Sequence Analysis of *Leuconostoc mesenteroides* Bacteriophage Φ 1-A4 Isolated from an Industrial Vegetable Fermentation^{∇}

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Vegetable fermentations rely on the proper succession of a variety of lactic acid bacteria (LAB). Leuconostoc mesenteroides initiates fermentation. As fermentation proceeds, L. mesenteroides dies off and other LAB complete the fermentation. Phages infecting L. mesenteroides may significantly influence the die-off of L. mesenteroides. However, no L. mesenteroides phages have been previously genetically characterized. Knowledge of more phage genome sequences may provide new insights into phage genomics, phage evolution, and phage-host interactions. We have determined the complete genome sequence of L. mesenteroides phage Φ 1-A4, isolated from an industrial sauerkraut fermentation. The phage possesses a linear, double-stranded DNA genome consisting of 29,508 bp with a G+C content of 36%. Fifty open reading frames (ORFs) were predicted. Putative functions were assigned to 26 ORFs (52%), including 5 ORFs of structural proteins. The phage genome was modularly organized, containing DNA replication, DNA-packaging, head and tail morphogenesis, cell lysis, and DNA regulation/modification modules. In silico analyses showed that Φ 1-A4 is a unique lytic phage with a large-scale genome inversion (\sim 30% of the genome). The genome inversion encompassed the lysis module, part of the structural protein module, and a cos site. The endolysin gene was flanked by two holin genes. The tail morphogenesis module was interspersed with cell lysis genes and other genes with unknown functions. The predicted amino acid sequences of the phage proteins showed little similarity to other phages, but functional analyses showed that Φ 1-A4 clusters with several *Lactococcus* phages. To our knowledge, Φ 1-A4 is the first genetically characterized L. mesenteroides phage.

Bacteriophages are the most abundant biological entities (estimated to be on the order of $\geq 10^{31}$) on the planet (9, 18). Phages are ubiquitous in nature and can influence the microbial ecology and genetics of bacteria. Because of their small (usually <60 kb) genomes, phages can provide an excellent model system for studying many biological processes, including DNA replication and genetic evolution. Despite this, many phages remain uncharacterized. Very little is known about phage diversity and phage-host interactions owing to the small number of sequenced phages. Furthermore, the existing phage sequence database is highly biased toward a limited spectrum of phage hosts, namely, Enterobacteriaceae, Bacillus, Staphylococcus, Pseudomonas, Vibrio cholerae, Lactococcus, Streptococcus thermophilus, and S. pyogenes. The majority of host species for sequenced phages are either pathogenic or dairy-related bacteria. Most of the newly sequenced phage genes have no assigned functions or matches in the GenBank database (7).

Vegetable fermentations rely on a variety of lactic acid bacteria (LAB). The proper succession of LAB directly determines the quality and safety of the final fermentation products. *Leuconostoc mesenteroides* initiates most vegetable fermentations. It converts the sugars in vegetables (primarily glucose and fructose) to lactic acid, acetic acid, ethanol, CO₂, and other flavor compounds (22, 58, 59, 60, 61). Acid production lowers the pH of fermenting vegetables and inhibits the growth of many microorganisms, including pathogens. CO_2 production promotes the establishment of an anaerobic environment which favors the growth of other LAB. The metabolites produced by *L. mesenteroides* largely determine the flavor characteristics of the final products. As fermentation proceeds, *L. mesenteroides* rapidly dies off. Other LAB, including *Lactobacillus plantarum*, take over and complete the fermentation.

It has been a widely held view that the disappearance of L. *mesenteroides* and the subsequent bacterial succession in sauerkraut fermentations are due to the inhibitory effect of acids that accumulate during fermentation (54, 61). Little is known about other factors that may play a role in bacterial succession. Recent studies have shown that phages are present in the vegetable fermentations (4, 47, 48, 74, 75). Because of the rapid lytic cycle of these phages, they may significantly impact starter cultures and bacterial succession in vegetable fermentations (56). Phages active against *L. mesenteroides* have been isolated and characterized (48); however, genome sequences have not been reported.

L. mesenteroides phage 1-A4 (designated Φ 1-A4) is of particular interest. Φ 1-A4 is a lytic phage that was repeatedly isolated during the initial stages of a commercial sauerkraut fermentation. As a result, Φ 1-A4 may significantly influence the survival of L. mesenteroides and flavor development during sauerkraut fermentation. It was found that Φ 1-A4 infects at least three different strains of L. mesenteroides (48), and there-

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TABLE 1. Primer pairs used in PCR and the sizes of PCR products

Primer seque	Size (bp)		
Forward	Reverse	Predicted	Estimated
AAATATCAGTAGCAACTACACCCGACAAGTCC	ATGAAACGAGAACGACAGCAACGTCTTA	3,149	3,100
AATACACAAACACAATCATCAACCCAGTTAG	ATGAAACGAGAACGACAGCAACGTCTTA	1,340	1,350
AAATATCAGTAGCAACTACACCCGACAAGTCC	TGAAGAGATGTTCCGTAATGAGACTGGTCA	1,224	1,250
TCCCACAATTCAAGTTCATTAATACTATCAG	ATTGTTGTGTTCATTATAGGCAAATAGC	978	950
AGCAGAGTTCATTGCAGTTGAGTTTACAAAGG	ATCATTCAAAGCAACGGCGTTCTTATTACTC	910	900
TTCTCAGCGTACATAAACTTAAGAACTTGG	TACGTACCTTTGGTCTCTGGAATATTGTATC	886	875
AATACACAAACACAATCATCAACCCAGTTAG	TATTGTACGTGCCATATTATTAGCCGTAATCC	738	725

fore it may also promote genetic exchange and genetic diversity in microbial communities (34).

The objectives of this study were to determine and analyze the complete genome sequence of Φ 1-A4, to experimentally identify the structural protein genes, and to compare the genome organization with that of related phages. To our knowledge, this study represents the first complete genomic and molecular characterization of *Leuconostoc* phage. The results from this study may provide new insights into our understanding of phage genetics. This study may aid the development of phage control technologies in vegetable and other fermentations that are susceptible to phage attack.

MATERIALS AND METHODS

Bacterial strain, phage, and media. Phage Φ 1-A4 and host *L. mesenteroides* 1-A4 were both isolated on the first day of fermentation in a 90-ton industrial sauerkraut fermentation tank (48). The phage and its host were maintained as separate glycerol stocks at -80° C for later use. *L. mesenteroides* 1-A4 was cultured statically in de Man-Rogosa-Sharpe (MRS) broth (Difco Laboratories, Detroit, MI) at 30°C. Φ 1-A4 was propagated on *L. mesenteroides* 1-A4 in MRS broth supplemented with 10 mM CaCl₂ at 30°C.

Phage lysate preparation. Phage lysate was prepared as previously described by Lu et al. (46, 47). Briefly, early-log-phase host 1-A4 cells were inoculated with a Φ 1-A4 stock at a multiplicity of infection of about 0.01 and then incubated at 30° C until complete cell lysis occurred (in about 6 h). The lysate was centrifuged at $4,000 \times g$ for 10 min and filtered (0.22- μ m pore size). The filtrate was treated with DNase I and RNase A. Phage particles were concentrated by polyethylene glycol precipitation and then purified by cesium chloride density gradient ultracentrifugation at $600,000 \times g$ for 6 h at 15°C.

Phage DNA preparation. Phage DNA was prepared from the ultracentrifugepurified phage preparation by phenol-chloroform extraction as described previously (46, 47). Briefly, the CsCl-purified phage suspension was extracted with phenol and chloroform-isoamyl alcohol, followed by extraction with chloroformisoamyl alcohol. Phage DNA was precipitated with sodium acetate and ethanol and then pelleted with a microcentrifuge. After being washed with 70% ethanol, the DNA pellet was air dried and then resuspended in Tris-EDTA buffer.

Phage genome sequencing and verification of the assembly. Initial sequencing was done using plasmid libraries of short phage fragments. Φ 1-A4 DNA was digested with MseI, and the resulting fragments were ligated into pUC18. Recombinant clones (n = 24) were sequenced using vector primer sequences. The sequences of 184 fragments were used to design additional primers. Sequencing reactions were then performed directly on uncut phage DNA using Thermo-Fidelase and Fimer technology (63, 67). This resulted in one linear fragment representing the completed phage genome. Trace assembly was done with Phred/Phrap software (21). Selected genome regions were verified by PCR using the primers listed in Table 1.

DNA sequence and bioinformatic analyses. The finished phage genome sequence was subjected to an automated two-step annotation using GAMOLA software (global annotation of multiplexed on-site blasted DNA sequences engine [2]). First, the predicted open reading frames (ORFs) were analyzed using a gapped BLASTP (basic local alignment search tool for protein sequences) algorithm and the nonredundant protein database provided by the National Center for Biotechnology Information (November 2009). This revealed common unbiased protein similarities to all known protein sequences. Second, a customized database generated from known phage genomes (E. Altermann, unpublished data) was used to reveal phage-specific protein similarities. Additionally, the deduced amino acid sequences were analyzed by using the Pfam (Protein Family) (5), TIGRfam (The Institute for Genomic Research, family) (27), and COG (Clusters of Orthologous Groups of Proteins) (70) functional databases to detect conserved motifs in the proteins. The automated annotation was augmented by structural analyses including the prediction of Rho-independent terminator structures (20), transmembrane helices (41), and signal peptide sequences (6). Subsequently, a manual verification was performed based on the functional analyses, physical genome properties (e.g., the expected modular makeup of the phage genome), physicochemical attributes of the predicted ORFs, and their deduced amino acid sequences.

The Molecular BioComputing Suite (57) was used to calculate the molecular masses and isoelectric points of predicted proteins of Φ 1-A4.

Analysis of structural proteins. Sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) was performed as described by Lu et al. (46). Briefly, CsCl-purified phages in SDS-PAGE sample buffer were heated in a boiling water bath for 10 min and then applied to a NuPAGE precast gradient minigel (4 to 12% Bis-Tris; Invitrogen Corporation, Carlsbad, CA). After electrophoresis, the protein bands were transferred onto a polyvinylidene difluoride membrane (0.2- μ m pore size) and then stained with 0.1% Amido Black. Visible protein bands were excised, and their N-terminal amino acid sequences (10 or 11 residues) were determined by ProSeq, Inc. (Boxford, MA). Based on the N-terminal amino acid sequence of each protein, the corresponding ORF was identified in the Φ 1-A4 genome.

Functional analysis. A total of 112 phage genomes were analyzed, and a functional genome tree was generated using compACTor (Altermann, unpublished software) and Mega4 (69). The genomes were chosen from representative phages of hosts assigned to the phylogenetic class of bacilli in GenBank. Briefly, a unique genomic distance matrix was calculated based on round-robin ORF comparisons and subsequently imported into Mega4 for visualization. The functional distribution was visualized using the unweighted-pair group method using average linkages (UPGMA) (68).

Nucleotide sequence accession number. The final verified genome sequence and annotation of Φ 1-A4 were submitted to the publicly accessible GenBank database under accession number GQ451696.

RESULTS AND DISCUSSION

 Φ 1-A4 belongs to the *Siphoviridae* family in the order *Caudovirales* (Fig. 1). The average burst size of Φ 1-A4 is 24 particles per infective center. The latency period is about 30 min in MRS broth supplemented with 10 mM CaCl₂ at 30°C based on a one-step growth kinetics curve (56). In addition to infecting its natural host, *L. mesenteroides* 1-A4, Φ 1-A4 was also found to infect two other *L. mesenteroides* strains, 1-B8 and 1-E10 (48). To our knowledge, this is the first report of genomic sequencing and characterization of an *L. mesenteroides* phage at the molecular level.

Genomic features of Φ 1-A4. The genome sequence of Φ 1-A4 was determined in two steps. Plasmid libraries were used first to obtain initial phage sequences for primer design, and in the second step, several rounds of direct genomic sequencing were performed that yielded one linear contig (see

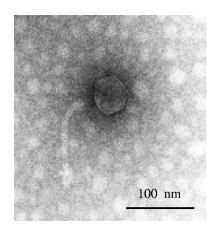


FIG. 1. Transmission electron micrograph of phage Φ 1-A4.

Materials and Methods for details). The resulting genome sequence and the predicted gene model were verified by PCR amplification for accuracy of sequence assembly. The measured sizes of PCR products agreed well with the corresponding predicted sizes (Table 1), which confirmed the unusual genome structure of Φ 1-A4. Furthermore, the repeats flanking the predicted origin of DNA replication were amplified and resequenced.

 Φ 1-A4 has a linear, double-stranded DNA genome consisting of 29,508 bp with a G+C content of 36%, which is very close to the G+C content (37%) of L. mesenteroides subsp. mesenteroides ATCC 8293 (53), which is closely related to the natural host of Φ 1-A4 (L. mesenteroides 1-A4). The origin of DNA replication and a cos site were identified in the genome (Fig. 2 and 3). Two adjacent runoff sequencing reactions obtained during Sanger sequencing of the phage genome identified the cos site boundaries (Fig. 3a and c). Manual enhancement of the electropherograms resulted in the identification of the 22-nucleotide (nt) cos site sequence (GGTTAATAGTAG TCTTTTTTAA) (Fig. 3b). The presence of a cos site indicated that the phage DNA may be circularized upon entry into the host. The origin of DNA replication was identified based on its similarities in structure to other phage replication origins (3, 38). Briefly, a noncoding region was identified harboring an AT-rich domain (50 to 60 nt, >95%) flanked by direct and indirect repeat structures. In Φ 1-A4, the noncoding region spans over 410 nt and contains a 55-nt AT-rich domain (98.19%) flanked by nine different types of direct and indirect repeats.

Bioinformatic analysis of the Φ 1-A4 genome revealed 50 possible ORFs. The ORFs are numbered consecutively starting from the ORF immediately downstream of the origin. Twenty-two ORFs were located on one strand, and 28 ORFs were on the complementary strand (Table 2 and Fig. 2). Of the 50 ORFs, 48 apparently initiated translation with the ATG start codon, 1 initiated with TTG, and 1 initiated with GTG (Table 2). Forty-six of the 50 ORFs were preceded by potential Shine-Dalgarno sequences (64) complementary to the 3' end of the 16S rRNA gene of *L. mesenteroides* ATCC 8293 (5'-C ACCTCCTTTCT-3'). The consensus sequence (AGGAGG) of Shine-Dalgarno elements is generally conserved and was chosen as the recognition sequence for ribosome binding site

(RBS) prediction (64). An *in silico* restriction site analysis of the nucleotide sequence agreed well with the experimentally determined restriction pattern (Fig. 4).

A total of 26 (52%) ORFs showed homology with previously characterized genes or were functionally annotated based on their respective positions in the phage genome. In many cases, homology to sequences of phages infecting gram-positive bacteria was found (Table 2). Twelve ORFs exhibited similarities to uncharacterized database entries (conserved hypotheticals), and 12 other ORFs had no homology to sequenced genes.

The phage genome was found to be modularly organized, consisting of DNA replication, DNA-packaging, head and tail morphogenesis, cell lysis, and gene regulation modules (Fig. 2). In silico analyses revealed that Φ 1-A4 contains a large-scale genome inversion ($\sim 30\%$ of the genome) compared to other phages, including the lysis module, part of the structural module, and the likely cos site. The predominant genome of phages with hosts phylogenetically closely related to Φ 1-A4 is linear, with nearly all of the genes located on the sense strand (for example, Leuconostoc phage KM20 and L. mesenteroides ATCC 8293 prophage, Fig. 5). The genome of Φ 1-A4 has a significant number of genes on the antisense strand, which is similar to the organization of Lactococcus phage genomes (Fig. 5). Large-scale genome inversions can significantly alter transcription patterns. We have identified putative promoter structures in the genome sequence of Φ 1-A4 that are likely to initiate the transcription of the inverted genome region (not shown). While we cannot exclude the possibility that the apparent inverted sequences are the result of recombination with another phage, we found no evidence to support this hypothesis. Both sections of the genome of Φ 1-A4 show the same GC content and codon usage. Further analyses will be carried out when the genome sequences of other *Leuconostoc* phages (or phages isolated from the same ecosystem) become available.

No evidence of a lysogeny module was found in the Φ 1-A4 genome. However, ORF22 exhibited very weak BLASTP similarities to the integrase subunit predicted in the *Solibacter usitatus* genome (GenBank accession no. CP000473). A twoiteration PSI-BLAST supported the initial finding. No significant functional Pfam or TIGRfam domains were identified. Only a very weak Pfam match (E value, 0.67) to SNF5, as well as a weak Pfam match to IN11, was detected. It is possible that ORF22 represents the remnants of an integration module or that it encodes a unique phage-specific protein of unknown function.

DNA replication module. Downstream of the putative origin of DNA replication (Fig. 2) are five genes (labeled in red) encoding proteins that may be involved in phage DNA replication. The predicted gene product of ORF1 (gpORF1) shows strong similarity to the putative DNA helicase of *Streptococcus pneumoniae* SP6-BS73 (Table 2) and that of *Lactobacillus reuteri* (data not shown). The derived protein product of ORF2 displays homology to the adenine-specific DNA methyltransferase from *Streptococcus* phage 9429.1. Adenine-specific DNA methyltransferase is often involved in chromosomal sitespecific DNA modification systems. In prokaryotes, the major role of DNA methylation is to protect host DNA against degradation by restriction enzymes (12). ORF3 encodes a protein homologous to various hypothetical bacterial proteins that have been shown to be bifunctional, having both DNA primase

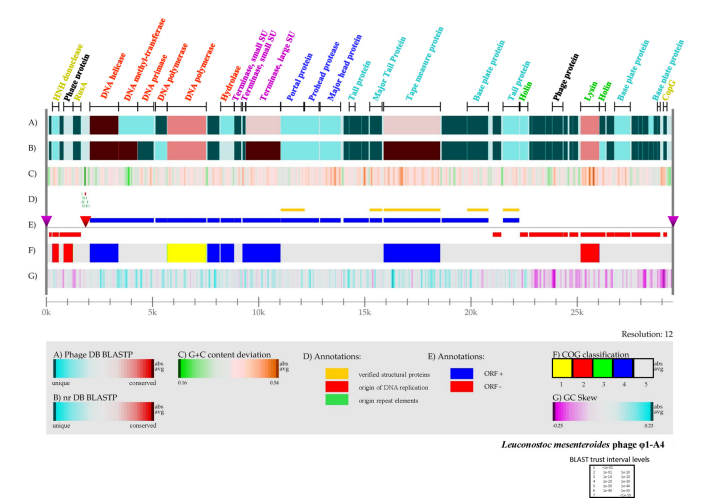


FIG. 2. Genome atlas of phage Φ 1-A4. The linear map was created using Genewiz, developed by Pederson et al. (60a; Center for Biological Sequence Analysis), and software developed in house. The legend at the bottom describes the individual panels corresponding to their respective designations (A to G). (A) Gapped BLASTP results obtained using the nonredundant database. (B) Gapped BLASTP results obtained using the custom phage genome database. In both panels A and B, blue regions represent unique proteins in Φ 1-A4 and highly conserved features are red. The color intensity corresponds to the level of similarity. Individual E values were grouped into trust interval levels as depicted in the legend. (C) G+C content deviation: green shading, low-GC regions; orange shading, high-GC islands. (D) Annotation shows the absolute positions of functional features as indicated. (E) ORF orientation. ORFs in the sense orientation (ORF +) are blue; ORFs oriented in the antisense direction (ORF –) are red. The origin of DNA replication is indicated by a red inverted triangle. The cos site, depicting the phage genome boundaries, is indicated by two purple inverted triangles on the genome line (gray line). (F) COG classification. COG families were assembled into five major groups: 1, information storage and processing; 2, cellular processes and signaling; 3, metabolism; 4, poorly characterized; 5, ORFs with uncharacterized COGs or no COG assignment. (G) GC skew. ORFs with predicted functional annotation are shown at the top of panel A with bars indicating their absolute sizes. The color coding corresponds to predicted functional phage modules. Red, DNA replication; purple, DNA packaging; dark blue, phage head; sky blue, tail structure; green, lysis; greenish yellow, DNA regulation and modification; black, not assigned to a specific phage module.

and polymerase activities (44). This suggests that the ORF3 protein is possibly involved in DNA synthesis. The predicted protein products of ORF4 and ORF5 exhibit sequence similarities to the hypothetical DNA polymerases from *L. mesenteroides* subsp. *cremoris* ATCC 19254 and to the putative DNA polymerase from *Listeria* phage P40 (Table 2) and *Lactobacillus* phage SalI (15), suggesting that these two proteins may be required for DNA replication.

The predicted product of ORF7 was found to be homologous to tRNA 3' endonucleases from *Listeria* phage P40 and *Lactobacillus delbrueckii* subsp. *bulgaricus* ATCC 11842. tRNA 3' endonuclease is a hydrolase that belongs to the metallo- β lactamase superfamily. Members of this family are ribonucleases that process the 3' end of tRNA precursors by cleaving the 3' trailing sequence, thereby preparing the pre-tRNA for the addition of a CCA tail (72).

DNA-packaging module. The protein specified by ORF10 exhibited similarity (Table 2) to a putative terminase large subunit from prophage lambda Sa2 harbored by *Streptococcus pneumoniae* SP14-BS69 (71). Terminases are enzymes involved in phage DNA packaging into proheads (11). These enzymes function as heteromultimers consisting of a large subunit and small subunits. In tailed phages, the small subunits of the terminase are responsible for specific DNA binding and the large subunit of the terminase mediates cleavage of the phage DNA into genome size units to be packaged into the prohead

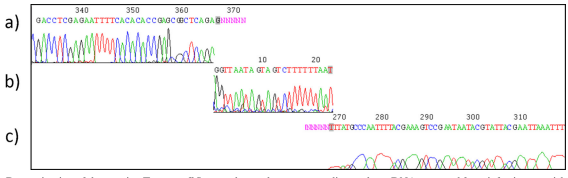


FIG. 3. Determination of the cos site. Two runoff Sanger electropherograms on linear phage DNA prepared from infectious particles identified the cos site boundaries. (a and c) Runoff electropherograms. (b) Manually enhanced traces identified the cos sequence. The figure is drawn to scale, and individual boxes are placed in relation to their expected sequence positions. Trace files were visualized using Chromas (http://www.technelysium.com.au/).

(19). Generally, the genes encoding terminase small subunits are located upstream of the terminase large subunit gene. Since the gene positions of ORF8 and ORF9 are upstream of the terminase large-subunit gene (Fig. 2), the products derived from these two ORFs are likely to be the small terminase subunits. Experimental data are needed to confirm if these two proteins are involved in phage DNA packaging.

Head and tail morphogenesis modules. Immediately downstream of the DNA-packaging module are the morphogenesis modules, including the head and tail modules. SDS-PAGE with a 4 to 12% Bis-Tris gradient gel revealed several structural protein bands (Fig. 6). Five bands were collected for the determination of N-terminal amino acid sequences (10 or 11 amino acids at the N terminus). Based on the N-terminal sequences, ORF11, ORF18, ORF19, ORF21, and ORF23 from the Φ 1-A4 genome were identified as the genes encoding the proteins extracted from the bands. An N-terminal methionine was absent from four of the five protein sequences (Fig. 6). Processing of the N-terminal methionine during protein maturation has been observed for many phages and occurs via host methionine aminopeptidase activity (45, 50).

The phage head morphogenesis module (dark blue in Fig. 2) includes ORFs encoding the experimentally determined minor head protein (also a putative portal protein), a putative prohead protease, and a putative head protein. The head genes are clustered together and precede the tail genes. The 42.9-kDa protein revealed on the SDS-PAGE (Fig. 6) has an N-terminal amino acid sequence (SQFDDINTRI) identical to residues 2 to 11 of the predicted gpORF11. This protein shows homology to the portal protein of *Escherichia coli* phage HK97 (35) (Table 2). Portal proteins typically function as dodecamers of a single polypeptide with an average molecular mass of 40 to 90 kDa. They form a hole (or portal) that enables DNA passage into the phage head during packaging (43).

gpORF12 showed similarities to the prohead proteases of the tailed phages of *Aggregatibacter aphrophilus* and *Burkholderia thailandensis* (Table 2) and, to a lesser degree, to the phage prohead peptidase of *Enterococcus faecium* phage HK97 (32). Prohead proteases are involved in the processing of the prohead protein (31). The prohead protease gene is generally located next to the gene for the prohead protein that is processed. In some cases, a prohead protease gene may be fused to the prohead protein gene. Immediately downstream of ORF12 is ORF13. gpORF13 showed homology to a major capsid protein of lactococcal phage Q54 (23). Based on the location of its gene in the genome, the putative major head protein is likely to be the protein processed by prohead protease. The cleavage of the head protein by the prohead protease generates the scaffold protein which aids prohead construction (28, 29).

Downstream of the head morphogenesis module is the tail morphogenesis module (sky blue in Fig. 2) including genes encoding a putative tail protein, four experimentally identified tail proteins (major tail protein, "tape measure" protein [TMP], baseplate protein, and receptor binding protein), and two putative baseplate proteins.

The product of ORF15 was found to have some similarity (E value, 1.2) to a tail protein of *Lactococcus lactis* phage TP901-1 (IPR011855) in the TIGRfam database. ORF15 is located between the major head and major tail genes (Fig. 2). In general, the genes located in this region are involved in the formation and connection of the head and tail structures and in DNA packaging (8).

The smallest structural protein observed by SDS-PAGE (Fig. 6) had a predicted molecular mass of 21.3 kDa. This protein appears to be the major phage protein, based on the amount of the protein observed in the SDS-PAGE gel. This protein is encoded in the tail protein gene region (Fig. 2) and displayed homology (although it was weak) to a phage major tail protein from the Phi13 family of *Streptococcus pneumoniae* CDC1873-00 (data not shown). The N-terminal sequence (M YDTRKITHGS) of the protein was identical to predicted residues 1 to 11 of the Φ 1-A4 gpORF18.

A protein with a predicted molecular mass of 91 kDa was the largest structural protein observed on SDS-PAGE (Fig. 6). The N-terminal amino acid sequence of the 91-kDa protein was ANTTSYMLKF, matching predicted amino acids 2 through 11 of the protein encoded by ORF19. The protein sequence was similar to that of the Mu-like phage minor tail protein in *Bifidobacterium longum* DJO10A (42) (Table 2). In many Mu-like phages, the gene(s) encoding tail fibers is relatively long. The predicted protein product of ORF19 contains a TMP domain that is only found in phages and prophages. These types of proteins determine phage tail length. In phage

TABLE 2. ORFs and genetic features of L. mesenteroides Φ 1-A4^a

ORF ^b	Putative RBS ^e					Database search results ⁱ for predicted product							
	Start ^c	End ^d	3'-AGGAGG	Start codon	Size (aa)	Mass ^g (kDa)	pI^h	Predicted function ^{<i>j</i>}	Organism matched	E value ^k	Reference ¹		
1 2	361 1710	1713 2612	AGGAGG	ATG ATG	450 300	51.95 35.00	5.22 7.81	DNA helicase Phage-related adenine-specific DNA methyltransferase	S. pneumoniae SP6-BS73 Streptococcus phage 9429.1	1e-52 8e-86	ZP_01821431.1 YP_596295.1		
3 4	2614 3448	3369 3996	TGGTGAA AGGAGAA	ATG ATG	251 182	29.07 20.80	5.43 8.67	DNA primase/polymerase DNA polymerase	L. mesenteroides subsp.	3e-78	ZP_03913787.1		
5	4005	5840	ACGAGAA	ATG	611	69.27	6.39	DNA polymerase	cremoris ATCC 19254 Listeria phage P40	4e-42	YP_002261447.1		
6 7	5896 6521	6468 7153	AGGAGAA GGGAGGA	ATG	190 210	21.63 24.01	4.93	Cons. hypo.	Listeria phogo P40	80.19	VB 002261444 1		
8	7164	7490	AGGAGG	ATG ATG	108	12.40	5.57 4.81	Hydrolase (tRNA 3' endonuclease) Terminase small subunit,	Listeria phage P40	8e-18	YP_002261444.1		
9	7550				50			deduced from position					
9 10	7552 7694	7710 9340	AGGAGG	ATG ATG	52 548	5.94 62.91	11.62 5.02	Terminase small subunit, deduced from position Terminase large subunit	S. pneumoniae prophage	2e-72	ZP_01828913.1		
								-	lambda Sa2		-		
11	9353	10474	AGGAGAA	ATG	373	42.91	4.89	Portal protein, amino acid sequence verified	E. coli phage HK97	4.5e-4	This work and NP_037699		
12	10434	11153	AGGAGAA	ATG	239	26.18	4.43	Phage prohead protease	A. aphrophilus NJ8700 caudovirus phage	1e-04	YP_003006998.1		
13	11205	12170	AGGAGAA	ATG	321	34.98	4.89	Major capsid protein	Lactococcus phage Q54	2e-04	YP_762590.1		
14	12310	12582	AGGAGGTG	ATG	90	10.35	4.33	Cons. hypo.					
15	12572	12850	AGGAGG	ATG	92	10.59	10.06	Put. phage tail protein					
16 17	12850 13164	13167 13493	GGGAGG AGGTGT	ATG ATG	105 109	12.39 12.32	4.44 10.92	Cons. hypo. Cons. hypo.					
18	13542	14123	AGGAGAA	ATG	109	21.25	4.55	Major tail protein, amino acid sequence verified			This work		
19	14197	16866	AGGAAATG	ATG	889	90.98	5.34	Phage tail TMP, amino acid sequence verified	B. longum DJO10A	1e-74	This work and YP_001955090.3		
20	16930	18126	TGGAGGA	ATG	398	45.48	4.71	Cons. hypo.			-		
21	18129	19127	AGGAGA	ATG	332	36.69	4.49	Baseplate, partial amino acid sequence match			This work		
22 23	C19335 19816	C19736 20583	AGGAGAA AGGAGA	ATG ATG	133 255	15.74 27.84	4.75 5.17	Cons. hypo. Tail protein, receptor binding, amino acid sequence verified	Oenococcus oeni phage fOg44	0.015	This work and CAD19145.1		
24	C20616	C20987	AGGAGA	ATG	123	14.00	5.72	Holin	G. adiacens ATCC 49175	7e-07	ZP_05738230.1		
25	C21054	C21188	AGGAGA	ATG	44	5.50	10.37	Cons. hypo.	0. <i>uuueens</i> 11100 19175	10 01	21_00700200.1		
26	C21185	C21349	TGGAGA	ATG	54	6.39	8.34	Unknown					
27	C21349	C21579	TTGAGG	ATG	76	9.01	8.94	Cons. hypo.					
28	C21579	C21815	AGGAAAA	ATG	78	9.18	6.31	Unknown					
29	C21805	C22020	ATAGGA	ATG	71	8.93	6.94	Unknown					
30 31	C22020 C22139	C22142 C22639	GAGAGG AGGAGAA	ATG ATG	40 166	4.43 18.81	9.24 9.38	Unknown Phage-related hydrogenase	L. mesenteroides subsp. mesenteroides ATCC 8293	1e-46	YP_819431.1		
32	C22660	C22869	AGGAGAA	ATG	69	8.14	4.10	Unknown					
33	C22945	C23115	AAGAGGA	ATG	56	6.09	8.60	Unknown					
34	C23112	C23357	AGGAGAA	ATG	81	9.76	5.88	Unknown					
35	C23473	C24354	AGGAGGA	ATG	293	31.60	5.92	Phage-related amidase, lysin	E. faecalis phage phiEF24C	4e-41	YP_001504118.1		
36	C24356	C24640	TGGTGG	GTG	94	10.61	8.94	Holin					
37	C24696	C25061	AGGAGA	ATG	121	13.14	4.13	Cons. hypo.	I lastia aboos alt	0.022	ND 605072.1		
38 39	C25073 C25873	C25807 C26190	AGGAGGA AGGAGG	ATG ATG	244 105	26.08 11.91	5.85 9.11	Baseplate Unknown	L. lactis phage r1t	0.022	NP_695073.1		
39 40	C25875 C26192	C26190 C26401	AUUAUU	TTG	69	8.04	4.93	Unknown					
41	C26395	C26670	AGGAAGTG	ATG	91	10.63	4.59	Cons. hypo.					
42	C26667	C26918	AGGAGA	ATG	83	9.71	4.33	Cons. hypo.					
43	C26911	C27099	AGGAGA	ATG	62	7.58	6.94	Unknown					
44	C27086	C27220	GGGAGAA	ATG	44	5.35	4.66	Put. baseplate		0.00	TTD 010000 1		
45	C27368	C27538	AGGAAA	ATG	56	6.50	5.31	Repressor, CopG	Nocardioides sp. strain JS614	0.26	YP_919229.1		
46 47	C27947 C28103	C28066 C28405	AGGATA	ATG ATG	39 100	4.76 11.64	10.38 7.84	Unknown Phage HNH endonuclease	L. casei prophage lambda Ba04	4e-07	YP_001987049.1		
48	C28443	C28634	AGGAGAA	ATG	63	7.47	4.06	Unknown					
49	C28636	C29076	AGGAGC	ATG	146	16.95	8.66	Phage-related protein Endodeoxyribonuclease RusA	E. faecalis phage phiEF24C	3e-12	YP_001504167.1		
50	C29073	C29447	AGGAGGA	ATG	124	14.87	8.60						

^{*a*} See the text for details.

^a See the text for details.
^b ORFs were numbered sequentially starting from the ORF immediately downstream of the predicted origin of DNA replication.
^c Genes located on the complementary strand are indicated by the letter C.
^d The end position includes the stop codon.
^e The sequence shown matches at least 60% of the consensus RBS (AGGAGG) sequence.
^f RBS consensus sequence identified downstream of promoters.
^g Molecular masses were calculated with the Molecular Biocomputing Suite (57).
^h Isoelectric points were calculated with the Molecular Biocomputing Suite.
ⁱ Database searches based on homologies of deduced amino acid sequences to the nonredundant amino acid database provided by NCBI were performed with the gapped BlastP algorithm. The best phage-related hits are included here.
^j Abbreviations: Cons. hypo., conserved hypothetical: Put., putative.

⁴Abbreviations: Cons. hypo., conserved hypothetical; Put., putative. ^k Probabilities derived from BLAST scores for obtaining a match by chance.

¹ Accession number or source.

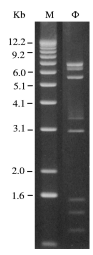


FIG. 4. HindIII restriction digestion analysis of Φ 1-A4 DNA. Lane M, 1-kb DNA ladder; lane Φ , Φ 1-A4.

lambda, the TMP is used as a template for tail polymerization and remains inside the tail tube (37). The length of the phage tail is directly proportional to the size of the TMP (36). TMPs are found in almost all *Siphoviridae* phages (62). Like most TMPs (800 to 5,006 amino acids long [55]), the 91-kDa protein (889 amino acids long) of Φ 1-A4 is the largest predicted protein in the Φ 1-A4 genome (Table 2) and is encoded in the putative tail protein gene region (Fig. 2). These features suggested that this protein is most likely to be a TMP.

The N-terminal amino acid sequence (ADTRTIYLHT) determined for the 36.7-kDa protein (Fig. 6) partially matched amino acids 2 through 11 (ADTSSIYLHT) of gpORF21. Residues 4 and 5 of gpORF21 were SS instead of RT. This result is possibly due to miscalled amino acids during the sequencing of this low-abundance protein. The protein appears to be a baseplate protein, based on its weak TIGRfam match to the baseplate assembly protein in *L. lactis* phage P2 (IPR013046). Kondou et al. (40) showed that baseplate protein is essential for phage Mu assembly and the generation of viable phages.

The N-terminal sequence of the 27.8-kDa protein (Fig. 6) is TLNTNEIVYT, matching predicted amino acids 2 through 11 of gpORF23. This protein may be responsible for host specificity because of the similarity (found by PSI-BLAST analysis) to a receptor binding protein in *Lactococcus* phage Q41 (accession number AAT81505.1). The protein also displayed sequence similarity to a hypothetical protein (ORF232) of *Oenococcus oeni* phage fOg44 (Table 2). This may suggest that the hypothetical protein of phage fOg44 is also involved in host range determination.

Figure 2 shows that the putative tail morphogenesis module was interspersed with ORFs belonging to the cell lysis module and other ORFs with unknown functions. This is not usually observed in other phages. This may have resulted from the large genome inversion as described above. Two ORFs, ORF38 and ORF44, are downstream of the lysis module and possibly encode baseplate proteins. The product of ORF38 displays some similarity to the structural protein from *L. lactis* phage r1t (73) (Table 2). It also shows weak similarity to the receptor binding protein from *Lactococcus* phage TP901-1 and to the putative tail-host specificity protein in *Lactococcus* phage P335 (E value, 1.4). Receptor binding proteins or tail-host specificity proteins are usually located on the baseplate. Based on Pfam database analysis, gpORF44 showed weak similarity to the baseplate structural protein domain.

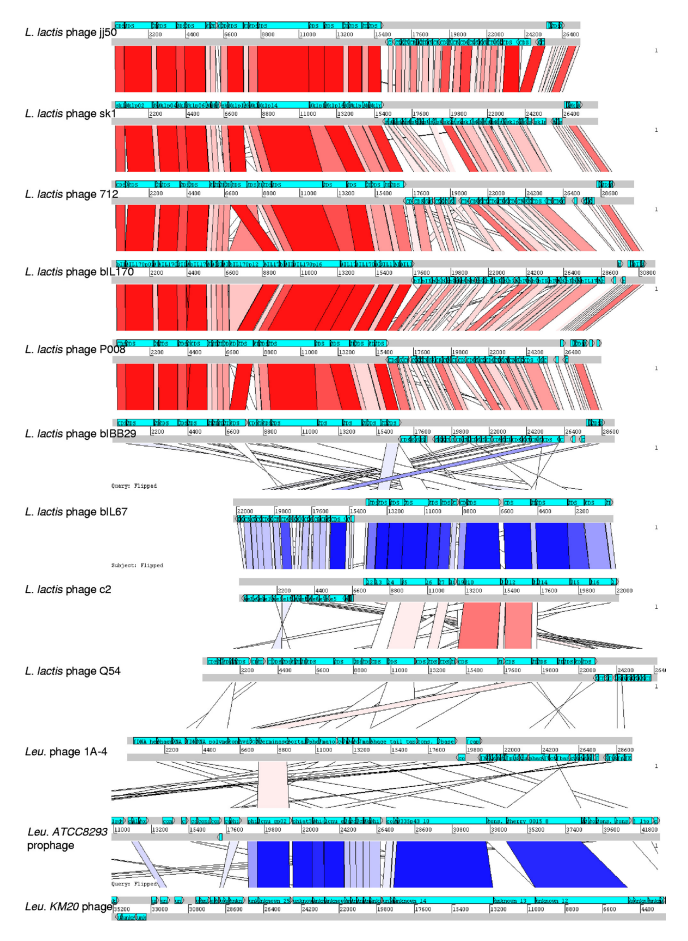
Host cell lysis module. The cell lysis module consists of ORF24, ORF35, and ORF36, encoding a putative endolysin and two holins, respectively (Table 2). In double-stranded DNA phages, the holin-lysin dual-lysis system is responsible for bacterial lysis and phage progeny release (17). Unexpectedly, this module is embedded in the tail morphogenesis module.

The predicted protein from ORF35 shows a strong sequence similarity to the putative *N*-acetylmuramoyl-L-alanine amidase of *Enterococcus faecalis* phage phiEF24C (Table 2). The protein appears to be a phage endolysin belonging to the amidase family. The amidase family includes zinc amidases that have *N*-acetylmuramoyl-L-alanine amidase activity. This enzyme domain cleaves the amide bond between *N*-acetylmuramoyl and L-amino acids (preferentially D-lactyl–L-Ala) in bacterial cell walls. The structure is known for phage T7 and indicates that two of the conserved histidines are zinc binding (13). Surprisingly, the putative endolysin gene in Φ 1-A4 was flanked by two predicted holin genes (ORF24 upstream and ORF36 downstream, Fig. 2).

The protein specified by ORF36 displays similarity to a hypothetical protein from Leuconostoc citreum KM20. The protein contains a consensus sequence (through hidden Markov model sequence analysis) which represents a putative phage holin from a number of phage and prophage regions of grampositive bacteria. Like other holins, gpORF36 is small (94 amino acids) with stretches of hydrophobic sequence and is encoded adjacent to the endolysin gene. The derived protein from ORF24 shows homology with the product of the holin gene which was found in the genomes of Granulicatella adiacens ATCC 49175 (Table 2) and Streptococcus pyogenes MGAS10394. However, unlike other holin genes, ORF24 is not adjacent to an endolysin gene. This has not been observed in other phage genomes. It is noteworthy that gpORF36 was not found to contain transmembrane domains, while gpORF24 had three transmembrane domains, similar to other holins (76). Further in vitro experiments are needed to elucidate whether ORF24 or ORF36 (or both) encodes a functional holin.

DNA regulation and modification module. The regulation and modification module contains ORF45, ORF47, and ORF50 (greenish color in Fig. 2). ORF45 encodes a protein that shows weak similarity to the transcriptional repressor, CopG, found in Nocardioides sp. JS614 (Table 2). The putative function of ORF45 as a regulator of transcription was supported by COG and Pfam database matches (data not shown). CopG (also known as RepA) is responsible for the regulation of plasmid copy number in Nocardioides. It binds to the repAB promoter and controls the synthesis of the plasmid replication initiator protein RepB. Many bacterial proteins that regulate transcription bind DNA through a helix-turn-helix (HTH) motif. CopG displays a fully defined HTH motif structure; however, it is not involved in DNA binding but is apparently responsible for the maintenance of the intrinsic dimeric functional structure and cooperativity (1, 25).

The product of ORF47 shows similarity to the HNH endo-



kDa 97 —	M	Φ	MW (kDa) Predicted		N-terminal sequence	ORF matched	aa position of matching ORF	Proposed function	
66 -			← 9	1.0	ANTTSYMLKF	ORF19	2-11	Tape measure protein	
55 —	-								
		-	← 42	2.9	SQFDDINTRI	ORF11	2-11	Portal protein	
36 _	_		← 30	6.7	ADTRTIYLHT	ORF21	2-11	Base plate protein	
31 -	-	inter la	← 2 ⁷	7.8	TLNTNEIVYT	ORF23	2-11	Receptor binding protein	
21 —	-'	-	← 2	1.3	MYDTRKITHGS	ORF18	1-11	Major tail protein	
14 —	-								

FIG. 6. Analysis of phage Φ 1-A4 structural proteins. After the purification of Φ 1-A4 by CsCl density gradient centrifugation, the phagecontaining band was analyzed by 4 to 12% SDS-PAGE. Lane M, molecular mass marker; lane Φ , proteins from Φ 1-A4. The N-terminal amino acid sequences of the five structural proteins were determined. The position of the N-terminal sequence in the corresponding Φ 1-A4 ORF and its proposed function are indicated.

nuclease of prophage lambda Ba04 in *Lactobacillus casei* BL23 (Table 2) and prophage L54a in *Staphylococcus aureus* subsp. *aureus* COL (24). The HNH motif was originally identified in the subfamily of HNH homing endonucleases, which initiate the process of the insertion of mobile genetic elements into specific sites (14). HNH endonucleases cleave phosphodiester bonds and are found in bacteria and viruses (16, 26, 66). This family includes pyocins, colicins, and anaredoxins.

Based on the Pfam database analysis, gpORF44 showed similarity (E value, 7.3e-04) to the endodeoxyribonuclease RusA. RusA functions as a Holliday junction resolvase (key enzyme in DNA recombination). The RusA protein of *E. coli* can resolve Holliday intermediates and correct the defects in genetic recombination and DNA repair associated with inactivation of RuvAB or RuvC (51).

Functional analysis. A functional distribution analysis of multiple genome sequences was performed using compACTor. The functional genome tree (Fig. 7) was constructed based on the ORFs from 112 phages. The hosts for the 112 phages include lactobacilli, lactococci, streptococci, enterococci, listeriae, bacilli, and clostridia. Φ 1-A4 clusters with several *Lactococcus* phages (functional cluster I [FC-I], 49.9 distance units [du]) and was part of another node in a separate group containing predominantly *Bacillus* phages (FC-II, 56.07 du, Fig. 7). Within FC-I, four separate groups could be observed (17.6, 18.6, and 49.9 du apart) and individual groups appeared to be tightly clustered.

As previously described in the literature, *L. lactis* 936-type phages show extensive genetic similarities, and it has been speculated that these phages have evolved from the same ancestral core genome (30, 52). The slight differences (0.47 du) within this subgroup supported the notion of a group of highly related phages. Based on the compACTor analysis, levels of

amino acid sequence similarities within FC-I were visualized in ACT (10) (Fig. 5). As expected, 936-type phages revealed a high level of sequence similarity and gene synteny in their late-gene modules (packaging, morphogenesis, and lysis); however, a greater level of diversity was observed for early and middle genes (replication).

A second group within FC-I (0.42 du, Fig. 7) comprised prolate-head c2-like lactococcal phages (49). Both phages c2 (33) and bIL67 (65) showed extensive genome similarity (Fig. 5). Interestingly, the c2-like group exhibited similarity in their structural module to L. lactis phage Q54 (23). In contrast, Q54 revealed a weaker similarity and gene synteny to 936-type phages (data not shown). The apparent differences in the ORFs of Q54 from 936- and c2-like phages were also illustrated by a functional distance of 18.6 du. With a distance of 49.9 du, Leuconostoc phage Φ 1-A4 represented a distinct member of FC-I. Only few ORFs showed amino acid sequence similarities to the other three subgroups (Fig. 5). Based on the similar functional distances between FC-I, FC-II, and Φ 1-A4 of 56.07 and 49.9 du, respectively, it is tempting to speculate that Φ 1-A4 might represent a separate functional cluster (FC-III) within this larger node.

Interestingly, Φ 1-A4 did not cluster with the two prophages harbored by *L. mesenteroides* ATCC 8293 (53) and *L. citreum* KM20 (39), respectively (Fig. 7). Instead, Φ 1-A4 clustered with several *Lactococcus* phages. The two *Leuconostoc* prophages show significant sequence similarity and gene synteny within their structural modules. They formed a distinct functional cluster (Fig. 7), and they were grouped together with several *Lactobacillus* phages. In contrast, Φ 1-A4 only displayed similarity to the predicted large terminase subunits of the two other phages (Fig. 5).

Overall, extensive genetic shuffling is apparent among the

FIG. 5. pACT alignment of Φ 1-A4 against phages of its functional cluster (F-Cluster 1 in Fig. 7), as well as against *Leuconostoc* phage KM20 and *L. mesenteroides* ATCC 8293 prophage. Based on the FGD algorithm, predicted phage ORFeomes were subjected to an all-versus-all analysis. Results were stored in pMSP data files (http://www.sanger.ac.uk/Software/ACT/v7/manual/start.html#COMPARISON-FILE-FORMAT) and visualized using the Artemis comparison tool (10). Red lines show direct similarities between amino acid sequences of predicted ORFs, and blue lines represent inverted similarity hits. Color intensities are proportional to the respective increasing levels of sequence similarity. Phage genomes are represented by two gray lines (showing sense and antisense strands), and light blue arrowed boxes indicate individual predicted ORFs, their relative lengths, and their orientations. Genomes are drawn to scale.

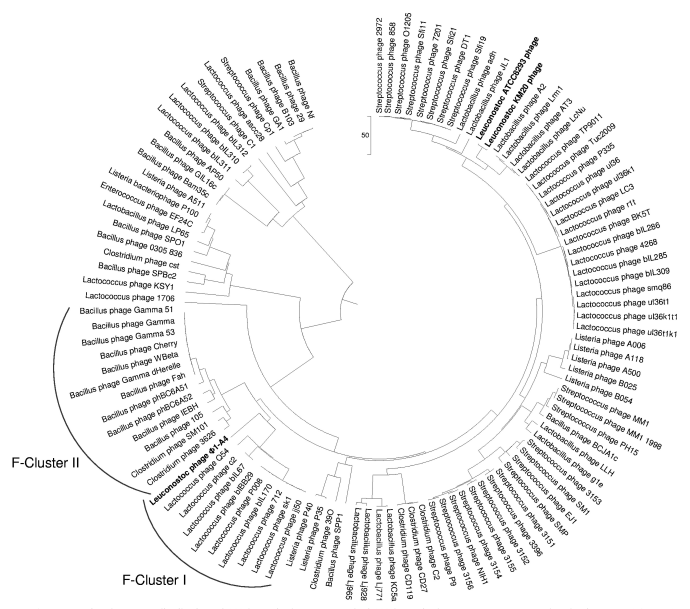


FIG. 7. Functional genome distribution of 112 bacteriophages. A total of 112 bacteriophage genomes were analyzed using compACTor. A distance matrix was calculated and subsequently imported into Mega4. The functional history was inferred using the UPGMA. The optimal tree with a branch length sum of 2,115.25304686 is shown. The tree is drawn to scale.

phages from different hosts, supporting the widely held view that tailed phages are genetic mosaics arising by the exchange of functional modules within a diverse genetic pool. The weak amino acid similarities among these phages are indicated by the long branches (\sim 200 du).

Conclusion. To our knowledge, this study represents the first complete genome sequence and genetic characterization of an *L. mesenteroides* phage. Bioinformatic analysis revealed that Φ 1-A4 is a unique lytic phage compared to phages infecting related species of LAB. Approximately 30% of the Φ 1-A4 genome is inverted compared to the genome organization of other sequenced phages. The endolysin gene was flanked by two holin genes. The tail morphogenesis module was interspersed with cell lysis genes. The overall amino acid sequences

of the phage proteins had little similarity to other sequenced phages. Functional analyses showed that Φ 1-A4 clusters with several *Lactococcus* phages. The results of this study may provide new insights that deepen our understanding of phage genetics and phage-host interactions in dynamic ecosystem such as vegetable fermentations.

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