C (1994) Cell binding and internalization by filamentous phage; displaying a cyclic Arg-Gly-asp-containing peptide. *The J Biol Chem* 269:12468–12474
- Havelaar AH, Hogeboom WM (1984) A method for the enumeration of male-specific bacteriophages in sewage. *J Appl Bacteriol* 56:439–447
- Havelaar AH, Hogeboom WM, Pot R (1984) F-specific RNA bacteriophages in sewage; methodology and occurrence. *Water Sci Technol* 17:645–655
- Hayes W (1968) The genetics of bacteria and their viruses, 2nd edn. Blackwell Scientific Publications, Oxford
- Hendrix RW (2002) Bacteriophages: evolution of the majority. *Theor Popul Biol* 61:471–480
- Higgins J, Higgins S, Guenther KL, Huff W, Donoghue MA, Donoghue DJ, Hargis BM (2005) Use of specific bacteriophage treatment to reduce salmonella in poultry products. *Poult Sci* 84:1141–1145
- Higuera G, Bastías R, Tsertsvadze G, Romero J, Espejo RT (2013) Recently discovered *Vibrio anguillarum* phages can protect against experimentally induced vibriosis in Atlantic salmon, *Salmo salar*. *Aquaculture* 392–395:128–133
- Huff WE, Huff GR, Rath NC, Balog JM, Donoghue AM (2005) Alternatives to antibiotics: utilization of bacteriophage to treat colibacillosis and prevent foodborne pathogens. *Poult Sci* 84: 655–659
- Hurst CJ, McClellan KA, Benton WH (1988) Comparison of cytopathogenicity, immunofluorescence and in situ DNA hybridization as methods for the detection of adenoviruses. *Water Res* 22:1547–1552
- Hwang I (2014) Virus outbreaks in chemical and biological sensors. *Sensors* 14(8):13592–13612. doi:10.3390/s140813592
- Hyman P, Abedon ST (2010) Bacteriophage host range and bacterial resistance. *Adv Appl Microbiol* 70:217–248
- Ignazitto G, Volterra L, Aulicino FA, D'angelo AM (1980) Coliphages as indicators in a treatment plant. *Water Air Soil Pollut* 13:391–398
- Imbeault S, Parent S, Lagacé M, Uhland CF, Blais JF (2006) Using bacteriophages to prevent furunculosis caused by *Aeromonas salmonicida* in farmed brook trout. *J Aquat Anim Health* 18: 203–214

- Inal JM (2003) Phage therapy: a reappraisal of Bacteriophages as antibiotics. *Archivum Immunologiae et Therapiae Experimentalis* 53:237–244
- Irving LG, Smith FA (1981) One-year survey of enteroviruses, adenoviruses, and reoviruses isolated from effluent at an activated-sludge purification plant. *Appl Environ Microbiol* 41:51–59
- Ishino Y, Shinagawa H, Makino K, Amemura M, Nakata A (1987) Nucleotide sequence of the *iap* gene, responsible for alkaline phosphatase isozyme conversion in *Escherichia coli*, and identification of the gene product. *J Bacteriol* 169:5429–5433
- Jado I, Lopez R, Garcia E, Fenoll A, Casal J, Garcia P (2003) Phage lytic enzymes as therapy for antibiotic-resistant *Streptococcus pneumoniae* infection in a murine sepsis model. *J Antimicrob Chemother* 52:967–973
- Jamalludeen N, She YM, Lingohr EJ, Griffiths M (2009) Isolation and characterization of virulent bacteriophages against *Escherichia coli* serogroups O1, O2, and O78. *Poult Sci* 88:1694–1702
- Jones JB, Vallad GE, Iriarte FB, Obradović A, Wernsing MH, Jackson LE, Balogh B, Hong JC, Momo MT (2014) Considerations for using bacteriophages for plant disease control. *Bacteriophage* 2:208–214
- Kamiko N, Ohgaki S (1993) Multiplication characteristics of FRNA phage and its utility as an indicator for pathogenic viruses. *Water Sci Technol* 27:133–136
- Karlsson F, Borrebaeck C, Nilsson N, Malmborg-Hager AC (2003) The mechanism of bacterial infection by filamentous phages involves molecular interactions between TolA and phage protein 3 domains. *J Bacteriol* 8:2628–2634
- Karunasagar I, Shivu MM, Girisha SK, Krohne G, Karunasagar I (2007) Biocontrol of pathogens in shrimp hatcheries using bacteriophages. *Aquaculture* 268:288–292
- Keen EC (2012) Paradigms of pathogenesis: targeting the mobile genetic elements of disease. *Front Cell Infect Microbiol* 2:1–3
- Kelly P, Anand P, Uvaydov A, Chakravartula S, Sherpa C, Pires E, O'Neil A, Douglas T, Holford M (2015) Developing a dissociative Nanocontainer for peptide drug delivery. *Int J Environ Res Public Health* 12:12543–12555
- Kirsch J, Siltanen C, Zhou Q, Revzin A, Simonian A (2013) Biosensor technology: recent advances in threat agent detection and medicine. *Chem Soc Reviews* 42:8733–8768
- Kodikara CP, Crew HH, Stewart GS (1991) Near on-line detection of enteric bacteria using lux recombinant bacteriophage. *FEMS Microbiol Lett* 67:261–265
- Krikelis V, Markoulatos P, Spyrou N, Serie C (1985a) Detection of indigenous enteric viruses in raw sewage effluents of the city of Athens, Greece, during a two-year survey. *Water Sci Technol* 17:159–164
- Krikelis V, Spyrou N, Markoulatos P, Serie C (1985b) Seasonal distribution of enteroviruses in domestic sewage. *Can J Microbiol* 31:24–25
- Kruger DH, Barcak GJ, Smith HO (1988) Abolition of DNA recognition site resistance to the restriction endonuclease EcoRII. *Biomed Biochim Acta* 47:K1–K5
- Krupovic M, Dutilh BE, Adriaenssens EM, Wittmann J, Vogensen FK, Sullivan MB, Rummieks J, Prangishvili D, Lavigne R, Kropinski AM, Klumpp J, Gillis A, Enault F, Edwards RA, Duffy S, Clokie MR, Barylski J, Ackermann HW, Kuhn JH (2016) Taxonomy of prokaryotic viruses: update from the ICTV bacterial and archaeal viruses subcommittee. *Arch Virol*. doi:10.1007/s00705-015-2728-0
- Kumari S, Harjai K, Chhibber S (2011) Bacteriophage versus antimicrobial agents for the treatment of murine burn wound infection caused by *Klebsiella pneumoniae* B5055. *J Med Microbiol* 60:205–210
- Laanto E, Sundberg LR, Bamford JKH (2011) Phage specificity of the freshwater fish pathogen *Flavobacterium columnare*. *Appl Environ Microbiol* 77:7868–7872
- Labrie SJ, Samson JE, Moineau S (2010) Bacteriophage resistance mechanisms. *Nat Rev Microbiol* 8:317–327
- Lakshmanan RS, Guntupalli R, Hu J, Kim DJ, Petrenko VA, Barbaree JM, Chin BA (2007) Phage immobilized magnetoelastic sensor for the detection of salmonella typhimurium. *J Microbiol Meth* 71:55–60
- Lang JM, Gent DH, Schwartz HF (2007) Management of *Xanthomonas* leaf blight of onion with bacteriophages and a plant activator. *Plant Dis* 91:871–878
- Le Romancer M, Gaillard M, Geslin C, Prieur D (2007) Viruses in extreme environments. *Reviews in Environ Sci Biotech* 6:17–31
- Lee CY, Kim SJ, Park BC, Han JH (2016) Effects of dietary supplementation of bacteriophages against enterotoxigenic *Escherichia coli* (ETEC) K88 on clinical symptoms of post-weaning pigs challenged with the ETEC pathogen. *J Anim Physiol Anim Nutr (Berl)*. doi:10.1111/jpn.12513
- Leiman PG, Kanamaru S, Mesyanzhinov VV, Arisaka F, Rossmann MG (2003) Structure and morphogenesis of bacteriophage T4. *Cell Mole Life Sci* 60:2356–2370
- León M, Bastias R (2015) Virulence reduction in bacteriophages resistant bacteria. *Front Microbiol* 6:343. doi:10.3389/fmicb.2015.00343
- Leverentz B, Conway WS, Alavidze Z, Janisiewicz WJ, Fuchs Y, Camp MJ, Chighladze E, Sulakvelidze A (2001) Examination of bacteriophage as a biocontrol method for salmonella on fresh-cut fruit: a model study. *J Food Protect* 64:1116–1121
- Li W, Caberoy NB (2010) New perspective for phage display as an efficient and versatile technology of functional proteomics. *Appl Microbiol Biotechnol* 85:909–919
- Lin L, Honh W, Ji X, Han J, Huang L, Wei Y (2010) Isolation and characterization of an extremely long tail *Thermus* bacteriophage from Tegchong hot springs in China. *J Basic Micro* 50:452–456
- Liu Y, Zhang Q, Fang C, Zhu S, Tang Y, Huang S (2005) Effect of glutathione on UV induction in prophage lambda. *Arch Microbiol* 183:444–449
- Lone A, Anany H, Hakeem M, Aguis L, Avdjian AC, Bouget M, Atashi A, Brovko L, Rochefort D, Griffiths MW (2016) Development of prototypes of bioactive packaging materials based on immobilized bacteriophages for control of growth of bacterial pathogens in foods. *Int J Food Microbiol* 217:49–58
- Luria SE, Anderson TF (1942) The identification and characterization of bacteriophages with the electron microscope. *Proc Natl Acad Sci U S A* 28:127–130. doi:10.1073/pnas.28.4.127
- Madigan M, Martinko J (2006) Brock biology of microorganisms, 11th edn. Prentice Hall, USA
- Mai V, Ukhanova M, Visone L, Abuladze T, Sulakvelidze A (2010) Bacteriophage Administration Reduces the Concentration of *Listeria monocytogenes* in the Gastrointestinal Tract and Its Translocation to Spleen and Liver in Experimentally Infected Mice. *Int J Microbiol* 2010:624234. doi:10.1155/2010/624234
- Makarova KS, Grishin NV, Shabalina SA, Wolf YI, Koonin EV (2006) A putative RNA-interference-based immune system in prokaryotes: computational analysis of the predicted enzymatic machinery, functional analogies with eukaryotic RNAi, and hypothetical mechanisms of action. *Biol Direct* 1:7
- Maranger R, Bird DF (1995) Viral abundance in aquatic systems: a comparison between marine and fresh waters. *Mar Ecol Prog Ser* 121: 217–226
- Marcó MB, Moineau S, Quiberoni A (2012) Bacteriophages and dairy fermentations. *Bacteriophage*. 2:149–158
- Markoishvili K, Tsitlanadze G, Katsarava R, Morris G, Sulakvelidze A (2002) A novel sustained-release matrix based on biodegradable poly (esteramide)s and impregnated with bacteriophages and an antibiotic shows promise in management of infected venous stasis ulcers and other poorly healing wounds. *Int J Dermatol* 41:453–458
- Marraffini LA (2015) CRISPR-cas immunity in prokaryotes. *Nature* 526: 55–61
- Marraffini LA, Sontheimer EJ (2008) CRISPR interference limits horizontal gene transfer in staphylococci by targeting DNA. *Science* 322:1843–1845

- Martínez-Díaz SF, Hipólito-Morales A (2013) Efficacy of phage therapy to prevent mortality during the vibriosis of brine shrimp. *Aquaculture* 400:120–124
- Marza JA, Sothill JS, Boydell P, Collins TA (2006) Multiplication of therapeutically administered bacteriophages in *Pseudomonas* infected patients. *Burns* 32:644–666
- Maszevska A, Wójcik E, Ciurzyńska A, Wojtasik A, Piątkowska I, Dastych J, Różalski A (2016) Differentiation of polyvalent bacteriophages specific to uropathogenic *Proteus mirabilis* strains based on the host range pattern and RFLP. *Acta Biochim Pol*. doi:10.18388/abp.2015_1114
- Mathur MD, Vidhani S, Mehndiratta PL (2003) Bacteriophage therapy: an alternative to conventional antibiotics. *J Assoc Physicians India* 51:593–596
- Matsuzaki S, Yasuda M, Nishikawa H, Kuroda M, Ujihara T, Shuin T, Shen Y, Jin Z, Fujimoto S, Nasimuzzaman M, Wakiguchi H, Sugihara S, Sugiura T, Koda S, Muraoka A, Imai S (2003) Experimental protection of mice against lethal *Staphylococcus aureus* infection by novel bacteriophage phi MR11. *J Infect Dis* 187:613–624
- Mayer G (2016) Downloaded from: <http://www.microbiologybook.org/mayer/phage.htm>
- McKenna F, Tarabily KAE, Hardy GESTJ, Dell B (2001) Novel in vivo use of a polyvalent *Streptomyces* phage to disinfect *Streptomyces* scabies-infected seed potatoes. *Plant Pathol* 50:666–675
- Meaden S, Koskella B (2013) Exploring the risks of phage application in the environment. *Front Microbiol* 4:8
- Merabishvili M, Pimay JP, Verbeken G, Chanishvili N, Tediashvili M, Lashkhi N, Glonti T, Krylov V, Mast J, Parys LV, Lavigne R, Volckaert G, Mattheus W, Verween G, Corte PD, Rose T, Jennes S, Zizi M, Vos DD, Vaneechoutte M (2009) Quality-controlled small-scale production of a well-defined Bacteriophage cocktail for use in human clinical trials. *PLoS One* 4:1–104. doi:10.1371/journal.pone.0004944e4944
- Micallef SA, Goldstein RE, George A, Ewing L, Tall BD, Boyer MS, Joseph SW, Sapkota AR (2013) Diversity, distribution and antibiotic resistance of *Enterococcus* spp. recovered from tomatoes, leaves, water and soil on U.S. mid-Atlantic farms. *Food Microbiol* 36:465–474
- Mojica FJM, Diez-Villasenor C, Garcia-Martinez J, Soria E (2005) Intervening sequences of regularly spaced prokaryotic repeats derive from foreign genetic elements. *J Mol Evol* 60:174–182
- Mulani MS, Azhar S, Azharuddin S, Tambe S (2015) Harnessing the power of bacteriophage for pathogen reduction in wastewater. *Int J Curr Microbiol Appl Sci* 2:152–161
- Murphy FA, Fauquet CM, Bishop DHL, Ghabrial SA, Jarvis AW, Martelli GP, Mayo MA, Summers MD (eds) (1995) Virus taxonomy. Sixth report of the International Committee on Taxonomy of Viruses. Springer-Verlag, New York, N.Y
- Nagel B, Dellweg H, Gierasch LM (1992) Glossary for chemists of terms used in biotechnology—(IUPAC recommendations 1992. *Pure Appl Chem* 64:143–168
- Naidoo R, Singh A, Arya SK, Beadle B, Glass N, Tanha J, Szymanski CM, Evoy S (2012) Surface-immobilization of chromatographically purified bacteriophages for the optimized capture of bacteria. *Bacteriophage* 2:15–24
- Nakai T, Park SC (2002) Bacteriophage therapy of infectious diseases in aquaculture. *Res Microbiol* 153:13–18
- Nilsson AS (2014) Phage therapy—constraints and possibilities. *J Med Sci* 119:192–198
- Noorlis A, Ghazali FM, Cheah YK, Tuan Zainazor TC, Ponniah J et al (2011) Prevalence and quantification of *Vibrio* species and *Vibrio parahaemolyticus* in freshwater fish at hypermarket level. *Int Food Res J* 18:689–695
- Örmälä AM, Jalasvuori M (2013) Phage therapy: should bacterial resistance to phages be a concern, even in the long run? *Bacteriophage* 3:e24219. doi:10.4161/bact.24219
- Otte ML, Jacob DL (2006) Constructed wetlands for phytoremediation: rhizofiltration, phytostabilisation and phytoextraction. In: Mackova M, Dowling D, Macek T (eds) *Phytoremediation Rhizoremediation*. Springer, Dordrecht, pp. 57–67
- Pal C, Macia MD, Oliver A, Schachar I, Buckling A (2007) Coevolution with viruses drives the evolution of bacterial mutation rates. *Nature* 450:1079–1081
- Pande J, Szcwzyk MM, Grover AK (2010) Phage display: concept, innovations, applications and future. *Biotechnol Adv* 28:849–858
- Park S, Nakai T (2003) Bacteriophage control of *Pseudomonas plecoglossicida* infection in ayu *Plecoglossus altivelis*. *Dis Aquat Org* 53:33–39
- Park SC, Shimamura I, Fukunaga M, Mori KI, Nakai T (2000) Isolation of bacteriophages specific to a fish pathogen *Pseudomonas plecoglossicida* as a candidate for disease control. *Appl Environ Microbiol* 66:1416–1422
- Parracho HMRT, Burrows BH, Enright MC, McConville ML, Harper DR (2012) The role of regulated clinical trials in the development of bacteriophage therapeutics. *Journal of Molecular and Genetic Medicine* 6:279–286
- Paschke M (2006) Phage display systems and their applications. *Appl Microbiol Biotechnol* 70:2–11
- Peltomaa R, Perolio IL, Benito-Peña E, Barderas R, Moreno-Bondi MC (2015) Application of bacteriophages in sensor development. *Anal Bioanal Chem* 1–24
- Petrenko V (2008) Evolution of phage display: from bioactive peptides to bioselective nanomaterials. *Expert Opin Drug Deliv* 5:825–836
- Pfankuch E, Kausche GA (1940) Isolierung und übermikroskopische Abbildung eines Bakteriophagen. *Naturwissenschaften* 28:46. doi:10.1007/BF01486932
- Pietilä MK, Roine E, Sencilo A, Bamford DH, Oksanen HM (2016) Pleolipoviridae, a newly proposed family comprising archaeal pleomorphic viruses with single-stranded or double-stranded DNA genomes. *Arch Virol* 161(1):249–256. doi:10.1007/s00705-015-2613-x
- Pourcel C, Salvignol G, Vergnaud G (2005) CRISPR elements in *Yersinia pestis* acquire new repeats by preferential uptake of bacteriophage DNA, and provide additional tools for evolutionary studies. *Microbiology* 151:653–663
- Borah PK, Jindal JK, Verma JP (2000) Integrated management of bacterial leaf spot of mungbean with bacteriophages of Xav and chemicals. *J Mycol Plant Pathol* 30:19–21
- Pleška M, Qian L, Okura R, Bergmiller T, Wakamoto Y, Kussell E, Guet CC (2016) Bacterial autoimmunity due to a restriction-modification system. *Curr Biol* 26:404–409
- Popowska M, Rzeczycka M, Miernik A, Krawczyk-Balska A, Walsh F, Duffy B (2012) Influence of soil use on prevalence of tetracycline, streptomycin, and erythromycin resistance and associated resistance genes. *Antimicrob Agents Chemother* 56:1434–1443
- Prebus A, Hillier J (1939) The construction of a magnetic electron microscope of high resolving power. *Can J Res* 17:49–65. doi:10.1139/cjr39a-004
- Price LB, Stegger M, Hasman H, Aziz M, Larsen J, Andersen PS et al (2012) *Staphylococcus aureus* CC398: host adaptation and emergence of Methicillin resistance in livestock. *MBio* 3:00305–00311
- Prigent M, Leroy M, Confalonieri F, Dutertre M, DuBow MS (2005) A diversity of bacteriophages forms and genomes can be isolated from the surface sands of Sahara Desert. *Extremophiles* 9:289–296
- Ptashne M (2006) Lambda's switch: lessons from a modules wap. *Curr Biol* 16:459–462
- Quintin F, Kerrigan WC, Sothill JS (2005) Experimental bacteriophage protection against *Staphylococcus aureus* abscesses in a rabbit model. *Antimicrob Agents Chemother* 1220–1221

- Rackus DG, Shamsi MH, Wheeler AR (2015) Electrochemistry, biosensors and microfluidics: a convergence of fields. *Chem Soc Rev* 44: 5320–5340
- Rahmani R, Zarrini G, Sheikhzadeh F, Zadeh NAM (2015) Effective phages as green antimicrobial agents against antibiotic-resistant hospital *Escherichia coli* jundishapur. *J Microbiol* 8(2):e17744
- Rakhuba DV, Kolomiets EI, Dey ES, Novik GI (2010) Bacteriophage receptors, mechanisms of phage adsorption and penetration into host cell. *Pol J Microbiol* 59:145–155
- Ravat F, Jault P, Gabard J (2015) Bactériophages et phagothérapie: utilisation de virus Naturels pour traiter les infections bactériennes. *Ann Burns Fire Disasters* 28:13–20
- Ravensdale M, Blom TJ, Garza JAG, Svircev AM, Smith RJ (2007) Bacteriophages and the control of *Erwinia carotovora* subsp. *Carotovora* Can J Plant Pathol 29:121–130
- Reardon S (2014) Phage therapy gets revitalized. *Nature* 510:15–16
- Rhoads DD, Wolcott RD, Kuskowski MA, Wolcott BM, Ward LS, Sulakvelidze A (2009) Bacteriophage therapy of venous leg ulcers in humans: results of a phase I safety trial. *J Wound Care* 18:237–243
- Rivas L, Coffey B, McAuliffe O, Coffey A, Ross RP, McDonnell MJ, Duffy G, Burgess CM (2010) In vivo and ex vivo evaluations of Bacteriophages e11/2 and e4/1c for use in the control of *Escherichia coli* O157:H7. *Appl Environ Microbiol* 76:7210–7216
- Ruska H (1940) Über die Sichtbarmachung der bakterioophagen Lyse im Übermikroskop. *Naturwissenschaften* 28:45–46. doi:10.1007/BF01486931
- Saksida S, Constantine J, Karreman GA, Neville C, Sweeting R, Beamish R (2006) Evaluation of sea lice, *Lepeophtheirus salmonis*, abundance levels on farmed salmon in British Columbia, Canada. In The Proceedings from the International Symposium 6–11 august on Veterinary Epidemiology and Economics XI, Cairns, Australia
- Sangster W, Hegarty JP, Stewart DB (2014) Phage therapy for *Clostridium difficile* infection: an alternative to antibiotics? *Seminars in Colon and Rectal Surgery* 25(3):167–170
- Sangster W, Hegarty JP, Stewart DB Sr (2015) Phage tail-like particles kill *Clostridium Difficile* and represent an alternative to conventional antibiotics. *Surgery* 157:96–103
- Sarker SA, McCallin S, Barretto C, Berger B, Pittet AC, Sultana S, Krause L, Huq S, Bibiloni R, Bruttin A, Reuteler G, Brussow H (2012) Oral T4-like phage cocktail application to healthy adult volunteers from Bangladesh. *Virology* 434:222–232
- Säwström CH, Lisle J, Anesio AM, Priscu JC, Laybourn-Parry J (2008) Bacteriophage in polar inland waters. *Extremophiles* 12:167–175
- Servick K (2016) Beleaguered phage therapy trial presses on. *Science* 352:1506
- Sharma M, Patel JR, Conway WS, Ferguson S, Sulakvelidze A (2009) Effectiveness of bacteriophage in reducing *Escherichia coli* o157:H7 on fresh cut cantaloupes and lettuce. *J Food Protect* 72:1481–1485
- Sidhu SS (2000) Phage display in pharmaceutical biotechnology. *Curr Opin Chem Biol* 11:610–616
- Silva YJ, Costa L, Pereira C, Mateus C, Cunha Â, Calado R, Gomes NCM, Pardo MA, Hernandez I, Almeida A (2014) Phage therapy as an approach to prevent *Vibrio anguillarum* infections in fish larvae production. *PLoS One* 9:e114197. doi:10.1371/journal.pone.0114197
- Singh A, Arya SK, Glass N, Hanifi-Moghaddam P, Naidoo R, Szymanski CM, Tanha J, Evoy S (2010) Bacteriophage tail spike proteins as molecular probes for sensitive and selective bacterial detection. *Biosens Bioelectron* 26:131–138
- Singh A, Poshtiban S, Evoy S (2013) Recent advances in Bacteriophage based biosensors for food-borne pathogen detection. *Sensors* 13: 1763–1786. doi:10.3390/s130201763
- Smith GP (1985) Filamentous fusion phage: novel expression vectors that display cloned antigens on the virion surface. *Science* 228:1315–1317
- Sperinde JJ, Choi SJ, Szoka FC Jr (2001) Phage display selection of a peptide DNaseII inhibitor that enhances gene delivery. *The Journal of Gene Medicine* 3:101–108
- Stewart GSAB, Smith T, Denyer S (1989) Genetic engineering for bioluminescent bacteria. *Food Sci and Tech Today* 3:19–22
- Subasinghe RP, Bondad-Reantaso MG, McGladdery SE (2001) Aquaculture development, health and wealth. In *Aquaculture in the Millennium. Technica I Proceedings of the Conference on Aquaculture in the Third Millennium*; Suba singhe RP, Bueno P, Phillips MJ, Hough C, McGladdery SE, Arthur JR, Eds Bangkok Thailand
- Sulakvelidze A, Alavidze Z, Morris JG Jr (2001) Bacteriophage therapy. *Antimicrob Agents Chemother* 45:649–659
- Sulakvelidze A, Kutter E (2005) Bacteriophage therapy in humans. In: Kutter E, Sulakvelidze A (eds) *Bacteriophages: biology and applications*. CRC Press, USA, pp. 381–436
- Summers WC (1999) *Felix d'Herelle and the origins of molecular biology*. Yale University Press, New Haven, Conn
- Suttle CA (2005) Viruses in the sea. *Nature* 437:356–361
- Suttle CA (2007) Marine viruses - major players in the global ecosystem. *Nat Rev Microbiol* 5:801–812
- Sybesma W, Zbinden R, Chanishvili N, Kutateladze M, Chkhotua A, Ujmajuridze A, Mehnert U, Kessler TM (2016) Bacteriophages as potential treatment for urinary tract infections. *Front Microbiol* 7: 465. doi:10.3389/fmicb.2016.00465
- Tan D, Gram L, Middelboe M (2014) Vibriophages and their interactions with the fish pathogen *Vibrio anguillarum*. *Appl Environ Microbiol* 80:3128–3140
- Tartera C, Lucena F, Jofre J (1989) Human origin of *Bacteroides fragilis* bacteriophages present in the environment. *Appl Environ Microbiol* 55:2696–2701
- Thompson CC, Amaral GR, Campeão M, Edwards RA, Polz MF, Dutilh BE, Ussery DW, Sawabe T, Swings J (2015) Microbial taxonomy in the post-genomic era: rebuilding from scratch? *Archiv Microbiol* 197:359–370
- Tock MR, Dryden DT (2005) The biology of restriction and anti-restriction. *Curr Opin Microbiol* 8:466–472
- Toro H, Price SB, McKee AS, Hoerr FJ, Krehling J, Perdue M, Bauermeister L (2005) Use of bacteriophages in combination with competitive exclusion to reduce salmonella from infected chickens. *Avian Dis* 9:118–124
- Twort FW (1915) An investigation on the nature of ultra-microscopic viruses. *Lancet* 2:1241–1243
- Twort FW (1922) The bacteriophage: the breaking down of bacteria by associated filter passing lysins. *Br Med J* 2:293–296
- Twort FW (1949) The discovery of the bacteriophage. *Science News* 14: 33–34
- Van Helvoort T (1992) Bacteriological and physiological research styles in the early controversy on the nature of the bacteriophage phenomenon. *Med Hist* 3:243–270
- Verner-Jeffreys DW, Algoet M, Pond MJ, Virdee HK, Bagwell NJ, Roberts EG (2007) Furunculosis in Atlantic salmon (*Salmo salar* L.) is not readily controllable by bacteriophage therapy. *Aquaculture* 270:475–484
- Viertel TM, Ritter K, Horz HP (2014) Viruses versus bacteria-novel approaches to phage therapy as a tool against multidrug-resistant pathogens. *J Antimicrob Chemother* 69:2326–2336
- Vinod MG, Shivu MM, Umesha KR, Rajeeva BC, Krohne G, Indrani K, Iddya K (2006) Isolation of *Vibrio harveyi* bacteriophage with a potential for biocontrol of luminous vibriosis in hatchery environments. *Aquaculture* 255:117–124
- Wagner P, Acheson D, Waldor M (2001) Human neutrophils and their products induce Shiga toxin production by enterohemorrhagic *Escherichiacoli*. *Infect Immun* 69:1934–1937
- Wang Z, Wang D, Chen J, Sela DA, Nugen SR (2016) Development of a novel bacteriophage based biomagnetic separation method as an aid for sensitive detection of viable *Escherichia coli*. *Analyst* 141:1009–1016

- Watanabe R, Matsumoto T, Sano G, Ishii Y, Tateda K, Sumiyama Y, Uchiyama J, Sakurai S, Matsuzaki S, Imai S, Yamaguchi K (2007) Efficacy of bacteriophage therapy against gut-derived sepsis caused by *Pseudomonas aeruginosa* in mice. *Antimicrob Agents Chemother* 51: 446–452
- Watson BB, Eveland WC (1965) The application of the phage fluorescent antiphage staining system in the specific identification of *Listeria monocytogenes*. I. Species specificity and immunofluorescent sensitivity of *Listeria monocytogenes* phage observed in smear preparations. *Journal of Infectious Disease* 115:363–369
- Weitz JS, Wilhelm SW (2012) Ocean viruses and their effects on microbial communities and biogeochemical cycles. *F1000 Biology Reports* 4:17. doi:10.3410/B4-17
- Weitz JS and Wilhelm SW (2013) An ocean of viruses. *The Scientist* (July 1st issue). <http://www.the-scientist.com/?articles.view/articleNo/36120/title/An-Ocean-of-Viruses/>. Downloaded 3 Jan 2016
- Wicki M, Auckenthaler A, Felleisen R, Karabulut F, Niederhauser I, Tanner M, Baumgartner A (2015) Assessment of source tracking methods for application in spring water. *J Water Health* 13: 473–488
- Wildy P (1971) Classification and nomenclature of viruses. First report of the International Committee on Nomenclature of Viruses. S. Karger, Basel
- Willats WG (2002) Phage display: practicalities and prospects. *Plant Mol Biol* 50:837–854
- Williamson KE, Radosevich M, Wommack KE (2005) Abundance and diversity of viruses in six Delaware soils. *Appl Environ Microbiol* 71: 3119–3125
- Williamson KE, Wommack KE, Radosevich M (2003) Sampling natural viral communities from soil for culture-independent analyses. *Appl Environ Microbiol* 69:6628–6633
- Wills QF, Kerrigan C, Soothill JS (2005) Experimental bacteriophage protection against *Staphylococcus aureus* abscesses in a rabbit model. *Antimicrob Agents Chemother* 49:1220–1221
- Withey S, Cartmell E, Avery LM, Stephenson T (2005) Bacteriophages—potential for application in wastewater treatment processes. *Sci Total Environ* 339:1–18
- Włodarczyk KD, Vandenheuve D, Jang HB, Olszak T, Arabski M, Wasik S, Drabik M, Briers Y, Higgins G, Tyrrell J, Harvey BJ, Noben JP, Lavigne R, Kawa ZD (2016) A proposed integrated approach for the preclinical evaluation of phage therapy in *Pseudomonas* infections (2016). *Scientific Reports* 6:28115. doi:10.1038/srep28115
- Wright A, Hawkins CH, Anggård EE, Harper DR (2009a) A controlled clinical trial of a therapeutic bacteriophage preparation in chronic otitis due to antibiotic-resistant *Pseudomonas aeruginosa*; a preliminary report of efficacy. *Clin Otolaryngol* 34:349–357
- Young R, Wang IN, Roof WD (2000) Phages will out: strategies of host cell lysis. *Trends Microbiol* 8:120–128
- Zhu YG, Johnson TA, Su JQ, Qiao MG, Stedtfeld RD et al (2013) Diverse and abundant antibiotic resistance genes in Chinese wine farms. *Proceedings of the National Academy of Sciences USA* 110: 3435–3440
- Zschach H, Joensen KG, Lindhard B, Lund O, Goderdzishvili M, Chkonia I, Jgenti G, Kvatadze N, Alavidze Z, Kutter EM, Hasman H, Larsen MV (2015) What can we learn from a metagenomic analysis of a Georgian Bacteriophage cocktail? *Viruses* 7:6570–6589