

Emerging Aspects of Jumbo Bacteriophages

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Abstract: The bacteriophages have been explored at a huge scale as a model system for their applications in many biological-related fields. Jumbo phages with a large genome size from 200 to 500 kbp were not previously assigned a great value, and characterized by complex structures coupled with large virions with a wide variety of hosts. The origin of most of the jumbo phages was not well understood; however, many other prominent features have been discovered recently. In the current review, we strive to unearth the most advanced characteristics of jumbo phages, particularly their significance and structural organization that holds immense value to the viral life cycle. The unique characteristics of jumbo phages are the basis of variations in different types of phages concerning their organization at the genomic level, virion structure, evolution, and progeny propagation. The presence of tRNA and additional translation-related genes along with chaperonin genes mark the ability of these phages for being independent of host molecular machinery enabling them to have wide host options. A large number of jumbo phages have been isolated from various sources through advanced standard screening methods. The current review has summarized the available data on jumbo phages and discussed the genome orientation of jumbo phages, translational machinery, diversity and evolution of jumbo phages. In the studies conducted, jumbo phages possessed special additional genes that helps to reduce the dependence of jumbo phages on their hosts. Furthermore, their genomes might have evolved from smaller genome phages.

Keywords: bacteriophages, tRNA, large genome, wide host range, chaperonin

Introduction

Bacteriophages bearing tails with genome size beyond 200 kbp are termed jumbo bacteriophages.¹ Many studies have tried to explore the new jumbo phages that infect bacteria of different species.^{2–7} They are also different from other bacteriophages not only because of size but also due to evolutionary path, genetic orientation, structure, and progeny transmission.⁸ Thus, they attained scientist's attention to isolate and characterize the new species of jumbo phages, as well as detailed elucidation about their biological mechanisms.⁹ Specific factors in the evolution of these phages are needed to explore the tendency toward gigantism. There are only a few numbers of tailed bacteriophages which have such large genome size.¹⁰ Mostly, predicted proteins of jumbo bacteriophages do not show similarity to protein sequences available in databases. The genome size of jumbo phages is too large to compare traditionally with smaller phages, but functions can be assigned to genes based on evaluation from smaller phages through comparative analysis.⁸ Comparisons have been done to check the genome similarity as it might be suggested that these have been evolved from small genome phages possibly due to limitations posed by the capsid size of the genome.¹¹ N1M2 was selected as an example to further investigate by phylogenetic and genomic analyses and it was

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found to be similar to jumbo phage-related detached to *Pseudomonas* OBP. It was studied previously and exhibited to be possessing likeness with phiKZ-like jumbo phages.²

The ICTV's Bacterial and Archaeal Viruses Subcommittee has now classified tailed phages into the following structural groups according to genomic data. Three new families of myoviruses have been officially approved Ackermannviridae, Chaseviridae, Herelleviridae; two for the siphoviruses, Demereviridae and Drexelviriidae, and one of podoviruses, Autographiviridae.¹² This is a latest taxonomy of tailed bacteriophages. According to this classification cultured jumbo phages that are reported in the literature until now are Myo and Siphoviruses (Table 1), suggesting that jumbo phages spread independently in different phage groups.¹³ Jumbo phages showed many variations in their head and tail morphologies and hair-like projections and long whiskers from both the head and tail sheath present in some cases.¹⁴

Until now, over 150 jumbo phages have been isolated since the discovery of phages (GenBank database, last accessed 1 June 2021). More than 85% isolated in just past 4 to 5 years, as research on bacteriophages revitalized and advanced by high throughput sequencing techniques. Their genomes were completely sequenced, exhibiting various remarkable aspects of their biology.^{15,16} These could be isolated through already in-use methods like agar diffusion and filtration. Due to the large genome size and capsid, jumbo phages might not be passed from the membrane and removed with bacteria.^{17,18} This can be the possible reason for the isolation of only a few jumbo phages until now. Additionally, we discussed only about cultured jumbo phages reported in the literature and have genome size larger than 200 kbp, phages approaching genome size 200 kbp will not be discussed in this review. We are trying to discuss the latest aspects related to jumbo bacteriophages including their general characteristics, diversity, and evolution in this review.

Hosts and Distribution

Jumbo phages were extracted from a variety of atmospheres ranging from marine sediments, water to soil constituted habitats like composts, animal feces, silkworms, and plant tissues (Table 1). But most recurrently, these phages were isolated from aquatic environments which provides ease in some of their functions like diffusion and infection to host bacteria which explains their frequency in such environments.^{8,19}

Jumbo phages have been identified from gammaproteo bacteria, betaproteo bacteria, alphaproteo bacteria, zeta-proteo bacteria, bacteroidetes, cyanobacteria, sporulating firmicutes, and actinobacteria. Currently, mostly jumbo phages present in databases are infecting gammaproteo bacteria.⁸ Most of the time these phages were isolated from a variety of gram-negative bacteria, while only eleven jumbo phages were extracted from gram-positive, the host is a common one from *Bacillus* spp.²⁰ It is still needed to investigate whether the affinity of these eleven phages to *Bacillus* is because of their structural compatibility or just because of the discovery of a small number of jumbo phages studied currently.^{8,21} Further studies shed light on a phenomenon that why these four phages have an attraction toward gram-positive strain.

Genome Structure and Orientation

The most remarkable property that distinguishes the jumbo phages from smaller phages is their large genome size. The first ever jumbo phage with the biggest known size so far is phage G with capsid size 160 nm, tail 453 nm in length, and genome size 497 kbp.²² PhiKZ phage particles possess large sized hexagonal heads of about 120 nm and contractile tail of about 180×20 nm in size.²³ Due to their large capsids, jumbo phages can regulate larger genomes compared to the phages having smaller capsids.¹⁶ It has been observed that the genome of phage G is only smaller just by 87 kbp than *Mycoplasma genitalium*, the genome of the smallest known bacterium.²⁴ Big size enable the jumbo genomes to carry a lot of genes that normally are not possible to hold by smaller genome size phages. For instance, all reported jumbo phages possess more genome replication and nucleotide metabolism facilitating genes while some jumbo phages possessed more than one paralogous gene for DNA polymerase and RNA polymerase.²⁵ The RNA Polymerase (RNAP) enzymes mostly are multi-subunit RNAP which are encoded by the phage genomes, while some of those have been observed in phage virions.⁸ The structural fragments of RNAP constitute multiple subunits and early gene transcription could be mediated by injecting subunits to host bacteria even before the expression of phage and host RNAP.²⁶ Detailed transcriptomic-based analysis of this phage's injection to the host highlights the fact that the expression of phage genes is only dependent upon the phage's RNAP and cellular machinery rather than the host's RNAP.²⁷ These phages also have more enzymes and proteins to take on the

Table 1 General Characteristics of Jumbo Bacteriophages

Phage	Host	Family	Viron Size (nm)		Genome Size (nt)	No. of ORFs	No. of tRNA	Sample Source	Accession No.	References
			Head	Tail						
G	<i>Bacillus megaterium</i>	Myoviridae	160	453	497,513	675	20	NA	NC_023719	Donelli et al 1975
vB_CsaM_GAP32	<i>Cronobacter sakazakii</i>	Myoviridae	115	118	358,663	545	26	Wastewater	NC_019401	Abbasifar et al 2014
I21Q	<i>Escherichia coli</i>	Myoviridae	116	115	348,532	611	7	Sewage	NC_025447	Ackermann and Nguyen 1983
PBEO 4	<i>Escherichia coli</i>	Myoviridae	132	125	348,113	551	6	River water	NC_027364	Kim et al 2013
K64-1	<i>Klebsiella pneumoniae</i>	Myoviridae	NA	NA	346,602	64	NA	NA	NC_027399	Pan et al 2015
vB_KleM-RaK2	<i>Klebsiella</i> sp.	Myoviridae	123	128	345,809	534	5	NA	NC_019526	Šimoliūnas et al 2013
201phi2-1	<i>Pseudomonas chlororaphis</i>	Myoviridae	129	200	316,674	461	1	Soil	NC_010821	Thomas et al 2008
PhiPA3	<i>Pseudomonas aeruginosa</i>	Myoviridae	100	185	309,208	375	5	Sewage	NC_028999	Monson et al 2011
Phabio	<i>Pseudomonas fluorescens</i>	Myoviridae	134	174	309,157	469	3	Soil	MF042360	Joanna et al 2017
Xoo-sp13	<i>Xanthomonas oryzae</i> pv. <i>oryzae</i>	Myoviridae	60	121	309,023	401	28	Soil	MN047793	Nazir et al 2020
OBP	<i>Pseudomonas fluorescens</i>	Myoviridae	119	191	284,757	309	4	Compost	NC_016571	Cornelissen et al 2012b
Lu11	<i>Pseudomonas putida</i>	Myoviridae	124	200	280,538	391	0	Soil	NC_017972	Adriaenssens et al 2012
phiKZ	<i>Pseudomonas aeruginosa</i>	Myoviridae	145	160	280,334	306	6	Sewage	NC_004629	Mesyanzhinov et al 2002
Noxifer	<i>Pseudomonas fluorescens</i>	Myoviridae	130	191	278,136	334	4	Soil	MF063068	Joanna et al 2017
CcrColossus	<i>Caulobacter crescentus</i>	Myoviridae	292x95	65	279,967	448	28	Surface water	NC_019406	Meczker et al 2014
KTN4	<i>Pseudomonas aeruginosa</i>	Myoviridae	130	168	279,593	368	6	Irrigated fields	KU521356.1	Danis-Włodarczyk et al 2016
vB_EamM_Deimos-Minion	<i>Erwinia amylovora</i>	Myoviridae	NA	NA	273,501	326	NA	Tree	KU886225.1	Espin et al 2017

(Continued)

Table 1 (Continued).

Phage	Host	Family	Viron Size (nm)		Genome Size (nt)	No. of ORFs	No. of tRNA	Sample Source	Accession No.	References
			Head	Tail						
vB_EamM_Special G	<i>Erwinia amylovora</i>	Myoviridae	NA	NA	273,224	324	0	Branches and blossoms	KU886222.1	Esplin et al 2017
vB_EamM_RAY	<i>Erwinia amylovora</i>	Myoviridae	NA	NA	271,182	319	1	Tree	KU886224.1	Esplin et al 2017
vB_EamM_Simmy50	<i>Erwinia amylovora</i>	Myoviridae	NA	NA	271,088	321	1	Bark	KU886223.1	Esplin et al 2017
Ea35-70	<i>Erwinia amylovora</i>	Myoviridae	NA	NA	271,084	318	1	Soil	NC_023557	Yagubi et al 2014
PA7	<i>Pseudomonas aeruginosa</i>	Myoviridae	NA	NA	266,743	341	NA	Mudflat	JX233784.1	Kwan et al 2006
phiR1-37	<i>Yersinia enterocolitica</i>	Myoviridae	138	383	262,391	367	5	Sewage	NC_016163	Kiljunen et al 2005
vB_EamM_Yoloswag	<i>Erwinia amylovora</i>	Myoviridae	NA	NA	259,700	334	NA	Tree	KY448244.1	Esplin et al 2017
PaBG	<i>Pseudomonas aeruginosa</i>	Myoviridae	136	220	258,139	308	NA	Lake water	NC_022096	Sykilinda et al 2014
vB_BpuM_BpSp	<i>Bacillus pumilus</i>	Myoviridae	137	192	255,569	318	0	Soil	KT895374	Yuan et al 2016a, b
NIM2	<i>Klebsiella aerogenes</i>	Myoviridae	113	158	253,367	257	24	Fecal	MN642089	Rhea Lewis et al 2020
P-SSM2	<i>Prochlorococcus</i>	Myoviridae	115	123	252,401	334	1	Seawater	NC_006883	Sullivan et al 2005
AR9	<i>Bacillus subtilis</i>	Myoviridae	NA	NA	251,042	291	1	NA	NC_031039	Laysh et al 2016
ValKK3	<i>Vibrio alginolyticus</i>	Myoviridae	NA	NA	248,088	390	NA	Marine sediment	NC_028829	Lal et al 2016
nt-1	<i>Vibrio natriegens</i>	Myoviridae	NA	NA	247,511	405	28	Marine sediment	NC_021529	Comeau et al 2014
VH7D	<i>Vibrio harveyi</i>	Myoviridae	NA	NA	246,964	327	NA	Seawater	NC_023568	Luo et al 2015
phi-pp2	<i>Vibrio parahaemolyticus</i>	Myoviridae	90x50	110	246,421	383	30	Aquaculture waterway	JN849462.1	Lin and Lin 2012
vB_EamM_Kwan	<i>Erwinia amylovora</i>	Myoviridae	NA	NA	246,390	285	8	NA	NC_031010	Esplin et al 2017
vB_EamM_Asesino	<i>Erwinia amylovora</i>	Myoviridae	NA	NA	246,291	277	NA	NA	NC_031107	NA
vB_EamM_ChrisDB	<i>Erwinia amylovora</i>	Myoviridae	NA	NA	244,840	277	11	Tree	NC_031126	Esplin et al 2017
KVP40	<i>Vibrio parahaemolyticus</i>	Myoviridae	140x70	NA	244,834	381	30	Marine sediment	NC_005083	Miller et al 2003
phiEaH2	<i>Erwinia amylovora</i>	Siphoviridae	NA	NA	243,050	262	NA	Soil	NC_019929	Dömötör et al 2012
vB_EamM_Stratton	<i>Erwinia amylovora</i>	Siphoviridae	NA	NA	243,953	276	12	Tree	KX397373.1	Esplin et al 2017

vB_EamM_Machina	<i>Erwinia amylovora</i>	Myoviridae	NA	NA	241,654	272	9	Tree	KX397370.1	Esplin et al 2017
vB_EamM_Caitlin	<i>Erwinia amylovora</i>	NA	NA	241,147	271	7	Tree	NC_031120	Esplin et al 2017	
vB_EamM_Parshik	<i>Erwinia amylovora</i>	Myoviridae	NA	241,050	271	10	Tree	KX397371.1	Esplin et al 2017	
vB_EamM_Huxley	<i>Erwinia amylovora</i>	Myoviridae	NA	240,761	271	9	Tree	NC_031127	Esplin et al 2017	
SPN3US	<i>Salmonella enterica</i>	Myoviridae	NA	240,413	264	2	Chicken feces	NC_027402	Lee et al 2011	
VP4B	<i>Vibrio harveyi</i>	Myoviridae	NA	236,053	212	NA	Ocean	KC131130.1	NA	
vB_EamM_Joad	<i>Erwinia amylovora</i>	Myoviridae	NA	235,374	245	NA	Tree	MF459647.1	Esplin et al 2017	
65	<i>Aeromonas salmonicida</i>	Myoviridae	NA	235,229	437	16	NA	NC_015251	Petrov et al 2010	
vB_EamM_RisingSun	<i>Erwinia amylovora</i>	Myoviridae	NA	235,108	243	NA	Tree	MF459646.1	Esplin et al 2017	
vb_AbatM_ME3	<i>Acinetobacter baumannii</i>	Myoviridae	NA	234,900	326	4	Wastewater	KU935715.1	Buttimer et al 2016	
Aehl	<i>Aeromonas hydrophila</i>	Myoviridae	NA	233,234	352	27	NA	NC_005260	Chow and Rouf 1983	
S-SSM7	<i>Synechococcus</i>	Myoviridae	NA	232,878	319	5	Seawater	NC_015287	Sullivan et al 2010	
Xoo-sp14	<i>Xanthomonas oryzae pv. oryzae</i>	Myoviridae	NA	232,104	251	0	Soil	MT939492	Nazir et al 2020	
CC2	<i>Aeromonas hydrophila</i>	Myoviridae	NA	231,743	427	9	Sewage	NC_019538	Shen et al 2012	
phiRSL I	<i>Ralstonia solanacearum</i>	Myoviridae	150	231,255	343	3	Soil	NC_010811	Yamada et al 2010	
vB_EamM_Phobos	<i>Erwinia amylovora</i>	Myoviridae	NA	229,501	247	NA	Tree	NC_031043	Esplin et al 2017	
ACG-2014f isolate Syn7803C90	<i>Synechococcus</i>	Myoviridae	NA	228,143	292	NA	Ocean	NC_026927	Gregory et al 2016	
phiAS5	<i>Aeromonas salmonicida</i>	Myoviridae	121 x 71	225,268	343	24	River water	NC_014636	Kim et al 2012	
CR5	<i>Cronobacter sakazakii</i>	Myoviridae	NA	223,989	231	NA	Soil	NC_021531	Lee et al 2016	
vB_EamM_EarPhillipIV	<i>Erwinia amylovora</i>	Myoviridae	NA	223,935	241	NA	Tree	NC_031007	Esplin et al 2017	
RS12	<i>Ralstonia solanacearum</i>	Myoviridae	NA	223,932	224	NA	Soil	NC_028950	Bhunchoth et al 2016	
Rogue	<i>Caulobacter crescentus</i>	Siphoviridae	205x60	223,720	350	23	Surface water	NC_019408	Meczker et al 2014	

(Continued)

Table 1 (Continued).

Phage	Host	Family	Viron Size (nm)		Genome Size (nt)	No. of ORFs	No. of tRNA	Sample Source	Accession No.	References
			Head	Tail						
RSFI	<i>Ralstonia solanacearum</i>	Myoviridae	NA	NA	222,888	230	NA	Soil	NC_028899	Bhunchoth et al 2016
PX29	<i>Aeromonas salmonicida</i>	Myoviridae	NA	NA	222,006	330	25	NA	NC_023688	Petrov et al 2010
SP-15	<i>Bacillus subtilis</i>	Myoviridae	NA	NA	221,908	317	NA	Soil	NC_031245	Taylor and Thorne 1963
Karma	<i>Caulobacter crescentus</i>	Siphoviridae	205x61	314	221,828	353	26	Surface water	NC_019410	Meczker et al 2014
PAU	<i>Sphingomonas paucimobilis</i>	Myoviridae	NA	NA	219,372	295	7	Silkworms	NC_019521	White and Suttle 2013
Swift	<i>Caulobacter crescentus</i>	Siphoviridae	219x63	295	219,216	343	27	Surface water	NC_019411	Meczker et al 2014
0305phi8-36	<i>Bacillus thuringiensis</i>	Myoviridae	95	486	218,948	246	0	NA	NC_009760	Serwer et al 2007
Magneto	<i>Caulobacter crescentus</i>	Siphoviridae	211x58	293	218,929	347	27	Surface water	NC_019407	Meczker et al 2014
PhiEaHI	<i>Erwinia amylovora</i>	Siphoviridae	NA	NA	218,339	241	NA	Aerial tissue	NC_023610	Meczker et al 2014
S-CAM7	<i>Synechococcus</i>	Siphoviridae	NA	NA	216,121	266	NA	NA	NC_031927	NA
phiCbK	<i>Caulobacter crescentus</i>	Siphoviridae	205x56	300	215,710	338	26	Surface water	NC_019405	Meczker et al 2014
EL	<i>Pseudomonas aeruginosa</i>	Myoviridae	140	200	211,215	201	NA	NA	NC_007623	Hertveldt et al 2005
S-SKS1	<i>Synechococcus</i>	Siphoviridae	NA	NA	208,007	281	11	Seawater	NC_020851	NA
phiN3	<i>Sinorhizobium</i>	Myoviridae	NA	NA	206,710	402	6	NA	NC_028945.1	NA
vB_EamM_Y3	<i>Erwinia amylovora</i>	Myoviridae	NA	NA	261,365	338	0	Soil	KY984068.1	Buttimer et al 2018
XacN1	<i>Xanthomonas citri</i>	Myoviridae	NA	NA	384,670	592	56	Soil	AP_018399.1	Yoshikawa et al 2018
Joad	<i>Erwinia</i>	Myoviridae	143	206	235,374	246	0	Apple blossom	MF459647.1	Arens et al 2018
RisingSun	<i>Erwinia</i>	Myoviridae	143	206	235,108	244	0	Apple blossom	MF459646.1	Arens et al 2018
Atu_ph07	<i>Agrobacterium tumefaciens</i>	Myoviridae	146x152	136	490,380	714	33	Soil	MF403008.1	Attai et al 2018
RP12	<i>Ralstonia solanacearum</i>	Myoviridae	120	180	279,845	289	1	Soil	AP017924.1	Matsui et al 2017
RP31	<i>Ralstonia solanacearum</i>	Myoviridae	120	180	276,958	287	1	Soil	AP017925.1	Matsui et al 2017

JA11	<i>Dickeya solani</i>	Myoviridae	NA	NA	255,356	325	0	Sewage	MH389777.1	Day et al 2018
JA13	<i>Dickeya solani</i>	Myoviridae	NA	NA	254,061	326	0	Sewage	MH460460.1	Day et al 2018
JA29	<i>Dickeya solani</i>	Myoviridae	NA	NA	253,323	324	0	Sewage	MH460461.1	Day et al 2018
JA33	<i>Dickeya solani</i>	Myoviridae	NA	NA	255,356	325	0	Sewage	MH460462.1	Day et al 2018
ADI	<i>Dickeya solani</i>	Myoviridae	NA	NA	261,658	332	0	Sewage	MH460463.1	Day et al 2018
Bonaishi	<i>Vibrio coralliilyticus</i>	Myoviridae	120	190	303,340	301	0	Seawater	MH595538.1	Jacquemot et al 2018
MJ3	<i>Pseudomonas aeruginosa</i>	Myoviridae	130	140	288,170	417	12	Equine livery yard	LR588166.1	Imam et al 2019
Pa21	<i>Pseudomonas syringae</i>	Myoviridae	NA	NA	305260	420	8	Kiwi orchard	MK552327.1	Frampton et al 2019
SPFM	<i>Salmonella</i> spp.	Myoviridae	Multiple	Multiple	233 to 242 kb	Multiple	Multiple	Multiple	Multiple	Thanki et al 2019
DM	<i>Erwinia amylovora</i>	Myoviridae	Multiple	Multiple	271 to 275kb	Multiple	Multiple	Multiple	Multiple	Sharma et al 2019
PTm1	<i>Tenacibaculum maritimum</i>	Myoviridae	120	150	224,680	308	NA	Seawater	AP019524.1	Kawato et al 2020
PTm5	<i>Tenacibaculum maritimum</i>	Myoviridae	NA	NA	226,876	306	NA	Seawater	AP019525.1	Kawato et al 2020
PA5oct	<i>Pseudomonas aeruginosa</i>	Myoviridae	131	136	286,783	465	12	Sewage	MK797984.1	Cédric Lood et al 2020

defensive mechanism of a host such as glycoside hydrolase, chitinase, and many others.²⁸

Majority of the phage genes are not well studied so far to comprehend their true and complete function. But some advanced research conducted in this field has unlocked unique and specific biological functions of cellular machinery including a conserved protein tubulin homolog which supplements a proteinaceous nucleus-like compartment that provides space and separates phage DNA.^{29,30} This arrangement facilitates proper phage and bacterial host infection and an optimal rate of reproduction. This spindle-like structure presents physical protection against the bacterial immune system and offers phage with a broad level of resistance.³¹

A few available reports showed that jumbo phages can synthesize their own NAD⁺ essential for the enzymes involved in phage DNA replication and regulation.^{32,33} Jumbo phages specify different diverse mechanisms like methyltransferases and incorporation of uracil instead of thymine in the genome that helps them to evade restriction attacks as a result of DNA modification. Furthermore, tRNAs help to get over the host defense mechanisms such as those by utilizing the endoRNases that create hindrance in translation by cleaving tRNAs.¹³

Translational Machinery

Growing shreds of evidence indicate that jumbo phages possess a large number of tRNA and the enzymes involved in the translation process.³⁴ tRNAs are stable molecules, and many of these phages retain more than one tRNA gene and can have up to 28 tRNAs (Table 1). The sequences of these tRNAs are different from host tRNAs. For instance, PhiAS5 holds 56 tRNA genes that could give rise to anticodon sequences of 16 variable amino acids.³⁵ During phage infection, host tRNA molecules are compromised, at this stage, translation is mediated by phage-encoded tRNAs. It is also known that some phages belong to the *Myoviridae* family encoding tRNA genes have broad host range.³⁶ Therefore, it is suggested that phages with a large number of tRNAs have a wide host range.³⁷ Phage XacN1 showed a wider host range by infecting nine *Xanthomonas citri* strains than the other phages that did not encode tRNAs like Siphovirus phage Cp1 and podovirus phage Cp2.³⁴

Another important enzyme tRNA synthetase was observed to be present in more than one jumbo phage like *Yersinia* phage ΦR1-37 and many others.³⁸ Different tRNAs in the phage genome are present to facilitate the

translational process. The tRNAs are especially abundant to regulate the translation of proteins that comprise structure and smooth the translational performance of phage-specific genes.³⁹ Mostly, the number of tRNAs increases as genome length becomes large.⁴⁰ Jumbo phages have an average of 15 tRNAs per genome which is distinct but related to their hosts. Generally, Jumbo phages encode a larger number of translation-related genes than those encoded by smaller phages.⁴¹ Therefore, numerous genes encoded in viral genomes appear to signify an increased level of independence on host translational machinery. These genes rectify and provide an alternative for host genes that are necessary for the life cycle, which display jumbo phages are usually not dependent upon host proteins which are usually required for smaller phage genomes.³¹ This kind of less dependence of jumbo phages on host bacteria proves their importance as a complete functional unit and opens up new avenues to gather more genetic information from bacteria through horizontal gene transfer.^{42,43}

Genetic Characterization

Giant phiKZ-like bacteriophages belong to myoviruses, which include *Pseudomonas aeruginosa* phiKZ, EL, OBP, and *Pseudomonas chlororaphis* 201phi2-1 that are found to have circular genomes.^{44,45} These genomes are packaged inside the capsid through a head-full packaging mechanism ensuring that the entire interior space of the viral protein head is filled with DNA in the course of the DNA packaging. After genome sequencing of the giant phages, it is concluded that their genomes encode structural proteins of capsid and tail, RNA polymerases,^{26,46} chaperonins,^{45,47} and inner body proteins.⁴⁸ Virus inner body possessed an internal proteinaceous structure, specific for jumbo phages.⁴⁹ Its function is to protect the DNA inside the capsid, helping in phage infection and form the depot of phage proteins. During infection to bacterial cell, some proteins are co-injected required to build the imperative machinery for the transcription of early genes.⁴⁸

Some members of this class of viruses which are diverse as far as their phylogenetic organization is concerned, usually can give rise to two enzymes. One is non-virion DNA-dependent RNA polymerase (RNAPs) and virion RNAP.^{9,48} Contrary to many known virus-encoded enzymes, the enzymes from the virus of this category consist of enzymes of multi-subunit nature and are related to cellular organisms. The Jumbo RNAPs usually do not

have conserved subunits needed for enzyme-oriented assembly as previously studied enzymes.⁵⁰ Their working mechanism is also different in various aspects. For instance, the promoter recognition steps are different from mechanisms usually adapted by various cellular enzymes. This could be explained by an example of a non-virion RNAP from phage AR9 which needs uracil on the promoter region to start promoter-oriented transcription originating from ssDNA.⁹ A common ancestor has been found in the case of multisubunit RNAP from jumbo phage which makes them a different subgroup among the much-stretched group of jumbo phages.⁵¹

Another jumbo phage Atu-ph07 has 714 ORFs in its genome. Among 714 ORFs 214 were hypothetical proteins and 390 ORFs found no homologs. Proteins with assigned functions based on similarity with conserved regions were found to be 110. The predicted proteins with assigned

functional annotations share similarities with different proteins from bacteria and bacteriophage at the sequence level.⁵² These genes with annotated functionality are found to give rise to various kinds of proteins. Some of these proteins are involved in nucleotide metabolism, while some of these are lysis proteins, structural and in the metabolism of tRNA, while some genes are found to encode proteins regulating the DNA replication (Figure 1). Further characterization and analysis of proteins discover the relation of proteins with those of 16T4 core proteins, which are homologous and similar to RAK2-like phages. Applying the tandem mass spectrometry, most structural proteins were confirmed experimentally and 112 more proteins were predicted of virion-associated type. The study concludes that phage is very lytic and the mode of its interaction with the host reflects application as a biocontrol agent.⁵²

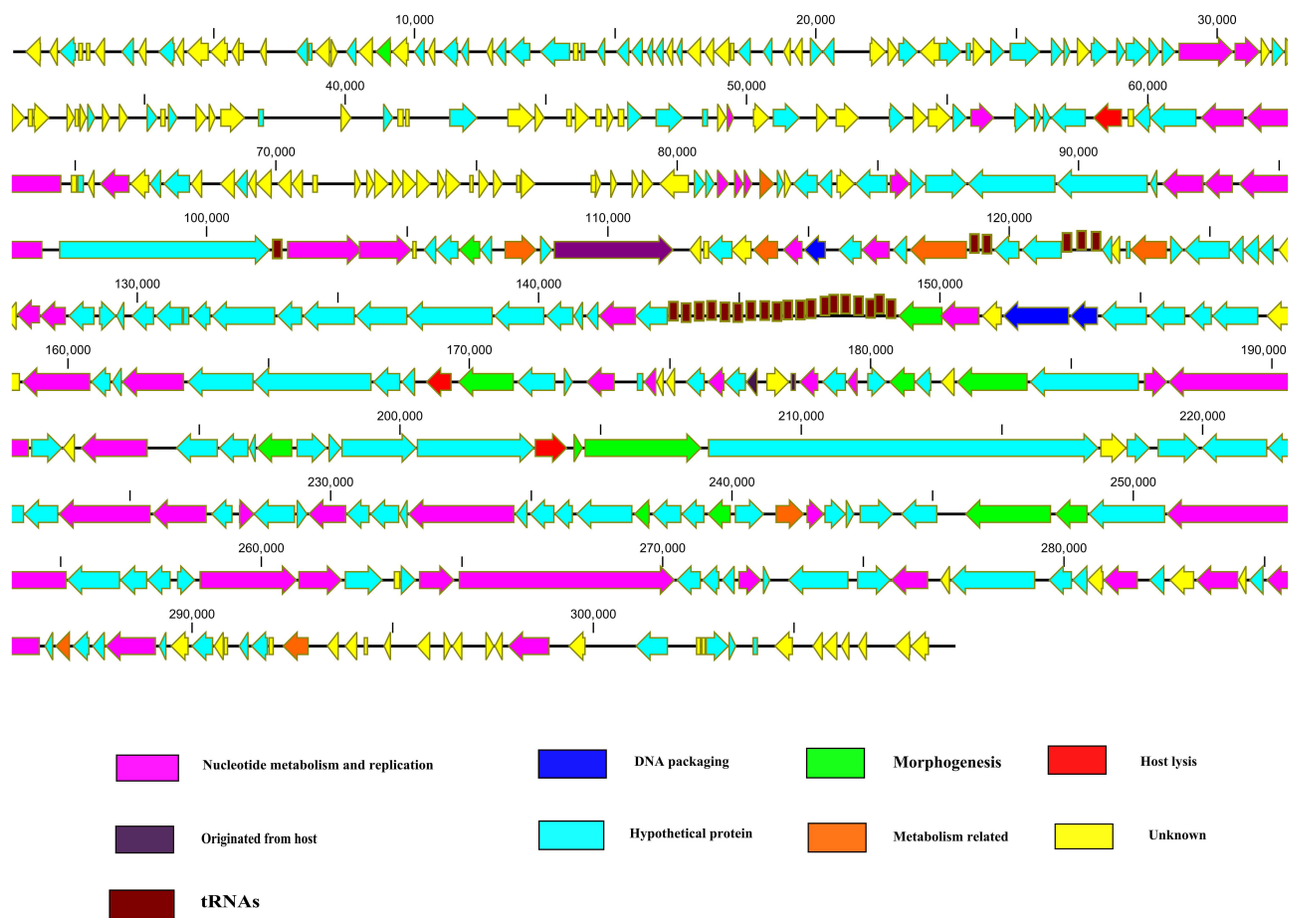


Figure 1 Schematic representation of the dsDNA genome of jumbo phages. Jumbo phage genes related to specific function scattered throughout the genome. Jumbo phages also contained a large number of tRNAs. Putative ORFs are presented as arrows, with predicted functions where available. Proposed modules are based on predicted functions. Turquoise, hypothetical protein; yellow, unknown; pink, nucleotide metabolism and replication; green, morphogenesis; red, lysis; blue, DNA packaging; brown, tRNA. The map was drawn with CLC Genomics main Workbench version 3.6.1 (CLC bio, Aarhus, Denmark).

Another study was conducted to characterize the novel jumbo phage which is lytic along with head fiber-like appendages. The full genome characterization of two phages (PTm5 and PTm1) of this type were found to be 226,876 and 224,680 bp in size respectively, with a genome compactness of about 29.7% and encoding 306 and 308 ORFs respectively. The sequences were found to be 99.5% identical between PTm5 and PTm1, highlighting the striking similarity between both of these phages.⁵³

The genetic similarity was observed in predicted ORFs between PTm1 and PTm5 belonging to *Tenacibaculum maritimum*, PT24 from *tenacibaculum*, and phage PAU from *Paucimobilis* till 15.0 to 16.6%. The close relation between these phages (PT24, PTm1, PAU, PTm5) was determined based on sequence similarity at the terminase large subunit genes through phylogenetic analysis. This phylogenetic analysis-based similarity was found to be more than those jumbo myoviruses which have equal genome sizes. Such genomes of equal size but differing at the phylogenetic level are ten in number.⁵³

A most recent study analyzed the genetic analysis of PALS2 and found its genome of 268,746 bp enabling it to be classified as a jumbo phage and its genome contained about 279 ORFs and 1 tRNA encoding asparagine while many of its predicted genes whose proteins cannot be assigned a function.⁵⁴

Gene Expression System

The smaller phages normally possess a modular genome structure, and genes having associated functions form clusters. However, the genes having alike functions in jumbo phage genomes are scattered in all over the genome (Figure 1).^{38,41,55} Efficient production of phage progeny required the timely expression of phage genes. Different phages adopt different strategies for the timely expression of genes. Similar to the small-genome phage, Φ KZ, a jumbo phage, genes are transcribed in a typical pattern, promptly by the phage-encoded RNAP.²⁶ Contrarily, the transcriptions of phage Φ R1-37 genes do not follow the typical pattern and expressed throughout the infection process by the phage-encoded RNAPs.²⁷ Notably, in both strategies, phage genes regulate under phage-encoded RNAPs instead of host RNAPs.

A research was conducted to find out some of the neglected, unexplored features of DNA transcription, recombination, and replication of jumbo phages.¹³ It also sheds light on the virion maturation system of jumbo viruses that infect bacteria. A new enzyme was found to

have an important role in controlling host cellular machinery and its orientation is protein modifying as far as its nature of action is concerned. Mechanisms were found to deal with the bacterial immune system by keeping the focus on host–virus interactions (R-M modification system, CRISPR–Cas systems).^{56,57} For example, effector activation is based on NAD and cyclic nucleotide mechanism of protection from superinfection during the process of pseudolysogeny. Overall, this research concludes that many proteins in jumbo viruses reflect the living systems observed in prehistoric replicons.¹³

According to Gaun (2020), most genes found in jumbo phages have not been studied well so far as their proper function is concerned. But now, some of the current research has highlighted the rare biological features which include conserved protein homologs which integrate a proteinaceous nucleus-like section that resides and separates phage DNA.¹³ Phages encoded a tubulin like protein PhuZ that possessed dynamic instability. The spindle of tubulin protein displays the instability at the dynamic level and positions the nucleus of phage within the bacterial host during infection of phage for efficient reproduction.³¹ The shell extends protection at the physical level for safety of phage genome from potential assault DNA targeting the bacterial immune system in this providing it huge resistance.³¹ CRISPR is an emerging tool for gene editing applied in broad fields and applications from biotechnology to medical-related fields. The CRISPR–Cas systems of jumbo phages can turn off genes encoding host transcription factors and translational genes. This mechanism of tuning the genes off offers a part of a larger interaction network that intercepts translation to redirect biosynthesis to phage-encoded functions.³² The CRISPR system enables bacteria to defend themselves from bacteriophage by presenting its adaptive immunity.¹³ But DNA modification and synthesis of anti CRISPR proteins coupled with some other mechanisms protect jumbo phages against the CRISPR–Cas system. A jumbo phage was found to be able to evade the CRISPR–Cas system.⁵⁸

Genetic analysis and characterization of PALS2 indicate the involvement of discovered genes in DNA repair, replication, metabolism of nucleotides, and multi-subunit RNA polymerase encoding genes, which is a genetic trait and a common characteristic of a jumbo virus. PALS2 was found to be like Φ KZ–like a virus in the result of comparative genomic analysis. The comparative genomic analysis also showed similarity to typical jumbo phages rather than staphylococcus phages.⁵⁴ The Cas protein from

bacteria is usually unable to approach virus nucleic acid during infection. But changing the location of an important restriction enzyme EcoR1 via engineering inside the proteinaceous compartment paves the way of targeting the virus and ensures the safety of the host cell. Φ KZ is also vulnerable to cas13a-CRISPR Cas enzyme that targets the virus nucleic acid. This study concludes that pseudomonas jumbo viruses defend themselves from quite a range of DNA targeting nucleases via making a protein barrier encapsulating their genomes.^{29,30,59,60}

Moreover, as being virulent, jumbo phages possessed some extra host lysis genes. They also possessed DNA replication, transcription, and nucleotide metabolism genes which help them to wider host range and reduce their dependence on the host.^{8,61–64} Jumbo phages should be comprehensively characterized first. Furthermore, a large number of proteins of jumbo phages are hypothetical with unclear functions. These proteins could have any side effects is not predicted yet.^{9,56,65} Therefore, the application of jumbo phages needs to reach a compromise between host specificity and optimal treatment.³¹ In summarizing the whole story, further research of jumbo phages will unveil more attractive characteristics that useful for phage biology. They are also enriched in evolutionary perspectives and intensify the phage usage in diversified applications.

Evolutionary Aspects

The evolutionary status of jumbo phages has not been understood in well manner because of non-availability of sufficient number of jumbo phages and their high level of diversity at genomic level. Till now, some phages like Φ KZ-like phages⁶⁶ and T4-like phages⁶⁷ are classified on the basis of morphological similarity and range of host. Moreover, no clear evidence has been observed for the classification of jumbo phages. Because of less similarity with already discovered phages, jumbo phages have been described as new genetic lineage. Yuan and Gao had classified 93 jumbo phages using amino acid sequences of terminase large subunits in five singletons and 11 clusters.⁸

This study performed a phylogenetic analysis of terminase large subunits of 152 jumbo phages and showed that these jumbo phages could be classified into 17 clusters and 8 singletons (Figure 2). As per phylogenetic analysis some phages that classified as Φ KZ including phage OBP and phage Lu11, now classified differently in this study.

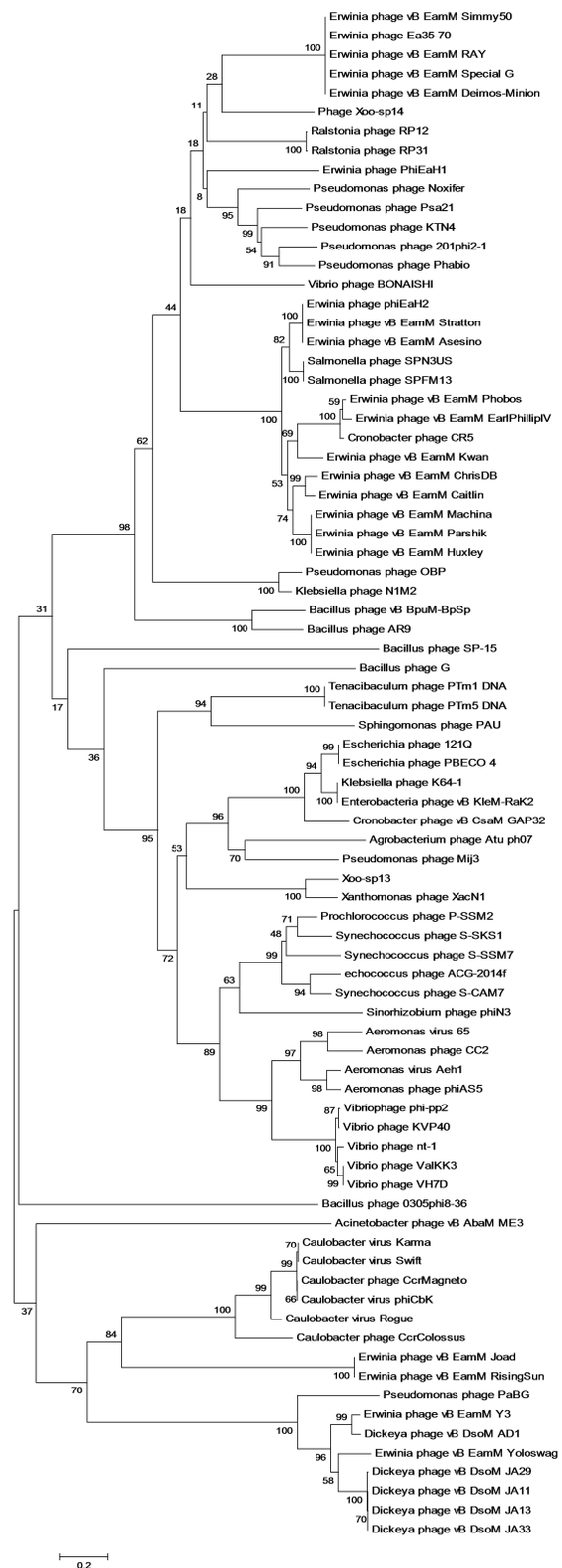


Figure 2 Phylogenetic analysis of jumbo phages. The evolutionary history was inferred using the neighbor-joining method. The evolutionary distances were computed using the Poisson correction method and are in the units of the number of amino acid substitutions per site. The analysis involved amino acid sequences of terminase large subunits from 152 jumbo phages. All positions containing gaps and missing data were eliminated. Evolutionary analyses were conducted in MEGA5.

According to core gene analysis of the jumbo phages, some phages should be classified differently as they previously classified as T4-like phages, eg Φ PAS5 which shares only 26% genes with T4-like phages and classified as T4-like phage. Contrarily, phage Φ PAS5 and Aeh1, which share 98% of their genes, have been classified in the same cluster in this study. The jumbo phages related to the same cluster tend to infect the host strains from similar genus and isolated from the same ecological environment. This study has assigned some phages to different clusters contrary to previous classification, on the basis of phylogenetic analysis.

The diverse foundation of jumbo phages is marked by very small level of similarity or no match at all among phages of different clusters. And phages from similar clusters are highly alike to each other exhibiting great degree or relatedness. As suggested in previous studies, evolution of jumbo phages might be a result of extracting novel genetic segments by smaller genome phages and increasing genome sizes over a certain period of time to give them their present shape.¹⁶ Essential genes imperative for phage life cycle are found in both small phages as well as jumbo phages as was concluded in the result of analysis of core genes.^{35,68} Genetic analysis of phage 0305 Φ 8-36 shed light on the origin that it has come into existence as the result of fusion of two viral genomes of ancestral nature by dint of the horizontal exchange of a genome module (block of genes) throughout the evolutionary course.⁶⁹ Whereas the main stream number of the jumbo phages are thought to get their genes from their respective hosts via horizontal gene transfer to formulate larger genomes.⁶⁵

Keeping the jumbo phages with unclear method of propagation on one side, many of the large dsDNA viruses comprising phycodnaviruses, asfarviruses, poxviruses, iridoviruses, and ascoviruses are characterized as nucleocytoplasmic large dsDNA viruses,⁷⁰ and gigantic sized viruses which normally infect amoeba, comprising *Mollivirus sibericum*, faustoviruses, mimiviruses, marseilleviruses, pandoraviruses, and pithoviruses.⁷¹ The replication cycles of aforementioned giant and large dsDNA viruses consist of the occurrence inside the host cytoplasm of viral factories which give rise to the progeny viruses.⁷² These viral factories were imagined to be the origin and starting point of the contemporary eukaryotic nucleus.⁷¹ Jumbo phages tend to display parallel replication physiognomies to the eukaryotic NCLDV. The protein of tubulin-like nature PhuZ of phage 20182-1 can arrange a spindle

apparatus and spot the phage genomic DNA to the mid-cell area of the bacterial host. Consequently, the encapsidated DNA arranges a rosette-like assembly encircled via a huge DNA-based mass, that looks alike to the viral factory of NCLDVs till some extent.⁷³ PhuZ related proteins have also been observed in the genomic structure of many jumbo phages coupled with phage genomes possessing near about 200 kbp. Evolutionary analysis of homologous proteins of PhuZ via genetic means marks similarity of jumbo phages with genome near 200 kbp, but is found to be different from the phages of small genome and the cellular microorganisms.⁸ Although phages having smaller genomes do not give rise to tubulin-like protein inside their own genomes, but they also facilitate engagement of the tubulin-like protein from the host bacteria to enable replication of the phage genome.⁷⁴ Development of viral factory-related assemblies via large viruses and jumbo phages paves the way for the creation of a podium to quintessence virus genomes, virus replication, host proteins required for replication, and associated proteins which assist in protecting viruses from host defenses,⁷² which in turn can smooth the process of virus propagation. Setting the features of forming viral factories aside, NCLDVs coupled with giant viruses from amoeba possess more genes affiliated with nucleotide metabolism, genome replication, and some other biochemical processes.⁷⁵ Even though NCLDVs, jumbo phages, and giant viruses of amoeba inclined to reflect many similar features, they are spaced far apart as far as evolutionary similarities are concerned.⁸ The jumbo phages have been observed more closely related to the archaea and bacteria, while the NCLDVs close evolutionary similarity is associated with the eukaryotes.^{76–80}

Conclusion

Isolation and discovery of jumbo viruses has greatly enriched our understanding of biological entity diversity and evolution. They have been isolated from diverse environments and showed high genetic diversity. However, genome sizes and their non-modular structures, scattered genes throughout the genome related to specific function, presence of RNAPs in phage virion that control the gene expression, and distance between jumbo phages and smaller phages are the prominent features that differentiate jumbo phages from the smaller-genome phages. Furthermore, genome similarity, infection and propagation mechanisms are more divergent characteristics that jumbo phages showed among each other. Several areas need to be

investigated deeply for a greater understanding of the jumbo phages. Although research showed that jumbo phages evolve from small phages but show bundle of differences. Mostly, functions of jumbo phages genes were predicted by using bioinformatics analysis. So, functional analysis of these genes will provide great understanding about phage interactions and evolution. Origin and evolution of jumbo phages need to be investigated more so that we can understand the origin and evolution of these cellular biological entities.

Author Contributions

All authors made substantial contributions to conception and design, acquisition of data, or analysis and interpretation of data; took part in drafting the article or revising it critically for important intellectual content; agreed to submit to the current journal; gave final approval of the version to be published; and agree to be accountable for all aspects of the work.

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Disclosure

The authors report no conflicts of interest for this work and declare that this research has been conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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