**Bacteriophages: An Illustrated General Review**

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**Table of Contents**

**Bacteriophages: An Illustrated General Review**

Abstract .......................................................................................................................... 52
Introduction ..................................................................................................................... 53
  Background .................................................................................................................. 53
  History ......................................................................................................................... 53
  Taxonomy ..................................................................................................................... 55
  Morphology .................................................................................................................. 55
Conclusions ...................................................................................................................... 59
Acknowledgments ........................................................................................................... 59
References ....................................................................................................................... 59

**Abstract**

Bacteriophages are the most widespread entities on earth, which are able to invade prokaryotes. Bacteriophages play roles in dynamics of bacterial population, pathogenicity, epidemics and evolution. Bacteriophages are taxonomically categorized into two major orders of Caudovirales and Ligamenvirales, five families, nine subfamilies, 145 genera and 684 species; from which, family members of Siphoviridae, Myoviridae, Podoviridae and Inoviridae are the most widespread bacteriophages in nature. Bacteriophages are morphologically categorized into two main categories of tailed and polyhedral, filamentous or pleomorphic bacteriophages. They can have a single or double stranded DNA or RNA genome. The current review generally introduces bacteriophages using illustrations.

**Keywords:** Bacteriophage, prokaryote, bacteria, virus, morphology, taxonomy
Introduction

Background

Bacteriophages (simply phages) are viruses that infect prokaryotes, including eubacteria and archaea (1). In other words, bacteriophages dramatically hunt prokaryotes as their prey (bacteriophagy). This term means bacterial eating in Latin. These predators are more abundant on earth (approx. population of $10^{31}$) than any other biological entities (2–4). They are found everywhere, in soil, water, food and even saliva. The bacteriophage population is estimated as being ten times greater than the bacterial population (5). The infection rate of bacteria by the bacteriophages is approximated $10^{25}$ per second, ongoing at least for the last three billion years. Bacteriophages destroy a significant portion (1/5) of bacterial populations every two days and are more important in adoption of pathogens to new hosts than any other mechanism, with new strains emerging and epidemics occurring (6–8). Therefore, host-pathogen interaction is redefined by a novel concept “host-pathogen-phage interaction”.

History

Bacteriophages were first discovered in the late 1800s. In 1896, Ernest Hankin, an English chemical examiner and bacteriologist, inactivated Vibrio cholae using filtrates of water from the Ganges and Yamuna Rivers in northern India (Figure 1) (9). This was repeated later in 1898 by Nikolay Gamaleya, a Russian physician and scientist, while working with Bacillus subtilis. In 1915, Frederick Twort, an English bacteriologist, suggested that the causative agent was a “bacterial virus” (Figure 2) (10). In 1917, Felix d’Herelle, a French-Canadian microbiologist, used a plaque method and reported the first bacterial lysis (Figures 3 and 4) (11). He named the causative agent a “bacteriophage”, which means bacteria eater in Greek. Most of the early applied knowledge of the bacteriophages are results of his studies (Figure 5). Clark and Clark (1927) made the first successful isolation of a streptococcal bacteriophage (12). Enterococcal bacteriophages were firstly isolated from sewage by Evans (1934) and later by Rakieten and Tiffany (1938), Tiffany and Rakieten (1939) and Evans (1941) (13–16).

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Kjems (1955) found evidence of lysogeny and isolated four strains of bacteriophages from Group D streptococci, including Streptococcus faecalis and S. faecium (17). Further isolations were carried out by Bleiweis and Zimmerman (1961), Brock et al. (1963) and Brock (1964) (18–20). Furthermore, the golden era of bacteriophage research has begun from 1938 when Max Delbruck, a German physicist and biologist, started a series of studies on physiology and ecology of the bacteriophages at the California Institute of Technology, USA. Soon, Italian Salvadore Luria and American Al Hershey joined him and carried out some most interesting genetic studies on bacteriophages. The genetic studies were continued by Luria and Human (1952) and Arber and Dussoix (1962) and many other researchers worldwide (21, 22).

### Table 1. Taxonomy and classification of bacteriophages

<table>
<thead>
<tr>
<th>Order</th>
<th>Family</th>
<th>Head</th>
<th>Tail</th>
<th>Env.</th>
<th>Gen.</th>
<th>Other</th>
</tr>
</thead>
<tbody>
<tr>
<td>Caudovirales</td>
<td>Myoviridae, A1</td>
<td>Isometric</td>
<td>Contractile</td>
<td>No</td>
<td>Linear dsDNA</td>
<td>Head L/W ratio = 1.3–1.8</td>
</tr>
<tr>
<td>Caudovirales</td>
<td>Myoviridae, A2</td>
<td>Elongated</td>
<td>Contractile</td>
<td>No</td>
<td>Linear dsDNA</td>
<td>Head L/W ratio ≥ 2</td>
</tr>
<tr>
<td>Caudovirales</td>
<td>Myoviridae, A3</td>
<td>Elongated</td>
<td>Contractile</td>
<td>No</td>
<td>Linear dsDNA</td>
<td></td>
</tr>
<tr>
<td>Caudovirales</td>
<td>Podoviridae, C1</td>
<td>Isometric</td>
<td>Short, noncontractile</td>
<td>No</td>
<td>Linear dsDNA</td>
<td></td>
</tr>
<tr>
<td>Caudovirales</td>
<td>Podoviridae, C2</td>
<td>Elongated</td>
<td>Short, noncontractile</td>
<td>No</td>
<td>Linear dsDNA</td>
<td>Head L/W ratio = 1.4</td>
</tr>
<tr>
<td>Caudovirales</td>
<td>Podoviridae, C3</td>
<td>Elongated</td>
<td>Short, noncontractile</td>
<td>No</td>
<td>Linear dsDNA</td>
<td>Head L/W ratio ≥ 2.5</td>
</tr>
<tr>
<td>Caudovirales</td>
<td>Siphoviridae, B1</td>
<td>Isometric</td>
<td>Long, noncontractile</td>
<td>No</td>
<td>Linear dsDNA</td>
<td></td>
</tr>
<tr>
<td>Caudovirales</td>
<td>Siphoviridae, B2</td>
<td>Elongated</td>
<td>Long, noncontractile</td>
<td>No</td>
<td>Linear dsDNA</td>
<td>Head L/W ratio = 1.2–2</td>
</tr>
<tr>
<td>Caudovirales</td>
<td>Siphoviridae, B3</td>
<td>Elongated</td>
<td>Long, noncontractile</td>
<td>No</td>
<td>Linear dsDNA</td>
<td>Head L/W ratio ≥ 2.5</td>
</tr>
<tr>
<td>Ligamenvirales</td>
<td>Lipothrixviridae</td>
<td>PFP</td>
<td>PFP</td>
<td>Yes</td>
<td>Linear dsDNA</td>
<td>Filaments, lipids</td>
</tr>
<tr>
<td>Ligamenvirales</td>
<td>Rudiviridae</td>
<td>PFP</td>
<td>PFP</td>
<td>No</td>
<td>Linear dsDNA</td>
<td>Helical rods</td>
</tr>
<tr>
<td>No order</td>
<td>Corticoviridae</td>
<td>No</td>
<td>Circular supercoiled dsDNA</td>
<td>Yes</td>
<td>Circular ssDNA</td>
<td>Icosahedral capsid, lipids</td>
</tr>
<tr>
<td>No order</td>
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<td>PFP</td>
<td>PFP</td>
<td>Yes</td>
<td>Circular supercoiled dsDNA</td>
<td>Icosahedral capsid, lipids</td>
</tr>
<tr>
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<td>Guttaviridae</td>
<td>PFP</td>
<td>PFP</td>
<td>Yes</td>
<td>Circular ssDNA</td>
<td>Pleomorphic capsid, protrusions, bead of long fibers</td>
</tr>
<tr>
<td>No order</td>
<td>Fuselloviridae</td>
<td>PFP</td>
<td>PFP</td>
<td>Yes</td>
<td>Circular supercoiled dsDNA</td>
<td>Pleomorphic capsid, lipids, protrusions, short tail-like fibers</td>
</tr>
<tr>
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<td>Inoviridae: Inovirus</td>
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<td>PFP</td>
<td>No</td>
<td>Circular ssDNA</td>
<td>Long filaments with helical symmetry</td>
</tr>
<tr>
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<td>Inoviridae: Plecovirus</td>
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<td>PFP</td>
<td>No</td>
<td>Circular ssDNA</td>
<td>Short rods with helical symmetry</td>
</tr>
<tr>
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<td>PFP</td>
<td>No</td>
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<td>Quasi-icosahedral capsid</td>
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<td>PFP</td>
<td>No</td>
<td>Circular ssDNA</td>
<td>Icosahedral capsid</td>
</tr>
<tr>
<td>No order</td>
<td>Plasmaviridae</td>
<td>PFP</td>
<td>PFP</td>
<td>Yes</td>
<td>Circular supercoiled dsDNA</td>
<td>Pleomorphic capsid, no capsid, lipids,</td>
</tr>
<tr>
<td>No order</td>
<td>Tectiviridae</td>
<td>PFP</td>
<td>PFP</td>
<td>No</td>
<td>Linear dsDNA,</td>
<td>Icosahedral capsid with inner lipoprotein vesicle, tail is produced for DNA injection</td>
</tr>
<tr>
<td>Unassigned</td>
<td>Salterprovirus</td>
<td>PFP</td>
<td>PFP</td>
<td>Yes</td>
<td>Linear dsDNA</td>
<td>Pleomorphic capsid, lemon-shaped, short tail-like fibers</td>
</tr>
</tbody>
</table>
Taxonomy

Since research on bacteriophages is a fast developing trend worldwide and because much genetic data from bacteriophages are published every year, bacteriophage taxonomy changes regularly. Like other viruses, bacteriophages are classified based on general criteria, including type of genetic material (DNA or RNA, single or double stranded, linear or circular, segmented or non-segmented), morphology (e.g. symmetry versus asymmetry, presence or absence of the tail), capsid components (e.g. existence of envelope, lipid molecules), virion dimensions and the bacteriophage life cycle and strategies (23). Of these criteria, the first criterion is critical in Baltimore classification. Currently, the International Committee on Taxonomy of Viruses (ICTV) classifies bacteriophages in two orders (Caudovirales and Ligamenvirales), five families, nine subfamilies, 145 genera and 684 species and further unassigned families mostly belong to archaea (Table 1) (24, 25). Bacteriophage families are alphabetically listed as follows: Corticoviridae, Cystoviridae, Fuselloviridae, Guttaviridae, Inoviridae, Levirviridae, Lipothrixviridae, Microviridae, Myoviridae, Plasmaviridae, Podoviridae, Rudiviridae, Siphoviridae and Tectiviridae. Of these families, four families seem to be more widespread in nature, namely Siphoviridae (Order Caudovirales), Myoviridae (Order Caudovirales), Podoviridae (Order Caudovirales) and Inoviridae (no order) (26, 27). A dilemma is seen here which is linked to the bacteriophage nomenclature. The question is that which prokaryotes (eubacteria or archaea) can be referred as the host of bacteriophages, since Ligamenvirales members have been suggested recently as archaean viruses by ICTV (28). However, most of the bacteriophage families have been reported in eubacteria. Therefore, the current review still virtually uses “bacteriophage” as a general name for all prokaryotic viruses to avoid conflict.

Morphology

Bacteriophages are morphologically categorized to two major categories of tailed (head-tail) and PFP (polyhedral, filamentous or pleomorphic) bacteriophages (Figures 6–14). A typical structure of tailed bacteriophages consists of an icosohedral head containing the virus capsid (including units of capsomers), genome and sometimes a lipid coat, a neck or collar, a tail (some contain multiple tails) and a baseplate (Figure 15) (29, 30). The tail includes a core covered by a helical sheath that ends to the base plate, which bears long tail fibers and short tail fibers or tail pins. The genome of bacteriophages can be composed of either single or double-stranded DNA or RNA and can range in size from a few to several hundred kilobases (2, 31). The largest bacteriophage genome belongs to Klebsiella Phage vB_KleM-RaK2, including 346 kb (32). Most genes in bacteriophages belong to lysis, replication and regulation, packaging and structural, antibiotic resistance and housekeeping genes. In general, these genes can be classified into two major groups: housekeeping and regulatory genes. Housekeeping genes encode proteins such as head-tail joining proteins, head-tail adaptor proteins, host specificity proteins, portal proteins, prohead protease, tail component, tail major proteins, tape measure proteins, terminase large subunit and terminase small subunit. Regulatory genes encode proteins such as amidase, holing, HNH endonuclease and DNA primase (33). Since 1959, thousands of bacteriophages have been examined by electron microscope (EM) (34). However, the first bacteriophage was observed by Helmut Ruska, a German...
Bacteriophages: An Illustrated General Review

Fig. 6. An isometric head, long tailed bacteriophage (TEM 130,000×) (courtesy of R. Mazaheri Nezhad Fard).

Fig. 7. Isometric head, long tailed bacteriophages (TEM 92,000×) (courtesy of R. Mazaheri Nezhad Fard).

Fig. 8. A tailed bacteriophage (TEM 130,000×). Circular double-stranded genome is removed from the bacteriophage (courtesy of R. Mazaheri Nezhad Fard).

Fig. 9. A long head, long tailed bacteriophage (TEM 180,000×) (courtesy of R. Mazaheri Nezhad Fard).

Fig. 10. An isometric head, long tailed bacteriophage (TEM 92,000×) (courtesy of R. Mazaheri Nezhad Fard).

Fig. 11. Isometric head, short tailed bacteriophages (TEM 92,000×) (courtesy of R. Mazaheri Nezhad Fard).

Fig. 12. A filamentous bacteriophage (TEM 92,000×) (courtesy of R. Mazaheri Nezhad Fard).

Fig. 13. A droplet shaped (pleomorphic) bacteriophage (TEM 92,000×). Circular double-stranded genome is leaking from the bacteriophage (courtesy of R. Mazaheri Nezhad Fard).

Fig. 14. A lemon shaped (pleomorphic) bacteriophage (TEM 130,000×) (courtesy of R. Mazaheri Nezhad Fard).

physician, biologist, in 1939 using a prototype EM invented by his brother Ernst Ruska (35). Research indicate that approximately 96% of all bacteriophages have a tail; of which, up to 90% feature a single tail while a minority are polyhedral, filamentous or pleomorphic (PFP) (34, 36). Most tailed bacteriophages (nearly 61%) feature long, flexible, noncontractile tails and are categorized in the Siphoviridae family (Order Caudovirales) (37, 38). Some examples include c2, L5, lambda, N15, phiC31, PsiM1, T1 and T5 bacteriophages (Figure 16). This is followed by the members of Myoviridae family (Order Caudovirales) with long, nonflexible, contractile tails (Figure 17) and Podoviridae family (Order Caudovirales) with long, nonflexible, noncontractile tails (Figure 18). Examples of Myoviridae include Bcep781, BcepMu, FelixO1, HP1, I3, KVP40, Mu, P1, P2, PB1, phiCD119, phiH, phiHAP-1, phiKZ, SP01, T4 and Twort bacteriophages and examples of Podoviridae include 44AHJD, BPP-1, epsilon15, LUZ24, N4, P22, phi29, phiEco32, phiKMV, SP6 and T7 bacteriophages. Filamentous bacteriophages are categorized in three families: Inoviridae (no order; e.g. L51, M13, Pf1) (Figure 19), Lipothrixviridae (Order Ligamenvirales; e.g.
TTV-1, SIFV, AFV-1, AFV-2 (Figure 20) and Rudiviridae (Order Ligamenvirales; e.g. SIRV-1, SIRV-2) (Figure 21) (39). These bacteriophages are found in both eubacteria (Inoviridae) (40) and archaea (Lipothrixviridae and Rudiviridae) (41). Inoviridae are either long slender tube or short rod bacteriophages with a maximum size of 2 μm and contain a single-stranded circular DNA genome with 4.5–8 kb encoding 4–10 proteins (42). They are distributed worldwide and can be isolated from a variety of bacteria such as enterobacteria, Vibrio spp., Pseudomonas spp., Xanthomonas spp., Acholeplasma spp. and Spiroplasma spp. Polyphagy is a characteristic of filamentous bacteriophages such as Inoviridae; in which, more than one phage genome are packed in a bacteriophage capsid. Other filamentous bacteriophage families belong to thermophilic and hyperthermophilic archaea. Lipothrixviridae (Order Ligamenvirales) are flexible, enveloped, rod-shaped bacteriophages with 24–38 and 410–1950 nm dimensions (28). The family includes four genera of Alphalipothrixivirus, Betalipothrixivirus, Gammalipothrixivirus and Deltalipothrixivirus (43). Genome consists of a linear dsDNA genome including 15.9–56 kb. Rudiviridae (Order Ligamenvirales) are rigid, non-enveloped, rod-shaped bacteriophages with three tail fibers at each end (44). The family includes one genus of Rudivirus. Genome consists of a linear dsDNA genome including 32–35 kb and inverted terminal repeats as well as direct repeats at both ends. Poliovirus-like (the semi-suffix “like” has recently been removed from the bacteriophage names by ICTV but may still be used in some papers) bacteriophage group belongs to Leviviridae family (no order) of bacteriophages (e.g. MS2, Qbeta) (Figure 22). These polyhedral bacteriophages contain a single-stranded linear molecule of RNA (45). Single-stranded RNA bacteriophages are widely found in bacterial isolates from sewage and feces of mammals but unlikely to be found in avian bacterial species (46). Leviviridae are small bacteriophages with a reported size of 23–30 nm (47, 48). Tectiviridae (no order; e.g. AP50, Bam35, P17-14, PRD1) are non-enveloped, icosahedral bacteriophages with capsid symmetry (Figure 23). Virion size is nearly 66 nm with inner membrane vesicle and 20-nm apical spikes. Although the bacteriophage is a non-tailed one, a tail is produced for DNA injection during the host
infection process (Figure 24). Genome consists of a ~15-kb coiled, linear, dsDNA molecule (49). Corticoviridae bacteriophages (no order; e.g. marine PM2) include ~56-nm icosahedral virions with an internal lipid core and a ~10-kb circular, supercoiled dsDNA genome encoding 21 proteins (Figure 25) (50). There is one genus within the family namely Cortocovirus. Cystoviridae family members (no order; e.g. Phi6–Phi14) are ~85-nm enveloped, spherical
virions with double capsid structure (Figure 26) (51). One genus has been described for the family, namely Cystovirus. The total 13.3-kb genome is segmented, linear dsDNA and contains three segments of large (L), medium (M) and small (S), encoding 12 proteins. Microviridae (no order; e.g. Chp-1, phiMH2K, phiX174, SpV4) includes ~30-nm non-enveloped, round, icosahedral bacteriophages with spikes (Figure 27) (52). Genome is described as circular ssDNA(+) with 4.5–6 kb in length, encoding A–H and J–K proteins. Plasmaviridae (no order, e.g. L2) is a ~80-nm enveloped, spherical/plasmoviral bacteriophage with one genus of Plasmavirus (Figure 28) (53, 54). Genome of the bacteriophage consists of a ~12-kb low-GC circular, supercoiled dsDNA. Droplet-shaped bacteriophages are classified in the Guttaviridae family (no order; e.g. Sulfolobus newzealandicus droplet-shaped virus) (Figure 29); firstly investigated by Zillig et al. (1998) and described by Arnold et al. (2000) in archaea and later by Mazaheri Nezhad Fard et al. (2010) in eubacteria (55–58). Two genera of Alphaguttavirus and Betaguttavirus are seen in the family. These bacteriophages bear long thin filaments (beard) at one end and contain a dsDNA genome (44, 59). The genome consists of single segment containing a circular dsDNA molecule (approx. size of 20 kb) (60–62). This segment forms a covalently closed circle, which is extensively methylated (60Sau and Deb, 2008; 61). Fuselloviridae family (no order; e.g. SSV-1) represents lemon/spindle-shaped bacteriophages having peripheral projections and short tail-like fiber(s) attached to one pole (Figure 30) (59, 63, 64). Two genera of Alphafusellovirus and Betafusellovirus are observed within the family. The bacteriophages contain a single molecule of circular, supercoiled dsDNA with 14.8–17.3 kb encoding 31–37 genes. As with Guttaviridae, Fuselloviridae have mostly been isolated on or from archaea but first reported in eubacteria by Mazaheri Nezhad Fard et al. in 2010 (55, 65, 66). However, detection of possible archaean bacteriophages in eubacteria cannot be completely ruled out as other bacteriophage families (Siphoviridae and Myoviridae) are found in both bacterial kingdoms (41, 55). Another lemon-shaped bacteriophage, the unassigned Salterproirus (e.g. His1 virus, His2 virus), is a pleomorphic, enveloped bacteriophage with short tail-like fibers and a ~14.5-kb linear dsDNA genome (Figure 31) (67). The bacteriophage habitats include high salted waters and lakes (the reason for naming).

Conclusions

In conclusion, bacteriophages are a diverse group of biological entities with great promising potentials of contributions to a variety of sciences. Study of bacteriophages is an interesting challenge, which opens new horizons in front of researchers. Not only are these subjects mostly unknown, they include many useful features from research to treatment which should be explored.

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Bacteriophages: An Illustrated General Review

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