

# Characterization of Two Virulent Phages of Lactobacillus plantarum

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We characterized two *Lactobacillus plantarum* virulent siphophages, ATCC 8014-B1 (B1) and ATCC 8014-B2 (B2), previously isolated from corn silage and anaerobic sewage sludge, respectively. Phage B2 infected two of the eight *L. plantarum* strains tested, while phage B1 infected three. Phage adsorption was highly variable depending on the strain used. Phage defense systems were found in at least two *L. plantarum* strains, LMG9211 and WCSF1. The linear double-stranded DNA genome of the *pac*-type phage B1 had 38,002 bp, a G+C content of 47.6%, and 60 open reading frames (ORFs). Surprisingly, the phage B1 genome has 97% identity with that of *Pediococcus damnosus* phage clP1 and 77% identity with that of *L. plantarum* phage JL-1; these phages were isolated from sewage and cucumber fermentation, respectively. The double-stranded DNA (dsDNA) genome of the *cos*-type phage B2 had 80,618 bp, a G+C content of 36.9%, and 127 ORFs with similarities to those of *Bacillus* and *Lactobacillus* strains as well as phages. Some phage B2 genes were similar to ORFs from *L. plantarum* phage LP65 of the *Myoviridae* family. Additionally, 6 tRNAs were found in the phage B2 genome. Protein analysis revealed 13 (phage B1) and 9 (phage B2) structural proteins. To our knowledge, this is the first report describing such high identity between phage genomes infecting different genera of lactic acid bacteria.

actobacilli are widely used in a variety of food fermentation processes, where they contribute to the flavor and texture of final products. They also produce organic acids, and the resulting low pH protects fermented products from degradation by spoilage microorganisms (15). In recent years, the industrial relevance of lactobacilli has been significantly enhanced by their increasing use as probiotics (12) or as a biotechnological tool (32).

Lactobacillus plantarum is commonly found as part of the natural microflora of fermented foods (dairy, vegetables, and meats) (12, 53, 68). This lactic acid bacterium may also be added as a starter or adjunct culture, in both cases improving the organoleptic characteristics of the final products (2, 12, 14, 15, 48, 49). Additionally, many L. plantarum strains possess documented probiotic properties, and marketed functional foods contain these strains (12, 53). L. plantarum can be used as a probiotic starter culture in the production of functional foods, taking advantage of, among others, its ability to grow in milk. However, the increasing use of L. plantarum as a starter or adjunct culture can lead to phage infections in industrial environments, with adverse effects on the final product (25, 51).

Phage infection is still one of the persistent causes of substandard dairy fermentation processes (60). Virulent phages can lyse starter cultures, yielding low-quality products that lead to economic losses. Consequently, efficient control measures to minimize problems caused by phage attacks become essential. In order to carry out successful antiphage strategies, knowledge about phage population and biology is needed (27, 39).

To date, over 30 *L. plantarum* phages, isolated from several sources, have been reported (16, 70, 72). All belong to the *Caudovirales* order (tailed phages, double-stranded DNA genome) (1, 70), and members belonging to each of the three *Caudovirales* families have been isolated: *Siphoviridae* (19 phages), *Myoviridae* (5 phages), and *Podoviridae* (1 phage). Other *L. plantarum* phages have been reported but not classified (70). Therefore, *L. plantarum* phages are relatively diverse and found in a wide variety of niches.

To our knowledge, only four *L. plantarum* phage genomes have been sequenced. Phage g1e (*Siphoviridae*, temperate) was isolated from plant materials and has a 42,259-bp genome with a G+C content of 43.1% and 62 open reading frames (ORFs) (37). Phage Sha1 (*Siphoviridae*, temperate) was isolated from kimchi and has a 41,726-bp genome with a G+C content of 40.6% and 58 putative ORFs (72). Phage JL-1 (*Siphoviridae*, virulent) was isolated from fermented cucumbers (43) and possesses a 36,700-bp genome with a G+C content of 39.4% and 52 ORFs. Finally, phage LP65 (*Myovidiae*, virulent) was isolated from fermented meat and has a very large genome of 131,573 bp with a G+C content of 37.3% and 165 ORFs (10).

Other *L. plantarum* phages have been analyzed in some detail; studies mainly included thermal and chemical sensitivities, and there were some preliminary genetics studies (9, 16, 44, 54, 65, 74). Overall, research on *Lactobacillus* phages has progressed over the past decade, but our knowledge of their biology and genetic composition is still limited and lags somewhat behind that of other industrially relevant phages (70).

The aim of this work was to carry out the characterization of two available *L. plantarum* phages. Phages ATCC 8014-B1 and ATCC 8014-B2 (herein referred to as B1 and B2, respectively) were previously isolated from corn silage and anaerobic sewage sludge (21).

## **MATERIALS AND METHODS**

Bacterial strains, phages, and culture conditions. L. plantarum strains were grown at 37°C in MRS broth (Difco). L. plantarum ATCC 8014 was

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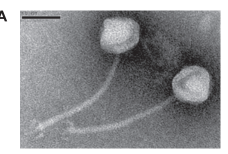
used as the host strain for phages B1 and B2. For phage amplification, MRS was supplemented with 10 mM CaCl<sub>2</sub> (MRS-Ca). Phage stocks were prepared as described previously (56) and stored as lysates at 4°C. Phage counts, expressed as PFU per milliliter, were obtained using the double-layer plaque titration method (64). Bacterial strains are maintained at the INLAIN Collection (Argentina) and the Félix d'Hérelle Reference Center for Bacterial Viruses of the Université Laval (Canada; www.phage.ulaval.ca) as frozen stocks in MRS broth containing 15% (vol/vol) glycerol. Phages B1 and B2 as well as the host *L. plantarum* ATCC 8014 were purchased from the American Type Culture Collection (Manassas, VA; www.atcc.org).

**Electron microscopy.** Ten microliters of 2% phosphotungstic acid (pH 7.0) was put in a clean sterile petri dish. A 200-mesh Formvar-carbon-coated copper grid (Pelco International) was deposited face down on the staining solution for 30 s. Then, 10  $\mu$ l of a purified phage suspension (10<sup>10</sup> PFU ml<sup>-1</sup>) was mixed with the stain by pipetting up and down. After 90 s, the grid was deposited face up on blotting paper. The grid was dried for 5 min and observed at 80 kV using a JEOL 1230 transmission electron microscope (62).

Microbiological assays. The host range of L. plantarum phages B1 and B2 was assessed by spotting 5 microliters of  $10^{-2}$  and  $10^{-4}$  dilutions of a high-titer lysate (10<sup>9</sup> PFU ml<sup>-1</sup>) on top of agar containing one of the eight L. plantarum strains tested (see Table 1). To study the phage adsorption process, L. plantarum cultures were grown in MRS to an optical density at 600 nm of 0.6 to 0.8, after which they were in contact with phage B1 or B2 at a final concentration of 10<sup>3</sup> PFU ml<sup>-1</sup>. The phage-containing cultures were incubated at 37°C for 15 min, and then we proceeded as described elsewhere (22). To determine the presence of active natural defense mechanisms against phages B1 and B2, the efficiency of plaquing (EOP) was calculated by dividing the phage titer on the test *L. plantarum* strain by the titer of the phage on the phage-sensitive host strain L. plantarum ATCC 8014. For phage-host systems showing reduced EOP values, two phage plaques obtained on the restrictive strain were purified and propagated on the same strain. The lysate obtained (modified phage) was titrated on both strains (original sensitive host and the restrictive strain) to determine a second EOP value. Modified phages were then propagated again on L. plantarum ATCC 8014, and the resulting lysate (unmodified phage) was titrated on both strains (4).

Phage DNA preparation and sequencing. Genomic DNA of phages B1 and B2 was isolated using a Maxi lambda DNA purification kit (Qiagen) with modifications (19). The restriction profiles of phage B1 and B2 DNA were compared to confirm differences. Restriction endonucleases (Roche Diagnostics) were used as recommended by the manufacturer. The DNA fragments were separated in a 0.8% agarose gel, stained with ethidium bromide, and photographed under UV illumination. Genome sequencing was performed at the Plateforme d'ADN génomique de l'Université Laval (Université Laval, Québec, Canada) using a GS-FLX Titanium apparatus (Roche) and the 454 pyrosequencing technique. For phage B1, 39,144 reads were generated and assembled into a single contig with a coverage of 430-fold. For phage B2, 4,670 reads were generated and assembled into a single contig with a coverage of 18-fold. The extremities of the genomes were determined by sequencing ligated phage DNA preparations using converging PCR primers at the genomic platform of the Centre Hospitalier de l'Université Laval with an ABI Prism 3100 apparatus.

Bioinformatics analysis. Sequence analyses were performed using BioEdit (30). Open reading frames (ORFs) were first identified using the GenMark program (46) and were further confirmed with ORFinder (http://www.ncbi.nlm.nih.gov/gorf/gorf.html). An ORF was considered valid if it had AUG, UUG, or GUG as the starting codon, encoded at least 29 amino acids (aa), and was preceded by an *L. plantarum* Shine-Dalgarno sequence (AGAAAGGAGGTGATC) (5). Function was attributed to an ORF using Blast2go (http://blast2go.bioinfo.cipf.es/start\_blast2go) and BLASTp (NCBI [http://blast.ncbi.nlm.nih.gov/Blast.cgi]). The annotations were supported by searching for protein functional domains using



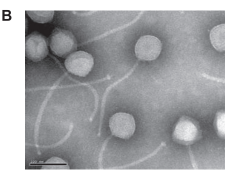


FIG 1 Electron micrographs of the phages B1 (A) and B2 (B). Bars, 50 nm (A) and 100 nm (B).

the NCBI Conserved Domain Database (http://www.ncbi.nlm.nih.gov/Structure/cdd/wrpsb.cgi) and EMBL InterProScan (http://www.ebi.ac.uk/Tools/InterProScan/). The tRNAs were identified using the tRNAscan-SE server (http://lowelab.ucsc.edu/tRNAscan-SE) and the ARAGORN program (41). Codon usage was determined through the DNA 2.0 Web server (Menlo Park, CA) and the Count-codon program available on the Kazusa DNA Research Institute Web page (http://www.kazusa.or.jp/codon/). The bacterial codon usage for the *L. planta-rum* host strains was obtained from the Kazusa DNA Research Institute database.

Analyses of phage B1 and B2 structural proteins. Phage lysates were concentrated with polyethylene glycol (PEG) and purified using two CsCl gradients (61). Purified phages were recovered by ultracentrifugation using a Beckman SW41 Ti rotor at 35,000 rpm (210,053  $\times$  g) for 3 h, followed by a second ultracentrifugation using a Beckman NVT65 rotor at 60,000 rpm (342,317  $\times$  g) for 18 h. The phage preparations were then dialyzed against phage buffer (0.05 M Tris-HCl [pH 7.5], 0.1 M NaCl, 8 mM MgSO<sub>4</sub>). Purified phages  $(4 \times 10^{11} \text{ PFU ml}^{-1})$  were treated as described elsewhere (62). Briefly, phages were mixed with  $4 \times$  loading buffer and boiled for 5 min. The samples were sonicated for 5 s with an ultrasonic Sonifier W-350 cell disrupter. Proteins were then separated by migration on a 12% SDS-polyacrylamide gel (1.5 mm thick). The Coomassiestained protein bands of interest were excised from the gel and identified by liquid chromatography-tandem mass spectrometry (LC-MS/MS) at the Centre Protéomique de l'Est du Québec (Université Laval, Quebec, Canada). These results were analyzed using the Scaffold Proteome software (13, 33, 55). Purified phage lysates were also directly analyzed by LC-MS/MS.

**Nucleotide sequence accession numbers.** The complete genome sequences of phages B1 and B2 have been deposited in GenBank under accession numbers JX486087 and JX486088, respectively.

#### **RESULTS AND DISCUSSION**

**Electron microscopy.** Both B1 and B2 phages have long noncontractile tails (Fig. 1) and belong to the *Siphoviridae* family, as do most characterized *L. plantarum* phages (70). Phage B1 has an icosahedral capsid with an estimated diameter of  $54 \pm 3$  nm and a

TABLE 1 Host range and adsorption rates of phages B1 and B2 on L. plantarum strains

		Phage B1		Phage B2	
Strain	Source <sup>a</sup>	EOP	Adsorption (%)	EOP	Adsorption (%)
ATCC 8014	ATCC	1.0	99.6 ± 4.8	1.0	90.8 ± 3.0
WCSF1	Human saliva (NCBIM collection)	$1.5 \times 10^{-2}$	$19.0 \pm 1.4$	b	0
LMG9211	Human saliva (BCCM collection)	$4.0 \times 10^{-3}$	$2.3 \pm 0.4$	_	$92.4 \pm 1.8$
PLN	NSLAB (INLAIN collection)	_	0	1.0	$98.5 \pm 1.5$
SMQ-1113	Industrial strain		$1.1 \pm 0.7$		$14.4 \pm 3.9$
SMQ-1114	Industrial strain		$8.2 \pm 2.3$		$3.2 \pm 1.0$
SMQ-1115	Industrial strain		$9.5 \pm 2.5$		$12.3 \pm 1.6$
SMQ-1116	Industrial strain		$13.7 \pm 3.6$		$10.4 \pm 6.4$

<sup>&</sup>lt;sup>a</sup> ATCC, American Type Culture Collection; BCCM, Belgian Coordinated Collections of Microorganisms; NCBIM, National Collection of Industrial and Marine Bacteria; NSLAB, nonstarter lactic acid bacteria; INLAIN, Instituto de Lactología Industrial.

tail of 157  $\pm$  10 nm in length and 8  $\pm$  1 nm in width. The baseplate appears somewhat complex, with spikes or fibers (Fig. 1A). Phage B2 has an icosahedral capsid with a diameter of 74  $\pm$  2 nm and a tail of 240  $\pm$  3 nm in length and 10  $\pm$  1 nm in width (Fig. 1B). Other investigators previously reported a larger capsid diameter (110 nm) and a much longer tail (500 nm) for *L. plantarum* phage B2 (54). Although dimensions may vary due to the use of different electron microscopes and methodologies (59), this cannot explain such large differences. At this time, it is unclear why such a discrepancy exists.

Microbiological assays. The results of the host range and adsorption tests are presented in Table 1. Each phage exhibited a distinctive host range but shared a common host (L. plantarum ATCC 8014). Phage B1 also replicated on L. plantarum strains WCSF1 and LMG9211, but the EOP was reduced. Surprisingly, under the conditions tested, the adsorption of phage B1 on strains LMG9211 and WCSF1 was very low, although clear plaques were formed. This low adsorption could be due to a limited number of phage receptors (in comparison with L. plantarum ATCC 8014) or their availability on the cell surface. Similar results were reported for Lactobacillus paracasei phages (8). Conversely, phage B2 was amplified on L. plantarum strain PLN and on its host ATCC 8014 (Table 1). Interestingly, phage B2 adsorbed well to strain LMG9211 without forming plaques (Table 1), suggesting the presence of phage resistance mechanisms in this strain (39). In general, phages were not able to adsorb on the other *L. plantarum* strains tested, suggesting the absence of receptors or perhaps adsorption blocking mechanisms (39).

Restriction/modification systems. As indicated above, *L. plantarum* LMG9211 and WCSF1 seemed to carry a natural defense system, as the EOP of phage B1 was reduced (Table 1). Phage plaques were recovered from these two hosts (LMG9211 and WCSF1), purified, and amplified on each strain. These amplified phages had an EOP of 1.0 on *L. plantarum* ATCC 8014. When these phages were propagated again in their original host, *L. plantarum* ATCC 8014, the EOP values were reduced and similar to those shown in Table 1. This temporary host-specific immunity suggests the presence of a classical restriction/modification (R/M) system in both strains (52). Besides, the same specificity might be involved in both systems, since an EOP value of 1 was obtained when LMG9211-amplified phage was tested on *L. plantarum* WCSF1 and when WCSF1-amplified phage was tested on *L. plantarum* LMG9211. A type I restriction/modification system was

previously identified in the genome of *L. plantarum* WCSF1, though its functionality was not demonstrated (36, 63).

Genome analysis. Phages B1 and B2 have linear doublestranded DNA genomes comprising 38,002 bp and 80,618 bp, respectively. Nes et al. (54) reported a relatively similar genome size for phage B2 (73 kbp), which was calculated from the addition of the molecular sizes of DNA restriction fragments. Phage B1 has the highest GC content (47.6%) reported to date for an L. plantarum phage. The GC content of phage B2 was much lower, at 37.0%, but is similar to the GC content of the L. plantarum myophage LP65 (10). The GC content of the host strain L. plantarum ATCC 8014 was previously estimated at 45.1% (50), whereas genome sequencing of strain WCSF1 revealed a GC content of 44.5% (36, 63). The genomes of two other L. plantarum strains also have GC contents of 44.5 to 44.7% (71, 75). The GC contents were similar throughout the genomic sequences of both phages, although some noncoding regions in phage B2 were AT rich. The lower GC content of phage B2 may suggest that some genetic elements were derived from phages infecting other hosts (23, 31).

The phage genomic DNA was also digested with various restriction enzymes (EcoRV, HindIII, MluI, and SalI), and the profiles obtained were similar to the theoretical profiles obtained from the genomic data (NEBcutter), suggesting the absence of modified nucleotides (data not shown). The profile obtained for phage B2 was similar to that reported elsewhere (54). Analysis of the genome extremities indicated that phage B1 is a *pac*-type phage, like *L. plantarum* phages fri, JL-1, and LP65 (10, 43, 65), whereas phage B2 was classified as a *cos*-type phage, similar to SC921 phage (74). The *cos* site is 11 nucleotides long (5'-TGAGC GCCCTA-3') (data not shown).

Sixty ORFs were identified for phage B1 and 127 ORFs for phage B2 (Tables 2 and 3; Fig. 2 and 3). They covered 93% (B1) and 87% (B2) of the genome length. A total of 56 ORFs (93%) for phage B1 and 65 ORFs (51%) for phage B2 had homology to previously characterized genes in public databases. However, a protein function could be attributed to products of only 25 ORFs (42%) for phage B1 and 37 ORFs (29%) for phage B2. The predominant starting codon was ATG for both phages (90% for B1, 86% for B2). Interestingly, four B1 ORFs share some identity with B2 ORFs, namely, B1 ORF15 and B2 ORF33, B1 ORF18 and B2 ORF36, B1 ORF22 and B2 ORF40, and B1 ORF35 and B2 ORF99. Of interest, ORF18 of phage B1 is likely involved in host recogni-

b -, not determined, as the phage does not infect the strain.

TABLE 2 Open reading frames deduced from the genome of L. plantarum phage B1 and their predicted functions

in	GenBank accession no.	YP_004934175	YP_004934174	YP_223885	YP_004934173	YP_223886	YP_004934172	YP_223887	YP_004934171	YP_223888	YP_004934170	YP_223889	YP_004934169	YP_223890	YP_004934168	YP_223891	YP_004934167	YP_223892	YP_004934166	YP_223893	YP_223894	YP_004934222	ZP_03964227
Aligned protein	Genl	YP_(	YP_(	YP_2	YP_(	YP_3	YP_(	YP_2	YP_(	YP_3	YP_(	YP_3	YP_(	YP_2	YP_(	YP	YP_(	YP_3	YP_(	YP_2	YP_2	YP_(	ZP (
Aligno	Size (aa)	139	431	440	513	206	270	273	176	184	286	286	75	64	115	113	92	94	125	125	117	208	199
	E value	3.0E-64	0	0	0	0	0	2.0E-72	1.0E - 118	1.0E - 27	0	7.0E - 138	9.0E-37	6.0E-11	6.0E-74	2.0E-42	5.0E-57	3.0E-27	5.0E-85	3.0E-46	2.0E - 40	5.0E - 150	4 OF-92
	Best match(es) (extent <sup>c</sup> ; % amino acid identity)	clP1_010 (P. damnosus phage clP1)	(39/103; 90) clP1_009 ( <i>P. damnosus</i> phage clP1) (430/431; 99)	ORF440 ( <i>L. plantarum</i> phage phiJL-1) (293/431; 68)	clP1_008 ( <i>P. damnosus</i> phage clP1) (508/513: 99)	ORF506 ( <i>L. plantarum</i> phage philL-1) (284/506: 56)	clP1_007 ( <i>P. damnosus</i> phage clP1) (266/270; 99)	ORF273 ( <i>L. plantarum</i> phage	clP1_006 ( <i>P. damnosus</i> phage clP1)	ORF184 ( <i>L. plantarum</i> phage	clP1_005 (P. damnosus phage clP1_084/286: 99)	ORF286 (L. plantarum phage	phJL-1) (198/286; 69) clP1_004 ( <i>P. damnosus</i> phage clP1	(74/75; 99) ORF64b ( <i>L. plantarum</i> phage	phiJL-1) (33/49; 67) clP1_003 (P. damnosus phage clP1)	(113/115; 98) ORF113 ( <i>L. plantarum</i> phage	clP1_002 (P. damnosus phage clP1) (90/92: 98)	ORF94 ( <i>L. plantarum</i> phage phiJL- 1) (49/93; 53)	dP1_001 ( <i>P. damnosus</i> phage clP1) (123/125; 98)	ORF125 ( <i>L. plantarum</i> phage philL-1) (77/121; 64)	ORF117 ( <i>L. plantarum</i> phage	clP1_057 (P. damnosus phage clP1) (208/208: 100)	ODE100 (I plantamim phase
	Predicted function	Terminase small subunit	Terminase large subunit		Portal protein		Capsid protein		Scaffold protein		Major capsid protein		Minor capsid protein		DNA packaging				Head to tail joining			Major tail protein	
	Putative RBS and start $\operatorname{codon}^b$	$\overline{A}\overline{A}\overline{A}\overline{A}\overline{A}\overline{G}\overline{G}\overline{A}\overline{A}\overline{A}\overline{G}\overline{G}\overline{g}a$	$\overline{A}AC\overline{A}TC\overline{G}G\overline{G}T\overline{1}TCC\overline{C}ga$ A $TG$		$\overline{A}T\overline{AA}C\overline{G}G\overline{A}G\overline{G}AGT\overline{T}A$ aaac $ATG$		C <u>GAAAGG</u> C <u>GG</u> ATTGAttatca <b>ATG</b>		<u>A</u> TTGAGGAGGAGAAccatc <b>ATG</b>		C <u>GGAGGAA</u> CT <u>T</u> A <u>AAC</u> a <b>ATG</b>		CA <u>AAAG</u> ACC <u>G</u> CT <u>A</u> G <u>C<b>ATG</b></u>		$\overline{ ext{A} ext{T} ext{A} ext{A} ext{T} ext{T} ext{A} ext{C} ext{G} ext{T} ext{A} ext{C} ext{C} ext{gtatgg} ext{G} ext{T} ext{G}$		G <u>G</u> GGTTC <u>AGGT</u> TC <u>T</u> T <b>ATG</b>		GA <u>A</u> CGT <u>G</u> CC <u>GT</u> T <u>ATC</u> a <b>ATG</b>		C <u>G</u> CT <u>AGGGGTG</u> TCAcaag <b>ATG</b>	$\overline{A}  ext{ATG} \overline{A}  ext{GGG}  ext{GGTGT} \overline{A}$ aaaaaat $ ext{ATG}$	
	Iq	6.3	8.2		4.8		8.8		4.7		4.8		4.4		5.0		9.5		9.3		8.0	4.3	
Predicted protein	Molecular mass (kDa)	11.4	48.9		58.4		30.6		19.1		30.9		7.2		14.1		10.2		15.1		13.2	22.7	
Predict	Size (aa)	103	431		513		270		176		286		75		123		92		137		117	208	
	Stop (bp)	553	1845		3402		4132		4762		5640		5915		6314		6859		66269		7388	8033	
	Start (bp)		550		1861		3320		4232		4780		5688		5943		6311		6546		7035	7407	
	ORF <sup>a</sup>		2		8		4		r.		9		7		∞		6		10		11	12	

(Continued on following page)

TABLE	TABLE 2 (Continued)	tinued)									
			Predic	Predicted protein						Aligne	Aligned protein
$\mathrm{ORF}^a$	Start (bp)	Stop (bp)	Size (aa)	Molecular mass (kDa)	Iq	Putative RBS and start $\operatorname{codon}^b$	Predicted function	Best match(es) (extent'; % amino acid identity)	E value	Size (aa)	GenBank accession no.
27	22616	24556	646	72.7	5.1	<u>AAA</u> GG <u>GGA</u> AA <u>T</u> AAAGcact <b>ATG</b>	DNA primase	clP1_042 ( <i>P. damnosus</i> phage clP1) (641/646, 99)	0	646	YP_004934207
								ORF637 ( <i>L. plantarum</i> phage phiJL-1) (285/634; 45)	7.0E-180	637	YP_223911
29	24921	25337	138	15.7	5.1	<u>A</u> TTG <u>AGGAGG</u> AA <u>AT</u> Gta <b>ATG</b>		clP1_040 ( <i>P. damnosus</i> phage clP1)	2.0E-68	138	YP_004934205
30	25340	25639	66	11.3	7.8	$\overline{A}A\overline{A}CG\overline{G}GAGG$ ATTATtaaat $f ATG$		clP1_039 ( <i>P. damnosus</i> phage clP1) (64/99, 65)	4.0E-39	66	YP_004934204
31	25600	26091	163	18.6	10.3	<u>A</u> CT <u>AAGGGGGTGA</u> AAac <b>ATG</b>	Replication protein	clP1_038 ( <i>P. damnosus</i> phage clP1)	4.0E - 102	150	YP_004934203
32	25959	26450	163	19.2	5.2	CGCGGGCACGTCATCtATT		clP1_037 ( <i>P. damnosus</i> phage clP1) (115/123, 93)	8.0E-80	123	YP_004934202
33	26488	26685	92	9.5	6.2	${ m TC} \overline{ m A} { m T} \overline{ m A} \overline{ m G} \overline{ m G} \overline{ m G} \overline{ m G} \overline{ m T} { m Tat} { m ATG}$		clP1_036 ( <i>P. damnosus</i> phage clP1) (62/65, 95)	2.0E-37	99	YP_004934201
34	26687	27151	154	17.8	5.1	$\underline{A}A\underline{A}AAGGGAATTAT$ taac $\pmb{A}\pmb{T}\pmb{G}$	Replicase	clP1_035 ( <i>P. damnosus</i> phage clP1) (153/154, 99)	4.0E - 106	154	YP_004934200
								ORF153 ( <i>L. plantarum</i> phage phiJL-1) (74/156; 47)	3.0E-35	153	YP_223913
35	27167	27487	106	11.8	4.6	<u>AGTAAAGGGGTAAA</u> AAcg <b>ATG</b>	DNA binding	clP1_034 ( <i>P. damnosus</i> phage clP1) (100/106, 94)	2.0E-67	106	YP_004934199
								ORF97 ( <i>L. plantarum</i> phage phiJL-1) (45/94; 48)	1.0E - 18	97	YP_223877
37	27725	29155	476	53.3	9.1	$\overline{1G}\overline{1A}\overline{CGGAGG}GAT\overline{1}Gca$ ATG	Helicase	clP1_033 ( <i>P. damnosus</i> phage clP1) (470/476, 99)	0	476	YP_004934198
								ORF467 ( <i>L. plantarum</i> phage phiJL-1) (291/446; 65)	0	467	YP_223915
38	29199	29471	06	10.2	4.8	$\overline{A}$ TC $\overline{A}\overline{A}\overline{G}$ C $\overline{A}\overline{A}\overline{G}$ G $\overline{G}\overline{A}$ GGta $f A$ T $f T$		clP1_032 ( <i>P. damnosus</i> phage clP1) (64/88, 73)	2.0E-39	102	YP_004934197
39	29458	29832	124	13.8	4.4	<u>AGAAAAGGGGT</u> AT <u>T</u> Ttg <b>ATG</b>		clP1_031 ( <i>P. damnosus</i> phage clP1) (119/124, 96)	3.0E-71	125	YP_004934196
40	29834	30253	139	16.0	5.4	T <u>GAAAGGA</u> TTGAT <u>T</u> Aac <b>ATG</b>		clP1_030 ( <i>P. damnosus</i> phage clP1) (125/143, 87)	3.0E-85	149	YP_004934195
41	30225	30659	144	16.4	5.3	<u>AAACTAAAAGTCA</u> CGa <b>ATG</b>		clP1_029 ( <i>P. damnosus</i> phage clP1) (98/139, 71)	4.0E-68	139	YP_004934194
42	30659	31102	147	16.2	0.6	$\overline{A}A\overline{A}AAGGGGTAAT\overline{T}GaataATG$		clP1_028 ( <i>P. damnosus</i> phage clP1) (129/147, 88)	5.0E-89	147	YP_004934193
43	31095	32072	325	36.9	5.3	$\overline{A}$ TT $\overline{A}$ T $\overline{G}GAGGT$ TG $\overline{L}$ Gaaag $A$ T $G$	Structural protein	clP1_027 ( <i>P. damnosus</i> phage clP1) (306/325, 94)	0	324	YP_004934192
44	32073	32468	131	14.3	4.5	$\overline{A}\overline{A}\overline{A}\overline{\Gamma}C\overline{G}G\overline{A}G\overline{\Gamma}\overline{\Gamma}\overline{A}\overline{\Gamma}$ Ttaa $m{A}m{T}m{G}$		clP1_026 (P. damnosus phage clP1) (129/131, 98)	1.0E-89	131	YP_004934191
45	32601	32840	79	9.1	0.9	C <u>GAAAGGA</u> C <u>G</u> AGGGAtaa <b>ATG</b>		clP1_025 ( <i>P. damnosus</i> phage clP1) (78/79, 99)	4.0E-50	79	YP_004934190

YP_004934189	YP_223917	YP_004934188	YP_004934187	YP_004934186	YP_004934185	YP_223874	YP_004934184	YP_004934183	YP_004934182	YP_004934181	YP_004934180	YP_004934179	YP_004934178	YP_004934177	YP_004934176
118	114	92	42	57	81	77	119	66	63	53	144	148	131	172	141
2.0E-81	8.0E-31	3.0E-35	3.0E-6	6.0E - 18	6.0E-51	2.0E-3	1.0E - 64	1.0E-65	4.0E-37	1.0E - 27	4.0E - 95	3.0E - 94	3.0E-87	1.0E - 119	4.0E - 93
clP1_024 ( <i>P. damnosus</i> phage clP1) (117/118, 99)	ORF114 ( <i>L. plantarum</i> phage phiJL-1) (54/104, 52)	clP1_023 ( <i>P. damnosus</i> phage clP1) (61/62, 98)	clP1_022 ( <i>P. damnosus</i> phage clP1) (39/42, 93)	clP1_021 ( <i>P. damnosus</i> phage clP1) (42/44, 95)	clP1_020 ( <i>P. damnosus</i> phage clP1) (78/81, 96)	ORF77 ( <i>L. plantarum</i> phage phi)L-1) (40/75, 53)	clP1_019 ( <i>P. damnosus</i> phage clP1) (94/97, 97)	clP1_018 ( <i>P. damnosus</i> phage clP1) (99/99, 100)	clP1_017 ( <i>P. damnosus</i> phage clP1) (61/63, 97)	clP1_016 ( <i>P. damnosus</i> phage clP1) (52/53, 98)	clP1_015 ( <i>P. damnosus</i> phage clP1) (136/144, 94)	clP1_014 ( <i>P. damnosus</i> phage clP1) (132/148, 89)	clP1_013 ( <i>P. damnosus</i> phage clP1) (1126/131, 96)	clP1_012 ( <i>P. damnosus</i> phage clP1) (167/172, 97)	clP1_011 (P. damnosus phage clP1)
														Endonuclease	
$\overline{A}$ TC $\overline{A}$ T $\overline{G}G\overline{A}GG$ ACGA $\overline{C}$ a <b>ATG</b>		<u>AGGAAAG</u> T <u>GGT</u> AA <u>T</u> Aaaa <b>ATG</b>	GGATATGAGGTGATCgaATG	<u>AACAAAGGGGT</u> CT <u>T</u> Atatt <b>ATG</b>	<u>AAAAAGGGGCCA</u> AGta <b>ATG</b>		<u>AGAA</u> C <u>G</u> TCATG <u>GGTC</u> g <b>ATG</b>	TA <u>AAAGGCGGCGA</u> GAtt <b>ATG</b>	$\overline{ ext{A}}\overline{ ext{A}}\overline{ ext{A}}\overline{ ext{G}}\overline{ ext{G}}\overline{ ext{G}}\overline{ ext{A}}\overline{ ext{A}} e$	CGAAAAGGGGTTTTTaa $ATG$	<u>A</u> ATT <u>AGGAGG</u> GTT <u>T</u> Tacc <b>ATG</b>	<u>A</u> CGG <u>AGG</u> TT <u>G</u> AA <u>ATC</u> a <b>ATG</b>	<u>A</u> TCG <u>AGGTG</u> AA <u>G</u> C <u>T</u> Ac <b>ATG</b>	TACTG <u>GGAGGTG</u> T <u>T</u> Atgac <b>ATG</b>	TGAAAGGTGATAATA <b>ATG</b>
9.3		5.2	8.5	12.1	5.6		9.5	8.6	4.4	0.9	5.0	6.1	8.6	10.0	9.1
13.7		7.6	4.7	6.2	9.3		11.5	11.1	7.2	6.1	16.8	17.1	14.9	19.1	15.7
118		62	42	57	81		26	66	63	53	144	148	131	172	138
33183		33706	33831	34030	34275		34634	34920	35108	35238	35685	36124	36512	37032	37441
32827 3		33518 3	33703 3	33857 3	34030 3		34341 3	34621 3	34917 3	35077 3	35251 3	35678 3	36117 3	36514 3	37025 3
46		48	49	50	51		52	53	54	55	56	57	28	59	09

<sup>&</sup>lt;sup>a</sup> Only the ORFs with significant hits to those of other proteins in the database are included.

<sup>b</sup> RBS, ribosome binding site. Underlined codons correspond to bases identical to the *L. plantarum* RBS consensus sequence; uppercase letters represent the RBS sequence; boldface indicates the starting codon; lowercase letters indicate spacer nucleotides between the RBS and start codon.

<sup>c</sup> Number of identical amino acids/total number of amino acids.

<sup>d</sup> NTP, nucleoside triphosphate.

 TABLE 3 Open reading frames deduced from the genome of L. plantarum phage B2 and their predicted proteins

			Predic	Dredicted protein			X X			Aliane	Alianed protein
i i	ć	ć	I Iculi	ced protein						- 10 mgm	Protein
ORF" or tRNA	Start (bp)	Stop (bp)	Size (aa)	Molecular mass (kDa)	pl	Putative RBS and start codon <sup>b</sup>	Predicted function	Best matches (extent'; % amino acid identity)	E value	Size (aa)	GenBank accession no.
12	3401	2769	210	25.1	5.2	TAAAAGGAGGAATAAaatt <b>ATG</b>	Fatty acid/phospholipid	PlsX gene product (Clostridium	9.0E-2	337	YP_003779451
15	4262	4393	43	4.8	3.9	<u>AAAA</u> G <u>GGAGG</u> ACT <u>T</u> Agc <b>ATG</b>	synthesis protein C.D.": transcription factor	fjungdahlii DSM 13528) (29/154; 19) EUBREC_3432 (Eubacterium rectale ATCC 33556) (15/43: 35)	2.0E-3	209	YP_002939292
16	4320	4802	160	18.6	9.3	<u>ACAA</u> GAA <u>AGATGAT</u> T <b>ATG</b>	C.D.: HNH endonuclease	pls32_p096 (Bacillus subtilis subsp.	8.0E-26	135	YP_004243694
17	4815	5321	168	18.7	5.1	TCT <u>AAGG</u> G <u>GGTGA</u> A <u>C</u> ac <b>ATG</b>	Terminase small subunit	pls32_p095 (Bacillus subtilis subsp.	3.0E - 22	163	YP_004243693
18 tRNA tRNA	5476 6246 6925	5988 6317 6997	170	19.7	4.9	<u>AAAA</u> G <u>G</u> T <u>AGGT</u> A <u>A</u> AGteaa <b>ATG</b>	tRNA-Asn tRNA-Leu	naro) (01/106; 37) gp089 ( <i>Lc. lactis</i> phage 949) (33/114; 29)	8.0E-11	171	YP_004306249
20	7143	7403	98	10.2	5.3	GA <u>AA</u> GA <u>GAGG</u> AA <u>A</u> AC <b>tATG</b>	INIVA-Wiel	ORF78 ( <i>L. plantarum</i> phage LP65) (81/ 86; 95)	1.0E - 40	109	YP_164713
tRNA 21	7742 7937	7814 9646	569	65.0	5.5	TAT <u>AA</u> C <u>GAGGTGAT</u> Atg <b>TTG</b>	tRNA-Gly Terminase large subunit	ORF5 (L. delbrueckii phage c5) (211/	9.0E-131	559	ACA63297
22	9830	11137	435	48.0	4.8	<u>ACAGAAGA</u> TAC <u>G</u> GTT <b>ATG</b>	Portal protein	522; 40) ORF5 ( <i>L. delbrueckii</i> phage LL-Ku) (106/384: 28)	3.0E-37	404	AAV30165
23	11133	12236	367	40.6	4.3	$T\underline{G}TT\underline{A}GGAGGTA\underline{A}TGacam{A}TT$	Major capsid	pls32_p090 (Bacillus subtilis subsp.	1.0E - 38	313	YP_004243688
24	12240	13526	428	45.9	5.3	GATTT <u>GGAGGT</u> CTAAtta <b>ATG</b>	protein/protease Major capsid protein	natto) (93/201; 46) ORF7 (L. delbrueckii phage LL-Ku)	1.0E-21	395	AAV30167
25	13657	14115	152	15.9	4.5	T <u>GA</u> T <u>AGGAGG</u> GA <u>AT</u> Acta <b>TTG</b>	Tail protein	(100/201; 28) SPC35_0138 (enterobacterial phage SPC35) (37/90-41)	5.0E - 9	162	YP_004306621
27	14453	14881	142	15.9	5.4	${ m TTT}_{\overline{ m A}}{ m GT}_{\overline{ m GAGGTGA}}{ m GA}{ m aa}{ m ATG}$	Head-tail adaptor	HMPREF9104_01875 (Lactobacillus kiconencis F0435) (23/87: 26)	1.0E-3	120	ZP_09556164
28	14832	15323	163	18.4	6.2	<u>AAA</u> CG <u>G</u> CG <u>GTTCATC</u> tg <b>GTG</b>	Head-tail joining protein	LSL_0288 (Lactobacillus salivarius phage Sal2) (31/126: 25)	7.6E-2	130	YP_535185
29 30	15447 15723	15704 16334	85 203	9.7 21.5	4.8	GTTC <u>AG</u> A <u>A</u> TAAA <u>AT</u> A <b>GTG</b> <u>A</u> AG <u>AAAGGTAAT</u> Tacta <b>ATG</b>	Tail protein Major tail protein	ORFIO ( <i>L. asei</i> phage phiAT3) (23/83; 28) OF33 ( <i>L. plantarum</i> phage Sha1) (61/	5.0E-2 8.0E-06	123	YP_025035 ADW01304
31	16442	16861	139	15.6	4.4	<u>AAAAA</u> TA <u>A</u> AA <u>T</u> TAT <u>T</u> Tag <b>ATG</b>		203; 29) LAR_1055 (Lactobacillus reuteri JCM 1112) (31/95; 33)	1.0E - 5	132	YP_001842051
33	17115	22778	1,887	199.5	10.1	$T\overline{G}CAAGGAGGGTT\overline{T}T$ aa $ATG$	Tail tape measure protein	LSL_0794 ( <i>L. salivarius</i> phage SalI) (2131/686: 35)	4.0E - 93	1,274	YP_535687
34	22802	24613	603	66.1	0.9	$\underline{A} \overline{1} \underline{A} C G A \underline{G} G \underline{G} \overline{G} \overline{1} \underline{A} \underline{A} \overline{1} \underline{C} c c c t c \boldsymbol{G} \boldsymbol{T} \boldsymbol{G}$	Minor structural protein	(245)(200, 52) ORF27 (L. plantarum phage Sha1) (321/ 596: 54)	1.0E-177	290	ADW01308
35	24669	27053	794	88.0	4.7	GATT <u>AGGAGGTAAT</u> Gga <b>ATG</b>	Minor structural protein	ORF28 (L. plantarum phage Sha1) (339/ 768: 44)	0	789	ADW01309
36	27065	30406	1,113	118.8	4.9	GAGT <u>AGGAGGT</u> T <u>ATC</u> aa <b>ATG</b>	Tail fiber/Host specificity	ORF29 (L. plantarum phage Sha1) (373/	0	929	ADW01310
37	30403	30624	73	8.0	4.7	CTGG <u>AGGA</u> TAAA <u>ATC</u> aa <b>ATG</b>	protein	2065-05) ORF30 ( <i>L. plantarum</i> phage Sha1) (61/ 71; 86)	3.0E-34	84	ADW01311

16.1         4.7         AAATAGGAGGAAATTaaactATG           41.8         4.6         AGAAAGGAATGATTFggtTTG           50.7         9.4         TAAAAGGAAGAAAagATG	A <u>AT</u> Taa AT <u>T</u> Tgg
5.3 $\underline{ATAAGG}$ 6.4.4 $\underline{AGGGAA}$	TCA <u>C</u> cacA
38.3 7.7 G <u>GGAATGGTGAGAT</u> Aca <b>ATG</b> 95.1 7.9 <u>ATTAGTCAGATGA</u> AGat <b>ATA</b>	<u>GAT</u> Aca <b>A</b> T 3AAGat <b>A</b> TA
54.9 5.7 <u>ATAGAGGAGG</u> AA <u>A</u> AAta <b>ATG</b>	\A <u>A</u> AAta <b>A</b> TG
7.8	C <u>AT</u> Att <b>ATG</b>
21.4 9.2 $\overline{\text{ATTAAGAACATTACCATG}}$	T <u>A</u> C <u>C</u> ATG
11.7 5.0 TGATTGGAGCAGTGAta <b>ATG</b>	<u>G</u> TGAata <b>A</b> T
24.3 5.2 <u>AGAAAG</u> AG <u>GTTTAT</u> Tttaa <b>ATG</b>	T <u>AT</u> Tttaa <b>AT</b> C
20.7 8.8 $AGGGAGAATTAAATcATG$	`A <u>A</u> ATc <b>ATG</b>
9.7 5.8 AGCGAGGAAAACGG <u>Ceg<b>GTG</b></u> 15.9 7.8 GA <u>AGGAGGTAA</u> ATaatg <b>ATG</b>	\CGG <u>C</u> cg <b>GTG</b> `A <u>A</u> ATaatg <b>AT</b> G
17.1 4.4 <u>AAGACGGAGGTAAAAATG</u>	AAAAtaATG
12.0 7.7 <u>AGATGGAGAGTGCT</u> Aaa <b>gATG</b> 13.2 4.1 G <u>GAAAAGTGATTGTAA<b>TG</b></u>	<u>'G</u> C <u>T</u> Aaag <b>ATG</b> TG <u>T</u> A <b>ATG</b>
26.5 9.4 CAGAGTCAGTTAGTCgg <b>GTG</b>	'AG <u>TCgg<b>GTG</b></u>
46.2 5.7 TACT <u>AG</u> AG <u>GGAGAAC</u> tta <b>ATG</b>	<u>GA</u> A <u>C</u> tta <b>ATG</b>
39.2 5.9 <u>AGGACTGAGACTAT</u> Aaaa <b>ATG</b> 36.0 4.8 <u>AACTAGGAGGAATTTgaa<b>TG</b></u>	TATATAaaaATG
13.1 4.4 CACG <u>A</u> TA <u>A</u> T <u>GTGA</u> AT <b>tATG</b>	<u>GA</u> ATt <b>ATG</b>
20.4 9.2 TA <u>AAA</u> TT <u>A</u> AAAATA <u>C</u> aa <b>ATG</b>	ΛΤΑ <u>C</u> aa <b>ΑΤG</b>
20.5 6.3 GAAGTGGAGTTGAGCgaatATG	JAGGast AT

GenBank gned protein

YP\_004243623

YP\_004306222

YP\_164649

YP\_164648

YP\_223877

YP\_164640 YP\_223879

ADM73619

YP\_004243622

YP\_002790752 accession no.

		(2001)								
			Predi	Predicted protein						Align
$ORF^a$ or	Start	Stop	Size	Molecular				Best matches (extent <sup>c</sup> ; % amino acid		Size
tRNA	(bp)	(bp)	(aa)	mass (kDa)	pl	Putative RBS and start $\operatorname{codon}^b$	Predicted function	identity)	E value	(aa)
83	58517	59224	235	27.2	4.8	<u>A</u> ATT <u>AGGAGG</u> AA <u>A</u> AAta <b>TTG</b>	Deoxyguanosine kinase	ORF73 ( <i>L. paracasei</i> phage Lb338-1) (80/239; 33)	2.0E-42	240
85	59491	59961	156	18.2	9.1	$\underline{A}A\underline{A}C\underline{A}GGAGG\underline{T}T\underline{A}AA$ acca $f ATG$		yorH (Bacillus subtilis subsp. natto) (49/ 162; 31)	3.0E-12	162
98	59954	61543	529	60.3	4.9	TT <u>A</u> GT <u>GGAGATGAT</u> Ttac <b>TTG</b>	DNA helicase	yorI (Bacillus subtilis subsp. natto) (186/ 517; 36)	2.0E-89	530
87	61745	62590	281	32.1	9.8	$ ext{CTC} \overline{ ext{AA}}  ext{ACT} \overline{ ext{G}}  ext{TG} \overline{ ext{G}}  ext{T} \overline{ ext{C}}  ext{a}  ext{T}  ext{G}$	DNA primase	ORF61 (Lc. lactis phage 949) (49/162; 31)	2.0E - 6	330
88	62587	64332	581	65.6	5.3	<u>ACAAGGAAGGTAAT</u> Gtc <b>TTG</b>	Single-stranded DNA exonuclease	ORF62 ( <i>Lc. lactis</i> phage 949) (141/572; 25)	2.0E - 18	593
91	64909	65292	127	14.4	9.9	CATT <u>AGGAGG</u> AA <u>A</u> AAgcg <b>ATG</b>		ORF14 ( <i>L. plantarum</i> phage LP65) (53/ 118; 45)	2.0E-22	120
86	67529	67873	114	13.5	4.8	TA <u>AAA</u> T <u>GA</u> C <u>G</u> AA <u>A</u> GAacta <b>ATG</b>		ORF13 ( <i>L. plantarum</i> phage LP65) (33/106; 31)	7.0E-9	124
66	67870	68175	101	11.7	5.3	T <u>GGAA</u> T <u>G</u> GA <u>GAGACATA</u>	DNA binding protein	ORF97 ( <i>L. plantarum</i> phage phiJL-1) (66/97; 68)	3.0E-41	26
100	68172	68573	133	15.6	4.4	$\overline{A}A\overline{A}G\overline{A}G\overline{A}G\overline{G}A\overline{1}\overline{A}A$ Gaagct $A\mathbf{T}G$		ORF5 ( <i>L. plantarum</i> phage LP65) (26/ 71: 37)	1.0E - 1	182
101	96889	69588	230	26.4	6.7	GGAGAGGAGTTTaaat $f ATG$		ORF93 ( <i>L. plantarum</i> phage phiJL-1) (35/85; 41)	1.0E - 08	93
102	69578	70021	147	16.9	5.9	<u>AGAAAGGTG</u> ACA <u>A</u> CG <b>ATG</b>		ORF157 ( <i>L. plantarum</i> phage LP65) (55/156; 35)	4.0E-10	146
								ORF142 ( <i>L. plantarum</i> phage phi)L-1) (52/149; 35)	2.0E-10	142
105	70682	71113	143	16.3	4.4	<u>AAAA</u> CGGAGGTGGCAacg <b>ATG</b>	DNA replication protein	ORF15 ( <i>L. plantarum</i> phage LP65) (55/ 161; 34)	2.0E-07	149
								ORF115 ( <i>L. plantarum</i> phage phig1e) (49/144; 34)	2.0E-10	115
108	71911	72354	147	16.6	4.9	TCG <u>A</u> T <u>GG</u> TA <u>GTGA</u> CGat <b>ATG</b>		ORF8 ( <i>L. plantarum</i> phage Sha1) (49/ 106; 46)	3.0E-18	140
110	72510	72953	147	17.4	5.7	<u>AGGAAGG</u> CA <u>GTG</u> G <u>T</u> Aatc <b>ATG</b>		ORF93 ( <i>L. plantarum</i> phage phiJL-1) (54/91: 59)	4.0E-21	93
1111	72964	73206	80	9.5	4.7	TGTT <u>AGG</u> GGAATAAt <b>ATG</b>		phiglep44 (L. plantarum phage phigle)	2.0E - 3	73

YP\_004301357

2.0E - 2

 $\begin{array}{l} \operatorname{gp24}\left(Brochothrix\;\mathrm{phage\;A9}\right)\,(37/125;\\ 30) \end{array}$ 

NP\_695176 ADW01289

YP\_164650

YP\_223880

YP\_164792

NP\_695175

YP\_223879

NP\_861801

193

1.0E - 12

tk (enterobacterial phage RB69) (56/

Thymidine kinase

<u>AGTAAGAAGGGAAAAa**ATG**</u>

5.0

25.0

217

74617

73964

114

196; 29)

YP\_239845

93

9.0E - 125.0E - 58

ORF40 (Staphylococcus phage 2638A)

ORF165 (L. plantarum phage LP65)

<u>AGAA</u>TA<u>GAGG</u>CTTATtaaa**ATG** <u>AAATAAGGGTTG</u>CAAttaa**GTG** 

10.6

14.5 15.8

127 138

78140 77731

4.3

79630

79214

TA<u>AAAGGGGTG</u>T<u>T</u>Gag**ATG** 

9.3

7.0

09

77549 77757

123 124 127

YP\_164800

135 198

TABLE 3 (Continued)

 $<sup>^{</sup>a}$  Only the ORFs with significant hits to those of other proteins in the database are included.  $^{b}$  Abbrewiation and presentation are as for Table 2.

<sup>&</sup>lt;sup>c</sup> Extent is as defined for Table 2. <sup>d</sup> C.D., conserved domains.

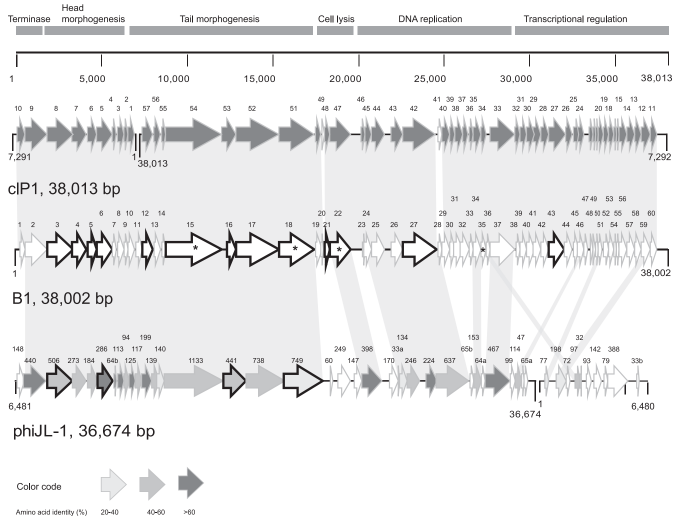


FIG 2 Genomic organizations of *L. plantarum* phages B1 and phiJL-1 as well as *P. damnosus* phage clP1. The scales above the genomes are in base pairs. Each arrow represents an ORF, and the numbering refers to Table 2 (for B1) and to the locus tags from phiJL-1 (accession number AY236756) and clP1 (accession number JN051154). The modules were based on the B1 organization. Genes coding for structural proteins experimentally determined by SDS-PAGE are indicated by thick outlines. Products of ORFs from phiJL-1 and clP1 sharing amino acid identity with those from B1 were drawn in a shade of gray according to the color code, and were linked by a shadow. White arrows represent products of ORFs sharing no identity. Phage phiJL-1 and clP1 genomes were split and reorganized in order to facilitate the alignment. ORFs sharing identity (>20%) with those of phage B2 are indicated by asterisks.

tion, and its identity with B2 ORF36 agrees with the observation that both phages infect the same host strain.

Presence of tRNA in the B2 genome. Six tRNAs were found in genome of phage B2 (Table 3) but none in B1. These six tRNAs deliver the amino acids asparagine (Asn, AAC), leucine (Leu, CTA), methionine (Met, ATG), glycine (Gly, GGA), and arginine (Arg, AGG and AGA). They were located in two genomic regions (6246 to 7814 and 42308 to 42522) of phage B2. Among *L. plantarum* phages for which the genomes are available, only the myophage LP65 contained tRNAs (14 tRNAs). The presence of tRNAs is often linked to large phage genomes (62).

The frequency of codon usage was then investigated for phages B1 and B2 (Table 4). The anticodons of some tRNAs found in the genome of phage B2 did not correspond to the codons most frequently used by the phage. For example, one tRNA matched the CTA codon, encoded a leucine, and had a frequency of 21.1% in the whole genome, whereas the most frequently used leucine

codon was TTA, which had a frequency of 42.5%. However, the CCT and TCT codons, which encoded arginine, were used more by phage B2 than other possible codons.

The codon usage of phage B2 was also compared to that of *L. plantarum* WCFS1 because no bacterial host strain for phage B2 has been sequenced yet (Table 4). Our results agreed with others (3) who suggested that phages encode tRNAs corresponding to codons that are less used by the host bacteria to increase specific phage protein expression (Table 4). The presence of tRNAs was reported for some *Lactococcus* phages: P087 (5 tRNAs) (69), KSY1 (3 tRNAs) (11), and 949 (6 tRNAs) (62). In contrast to the results observed here, the frequencies of codon usage by phage 949 tRNAs were similar for the phage and its host *Lactococcus lactis* IL1403.

Function assignment and genomic organization of phages B1 and B2. The ORF functions were assigned based on comparison with sequences in public databases (NCBI, InterProScan). Only the ORFs with the highest identity with those encoding other

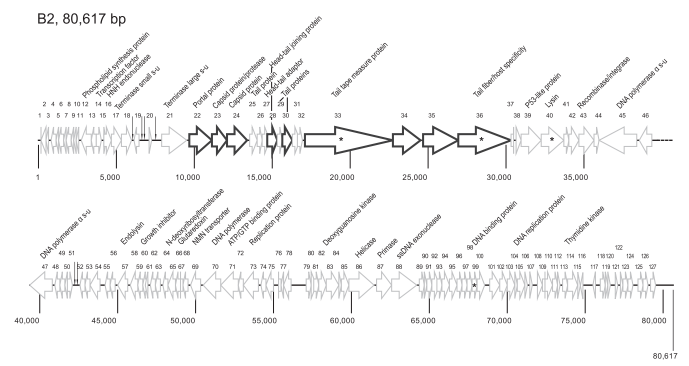


FIG 3 Genomic organization of *L. plantarum* phage B2. The scale under the genome is in base pairs. Each arrow represents an ORF, with its putative function, and the numbering refers to Table 3. Genes coding for structural proteins experimentally determined by SDS-PAGE are indicated by thick outlines. tRNAs are indicated by vertical arrows. ORFs sharing identity (>20%) with those of phage B2 are indicated by asterisks.

proteins in the database are shown in Tables 2 and 3. Although phages B1 and B2, isolated from corn silage and anaerobic sewage sludge, respectively, were similar according to morphological observations, genome sequencing confirmed wide differences be-

TABLE 4 Codon usage of L. plantarum strains and phage B2 for amino acids encoded by the B2  $tRNAs^a$ 

			Frequen	cy of codon	usage (%) for:
Amino acid	Anticodon	Codon	Phage B1	Phage B2	L. plantarun WCSF1
Asn	GTT	AAC	16.9	19.3	17.5
		AAT	36.8	25.9	26.6
Leu	TAG	CTA	14.3	21.1	11.5
		TTA	17.6	42.5	33.4
		TTG	14.1	41.0	25.3
		CTT	9.4	13.6	8.9
		CTC	5.1	5.3	8.7
		CTG	11.2	21.9	12.3
Met	CAT	ATG	32.0	38.4	26.1
Gly	TCC	GGA	12.9	8.5	10.0
,		GGT	22.7	12.4	26.4
		GGC	25.2	5.3	17.3
		GGG	11.9	6.8	12.3
Arg	TCT	AGA	3.7	20.7	1.7
· ·	CCT	AGG	2.9	11.3	0.8
		CGT	12.6	8.7	11.7
		CGC	12.3	4.1	8.8
		CGA	6.6	6.6	7.1
		CGG	11.2	7.1	12.9

<sup>&</sup>lt;sup>a</sup> Codons indicated in boldface are those encoded by the tRNAs in the phage B2 genome.

tween the phages. Diversity among *Lactobacillus* phages, due possibly to the high number of species in the *Lactobacillus* genus, was reported previously (74). However, a relatively conserved genome organization among them was evidenced (74). Yet, *L. plantarum* phages appear to be among the most diverse *Lactobacillus* phages. Distinct ecological niches and unrelated host strains may explain such diversity.

As for many siphophages, the genome of phage B1 is organized into the following functional clusters: DNA packaging, morphogenesis, lysis, and DNA replication (Fig. 2). No genes/proteins related to lysogeny were found, confirming its virulent nature. Interestingly, a high level of identity (97%) with the genome of phage clP1, infecting Pediococcus damnosus, followed by 77% identity with the genome of L. plantarum phage JL-1, was found. Of note, the genome of phage clP1 showed a GC content of 47.6%, which is much higher than those reported for pediococci (37.8 to 41.2%) (35). When each ORF was analyzed, high levels of identity with phage clP1 deduced proteins (65 to 100%) were also observed, while the levels of identity with proteins of phage JL-1 were always lower (29 to 80%) (Table 2). Pediococcus and L. plantarum strains are often found in the same ecological niches (cucumber fermentation, silage inoculants) (34, 73); thus, these comparative analyses support the notion that coexistence in the same environment can lead to the exchange of genetic elements (45). Others have shown that phages of *L. plantarum* were able to infect strains of other bacterial species isolated from the same habitat (10, 20, 45), although this was not tested here. L. plantarum myophage LP65 unexpectedly infected Carnobacterium strains associated with fermented meat (10), and some L. plantarum phages isolated from silage and sauerkraut were able to infect Lactobacillus pentosus and Lactobacillus brevis strains (20, 45). On the other hand, phages B1 and B2 have a narrow host range, as reported for other *L. plantarum* phages (11, 44, 69).

The genomic organization of phage B2 was also similar to those of other siphophages (Fig. 3). Some ORFs exhibited homology with *L. plantarum* myophage LP65. However, most were similar to ORFs of *Bacillus* and *Lactobacillus* strains and their phages (Table 3). Few proteins (Orf39, Orf43, and Orf105) were linked to prophage proteins, but phage B2 had the growth characteristics of a virulent phage. This observation was also reported for *L. plantarum* phages LP65 (10), g1e (70), and Sha1 (72). Overall, the genome assemblage of phage B2 was rather unique and appears to be made from parts of other characterized phages.

Phage DNA packaging. The deduced B1 proteins Orf1 and Orf2 share high similarity with the putative small and large terminase subunits from various phages, including *P. damnosus* clP1 and *L. plantarum* g1e and phiJL-1. Phage B2 Orf17 and Orf21 exhibited sequence similarities to the small and large subunits of the terminases from *Bacillus subtilis* subsp. *natto* and *Lactobacillus delbrueckii* phage c5, respectively. Of note, this B2 genomic region was interrupted by 4 tRNAs. In tailed phages, the small terminase subunit is responsible for specific DNA binding whereas the large terminase subunit mediates the cleavage of concatameric phage DNA into genome units as well as prohead binding (26). In particular, the large subunit usually provides the endonuclease and ATPase activities for packaging (38).

The Orf59 gene product of phage B1 was associated with endonuclease function due to its homology with Orf12 of *Pediococcus* phage clP1 and Orf51 of *Lactobacillus casei* phage phiAT3. Taking into account the position of the gene in the phage B1 genome, this protein might also be involved in the DNA packaging or replication (43). In phage B2, Orf16 was identified as an HNH endonuclease, which could be involved in DNA packaging since it precedes the small terminase subunit. The HNH family of proteins is associated with DNA binding and cutting functions and includes some phage packaging proteins (47).

Phage DNA replication. Orf24 and Orf26 of phage B1 have several characteristics in common with endonucleases and helicases (NTP binding). Orf27 exhibited homology to DNA primases, Orf31 to replication proteins, Orf34 to replicases, and Orf35 to DNA binding proteins. A helicase function was also attributed to Orf37 since it shared 99% identity with the putative helicase from phage clP1 (P. damnosus). These seven proteins may be involved in DNA replication. The phage B2 proteins Orf45 and Orf47 exhibited similarities to the DNA polymerase III protein ( $\alpha$ subunit) from Bacillus phage SPBc2 (42). A DNA polymerase function was also attributed to Orf70. It is tempting to speculate that phage B2 encodes its own DNA polymerase instead of relying on its host. Helicase and DNA primase functions were attributed to Orf86 and Orf87, respectively. The protein product of ORF88 may be an exonuclease, and Orf71 may be linked to ATP/GTP binding proteins. Other B2 proteins may have roles in nucleotide modification (Orf65, Orf69, Orf83, and Orf114).

**Host lysis.** A key step of the phage infection process is the release of new virions at the end of the lytic cycle. Orf21 of phage B1 has similarities with the holins of *P. damnosus* phage clP1 and of *L. casei* phage AT3. It has a transmembrane domain in the N-terminal part similar to holins of *Lactobacillus rhamnosus* phages Lc-Nu and Lmr1 (24, 66). Orf22 exhibited sequence similarity to the endolysins from various phages and was classified an endo-*N*-acetylmuramidase (muramidase). For phage B2, the

endolysin function was attributed to ORF40 (muramidase-like endolysin) as well as Orf57 (transglycosylase). Similarly, two endolysins were encoded by the *L. plantarum* myophage LP65 genome (Orf88 and Orf121) (10). No recognizable gene encoding a holin was found for phage B2. Of the four classes of bacterial endolysins recognized (muramidase, tranglycosylase, amidase, and peptidase), two are commonly found in *Lactobacillus* phages (muramidase and amidase) (70). Moreover, similarities found among lysins of phages infecting several bacterial species could suggest a common evolutionary origin. Endolysins from phages LL-H (*Lactobacillus delbrueckii* subsp. *lactis*) and 0303 (*Lactobacillus helveticus*) were able to hydrolyze the cell walls of some species from *Lactobacillus* and *Pediococcus* (17, 67).

Structural proteins of phages B1 and B2. Analysis of phage B1 using SDS-PAGE revealed at least five protein bands (Fig. 4A). Band B was associated with one phage protein (Orf3, portal), whereas two phage proteins were identified in the other four bands. Band A contained a minor tail protein (Orf18) and, surprisingly, a putative DNA primase (Orf27). Band C was made of Orf43 and Orf22 (endolysin). Band D contained two capsid proteins (Orf4 and Orf6). Finally, bands B and E contained two tail proteins, Orf12 and Orf16. Orf27 (primase) and Orf22 (endolysin) are likely nonstructural proteins that were carried over despite the phage purification procedure. Overall, the observed molecular masses of the phage proteins matched the theoretical values (Fig. 4A). Proteomic analysis of the complete phage particle revealed four other proteins (Orf5, Orf15, Orf17, and Orf21). Orf5 and Orf15 likely correspond to the scaffold and the tape measure proteins, respectively.

For phage B2, significantly more protein bands were observed by SDS-PAGE (Fig. 4B). Except for protein band G, which contained two phage capsid proteins (Orf23 and Orf24), all Coomassie-stained bands contained only one phage protein. Orf24 (major capsid protein), with a calculated molecular mass of 45.8 kDa, was associated with three protein bands (F, G, and H), with estimated molecular masses of 45, 35, and 30 kDa, respectively. In fact, when the peptides from Orf24 in bands G and H were analyzed, it was found that the N-terminal peptides of the protein were missing. This suggested that the B2 major capsid protein was processed, a phenomenon observed for other phages (28, 40). Orf23, found in band G, shared homology with a major capsid protein from Bacillus and peptidase U35, which can be fused with capsid proteins (28). This putative peptidase activity may be involved in cleavage of Orf24. Orf36, associated with band B, showed homology with the tail fiber protein of phage Sha1 (L. plantarum). However, tail fibers were not observed in the morphology of phage B2 by electron microscopy (72). In total, nine structural proteins were identified for phage B2 (Fig. 4B). Analysis of the complete phage B2 particles did not reveal any additional structural proteins.

Conclusions. Lactobacillus phages are understudied compared to other industrially relevant lactic acid bacteria (18, 29). One possible reason is that there are fewer reports of Lactobacillus phage infections than of Lactococcus lactis and Streptococcus thermophilus infections in the food industry. It is unclear if this lack of reported Lactobacillus phage infections is due their specific uses or due to their intrinsic properties. Understanding this paucity of Lactobacillus phage infections in industrial settings may provide novel tools to control phage populations in other susceptible environments. Still, phages infecting several Lactobacillus species represent a risk for industrial users (6, 7, 10, 58, 70). Knowledge of

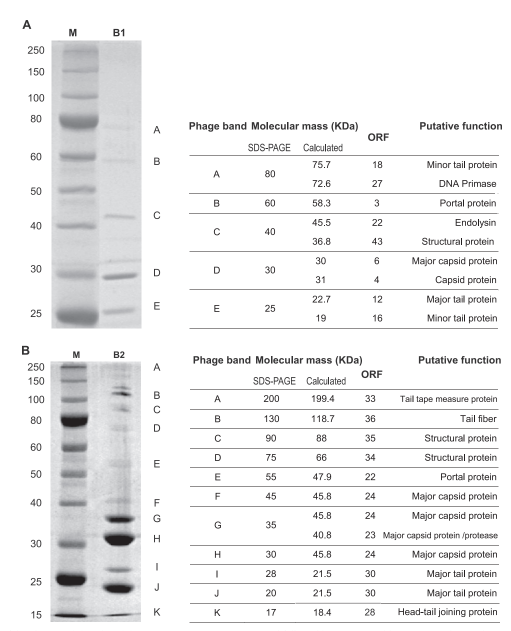


FIG 4 Migration of the phage B1 (A) and B2 (B) proteins on a 12% SDS-PAGE gel followed by Coomassie blue staining. The numbers on the left indicate the molecular masses of the ladder (protein ladder, 10 to 250 kDa; New England BioLabs). Letters on the right indicate bands cut out of the gel and identified by LC-MS/MS. Tables show the analysis of phage B1 and B2 structural proteins by LC-MS/MS.

their diversity is necessary to devise adapted control strategies. *L. plantarum* phages seem to have a relatively narrow host range, suggesting that strain rotation could be, whenever possible, an approach to limit phage multiplication. Moreover, some *L. plantarum* strains carry phage resistance mechanisms, which may be taken into account during the strain selection process. Comparative analysis of the phage B1 genome indicated that it is related to that of *L. plantarum* phage JL-1, suggesting that they form a phage group. On the other hand, analysis of the phage B2 genome suggested that this phage is currently unique among *L. plantarum* phages. The ever-increasing number of complete phage genome sequences has greatly improved our knowledge about phage diversity. The characterization of additional *L. plantarum* phages will help to determine the extent of their diversity.

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