

# Typing of bacteriophages by randomly amplified polymorphic DNA (RAPD)-PCR to assess genetic diversity

Diana Gutiérrez<sup>1</sup>, Antonio M. Martín-Platero<sup>2</sup>, Ana Rodríguez<sup>1</sup>, Manuel Martínez-Bueno<sup>2</sup>, Pilar García<sup>1</sup> & Beatriz Martínez<sup>1</sup>

<sup>1</sup>DairySafe Group, Instituto de Productos Lácteos de Asturias (IPLA-CSIC), Villaviciosa, Asturias, Spain; and <sup>2</sup>Departamento de Microbiología, Facultad de Ciencias, Universidad de Granada, Fuentenueva, Granada, Spain

Correspondence: Beatriz Martínez, DairySafe Group, Instituto de Productos Lácteos de Asturias (IPLA-CSIC), Apdo. 85, 33300-Villaviciosa, Asturias, Spain. Tel.: +34 985 89 33 59; fax: +34 985 89 22 33; e-mail: bmf1@ipla.csic.es

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#### Abstract

The recent boom in phage therapy and phage biocontrol requires the design of suitable cocktails of genetically different bacteriophages. The current methods for typing phages need significant quantities of purified DNA, may require a priori genetic information and are cost and time consuming. We have evaluated the randomly amplified polymorphic DNA (RAPD)-PCR technique to produce unique and reproducible band patterns from 26 different bacteriophages infecting Staphylococcus epidermidis, Staphylococcus aureus, Lactococcus lactis, Escherichia coli, Streptococcus thermophilus, Bacillus subtilis and Lactobacillus casei bacterial strains. Initially, purified DNA and phage suspensions of seven selected phages were used as a template. The conditions that were found to be optimal 8 uM of 10mer primers, 3 µM magnesium oxalacetate and 5% dimethyl sulfoxide. The RAPD genomic fingerprints using a phage titer suspension higher than 10<sup>9</sup> PFU mL<sup>-1</sup> were highly reproducible. Clustering by the Pearson correlation coefficient and the unweighted pair group method with arithmetic averages clustering algorithm correlated largely with genetically different phages infecting the same bacterial species, although closely related phages with a similar DNA restriction pattern were indistinguishable. The results support the use of RAPD-PCR for quick typing of phage isolates and preliminary assessment of their genetic diversity bypassing tedious DNA purification protocols and previous knowledge of their sequence.

### Introduction

Bacteriophages are ubiquitous in nature and found in all the habitats that their host bacteria colonize. It is now widely accepted that bacteriophages are the most abundant biological entities on Earth  $(10^{31} \text{ particles})$  (Brüssow & Kutter, 2005). They contribute largely to maintaining population densities and diversity of bacterial species, but also influence significantly biogeochemical and ecological processes including nutrient cycling, carbon flow and genetic transfer (Gill *et al.*, 2003; Thurber, 2009).

Classical bacteriophage taxonomy is based on their shape and size as well as their nucleic acid. Bacteriophages have been classified into 13 families; three of them (*Myoviridae*, *Siphoviridae* and *Podoviridae*) are members of the *Caudovirales* order that comprises about 96% of phages identified so far (5360 of 5568 reported to date, Ackermann,

2007). All these phages possess tail and double-stranded DNA.

The 500 bacteriophage genome sequences available at present in the NCBI phage database reveal the remarkable genetic diversity among phages, with genomes ranging from 15 up to 500 kb in size. Furthermore, bacteriophage genomes show a mosaic structure and each genome may be considered as a unique combination of modules whose size and rates of exchange vary considerably among the population. Nevertheless, despite the lack of similarity at the DNA level, phages encode proteins with significant sequence similarity, reflecting a common origin (Hendrix *et al.*, 1999). Recently, new phage classification schemes based on protein similarities have been developed for complementing the traditional classification (Lavigne *et al.*, 2008, 2009).

One of the main obstacles of phage biocontrol and phage therapy approaches is the narrow host range as a single

 Table 1. General features of bacteriophages, host bacteria and culturing conditions

		Bacteriophage	Genome		Propagation		
Phage	Host bacteria	family	size (kbp)	Life cycle	conditions	References	
vB_SepiS-philPLA4	S. epidermidis F12	Siphoviridae	39	Lytic	37 °C, shaking, TSB	D. Gutiérrez (unpublished data)	
vB_SepiS-philPLA5	S. epidermidis F12	Siphoviridae	39	Lytic	37 °C, shaking, TSB	Gutiérrez et al. (2010)	
vB_SepiS-philPLA6	S. epidermidis F12	Siphoviridae	38	Temperate	37 °C, shaking, TSB	Gutiérrez et al. (2010)	
vB_SepiS-philPLA7	S. epidermidis F12	Siphoviridae	33	Temperate	37 °C, shaking, TSB	Gutiérrez et al. (2010)	
vB_SauS-philPLA35	S. aureus Sa9	Siphoviridae	45.3	Lytic	37 °C, shaking, TSB	García <i>et al</i> . (2007)	
ΦH5	S. aureus Sa9	Siphoviridae	42.5	Temperate	37 °C, shaking, TSB	García et al. (2007)	
ФС2	L. lactis MG1614	Siphoviridae	22.1	Lytic	30 °C, static, GM17	Lubbers <i>et al</i> . (1995)	
Φ936	L. lactis IL1403	Siphoviridae	ND	Lytic	30 °C, static, GM17	Jarvis e <i>t al</i> . (1991)	
ΦΡ335	L. lactis F4.2	Siphoviridae	33.6	Lytic	30 °C, static, GM17	Braun <i>et al</i> . (1989)	
$\Phi$ FIPLA-1	S. thermophilus St5	Siphoviridae	35.2	Lytic	42 °C, static, GLM17	Magadán (2007)	
$\Phi$ FIPLA-3	S. thermophilus IPLA-10094	Siphoviridae	44.4	Lytic	42 °C, static, GLM17	Magadán (2007)	
$\Phi$ FIPLA-120	S. thermophilus IPLA-10094	Siphoviridae	45.5	Lytic	42 °C, static, GLM17	Magadán (2007)	
$\Phi$ FIPLA-122	S. thermophilus IPLA-10074	Siphoviridae	34.2	Lytic	42 °C, static, GLM17	Magadán (2007)	
ΦFIPLA-126	S. thermophilus LMD9	Siphoviridae	35.2	Lytic	42 °C, static, GLM17	Magadán (2007)	
ФА2	L. casei ATCC393	Siphoviridae	43.4	Temperate	37 °C, static, MRS	Herrero <i>et al</i> . (1994)	
ΦSPP1	<i>B. subtilis</i> 5B88G	Siphoviridae	44	Lytic	37 °C, static, 2 $\times$ YT	Riva <i>et al</i> . (1968)	
ΦSOM1	E. coli WG5	Siphoviridae	ND	Lytic	37 °C, shaking, 2 $\times$ YT	Muniesa <i>et al</i> . (1999)	
ΦSOM2	E. coli WG5	Myoviridae	ND	Lytic	37 °C, shaking, 2 $\times$ YT	Muniesa <i>et al</i> . (1999)	
ΦSOM4	E. coli WG5	Siphoviridae	ND	Lytic	37 °C, shaking, 2 $\times$ YT	Muniesa <i>et al.</i> (1999)	
ΦSOM7	E. coli WG5	Siphoviridae	ND	Lytic	37 °C, shaking, 2 $\times$ YT	Muniesa <i>et al.</i> (1999)	
ΦSOM8	E. coli WG5	Myoviridae	ND	Lytic	37 °C, shaking, 2 $\times$ YT	Muniesa <i>et al.</i> (1999)	
ΦSOM23	E. coli WG5	Siphoviridae	ND	Lytic	37 °C, shaking, 2 $\times$ YT	Muniesa <i>et al</i> . (1999)	
ΦSOM28	E. coli WG5	Siphoviridae	ND	Lytic	37 °C, shaking, 2 $ imes$ YT	Muniesa <i>et al</i> . (1999)	
ΦSCH10	E. coli WG5	Myoviridae	ND	Lytic	37 °C, shaking, 2 $ imes$ YT	Muniesa <i>et al</i> . (1999)	
ΦP1	E. coli WG5	Podoviridae	ND	Lytic	37 °C, shaking, 2 $ imes$ YT	Muniesa <i>et al</i> . (1999)	
ФХ174	E. coli WG5	Microviridae	ND	Lytic	37 °C, shaking, 2 $\times$ YT	Muniesa <i>et al</i> . (1999)	

ND, Not determined.

TSB, Tryptic soy broth (Scharlau Chemie S.A., Barcelona, Spain); GM17, M17 (Biokar, Beauvais, France) supplemented with glucose (0.5% p/v); GLM17, M17 supplemented with 0.5% glucose and 0.5% lactose; MRS, Man–Rogosa–Sharpe (Biokar); 2 × yeast extract and tryptone broth (YT), Sambrook *et al.* (1989).

phage may infect only specific strains. Thereby, the use of phage cocktails has been proposed (Sulakvelidze *et al.*, 2001). However, assessment of the genetic diversity among a large collection of phage isolates would require effective propagation of each phage to isolate enough DNA for sequencing or analysis of DNA restriction patterns, which is time consuming and not always successful. Thus, a quick and reproducible approach would be very valuable to type new phages whose genome sequences are unknown. Pioneering work has made use of fluorescence-labelled restriction fragment length polymorphism (fRFLP) to address bacteriophage typing (Merabishvili *et al.*, 2007).

Among other DNA-based approaches, random PCR amplifications of DNA segments using short primers of arbitrary nucleotide sequence have been used to generate specific profiles or genomic fingerprints that are used to compare the genotypic diversity among, for example, bacterial isolates (Johansson *et al.*, 1995; Guglielmotti *et al.*, 2006; Maiti *et al.*, 2009), or whole bacterial communities (Franklin *et al.*, 1999; Yang *et al.*, 2000). Randomly amplified polymorphic DNA (RAPD)-PCR using purified DNA has also been used to assess the genetic diversity of vibriophages (Comeau *et al.*, 2006; Shivu *et al.*, 2007) and phages infecting *Escherichia coli* (Dini & de Urraza, 2010) and *Pseudomonas aeruginosa* (Li *et al.*, 2010).

In this study, we have optimized a RAPD-PCR assay to evaluate whether reproducible patterns using phage lysates, instead of purified phage DNA, could be generated, as this would be more suitable for rapid screening of a high number of phage isolates.

# **Materials and methods**

#### Bacteriophage propagation and purification

Twenty-six bacteriophages were used in this study (Table 1). Phage propagation was performed in broth by infecting early exponential bacterial cultures supplemented with  $10 \text{ mM } \text{Ca}(\text{NO}_3)_2$  and  $10 \text{ mM } \text{MgSO}_4$ , at a multiplicity of infection of 1.0. Lysed bacterial cultures were centrifuged at  $10\,000 \text{ g}$ , the supernatants were filtered (0.45 µm, cellulose acetate membrane; VWR) and the phage titer was determined. Phage suspensions were dialyzed against distilled water for 1 h using 0.025-µm filters (MF-Millipore<sup>TM</sup> Membrane Filters; Millipore, Ireland) and stored at 4 °C.

Phage suspensions were also obtained from confluent lysis plaques on a solid medium. Appropriate phage dilutions were mixed with host bacteria in 0.7% top agar, poured on plates and incubated overnight. One milliliter of sterile-distilled water was added to plates and shaken for 1 h. The suspension was then centrifuged, and the supernatant was filtered and dialyzed as indicated above. Phage samples from both liquid and solid phage propagation were boiled for 10 min before the RAPD-PCR reaction.

Pure phage preparations were prepared by a CsCl continuous density gradient (Sambrook *et al.*, 1989). Briefly, 1 L of a bacterial lysate was centrifuged at 10 000 *g*. Phages were recovered from the supernatant by 10% polyethylene glycol8000 and 0.5 M NaCl precipitation. After centrifugation (13 000 *g*), phages were suspended in SM buffer (20 mg L<sup>-1</sup> Tris-HCl, 10 mg L<sup>-1</sup> MgSO<sub>4</sub>, 10 mg L<sup>-1</sup> CaCl<sub>2</sub>, 100 mg L<sup>-1</sup> NaCl, pH 7.5) containing RNase 40  $\mu$ g mL<sup>-1</sup>. Finally, phages were further purified by adding CsCl, followed by ultracentrifugation at 100 000 *g* at 4 °C for 20 h.

#### **Phage DNA isolation**

Phage DNA was extracted as described previously (García *et al.*, 2003) from 100  $\mu$ L of purified phage stocks previously dialyzed against SM buffer.

#### Genomic fingerprinting by RAPD analysis

Random amplification of polymorphic DNA was carried out according to a modification of the method described previously (Johansson *et al.*, 1995). Primers OPL5 (5'-ACGCAGGCAC-3'), RAPD5 (5'-AACGCGCAAC-3'), P1 (5'-CCGCAGCCAA-3') and P2 (5'-AACGGGCAGA-3') were assayed at three different concentrations (1, 4 and  $8 \mu M$ ).

PCR reactions were performed using PureTaq<sup>TM</sup> Ready-To-Go<sup>TM</sup> PCR Beads (GE Healthcare, Munich, Germany) adding 10 ng of purified phage DNA or  $10^7-10^8$  plaque forming units (PFU) of phage suspensions. Reactions were supplemented with 3 mM magnesium oxalacetate and/or 5% v/v dimethyl sulfoxide (DMSO). PCR was performed in a thermocycler (Bio-Rad, Hercules) under the following thermal cycling conditions: four cycles at 94 °C for 45 s, 30 °C for 120 s and 72 °C for 60 s; 26 cycles at 94 °C for 5 s, 36 °C for 30 s and 72 °C for 30 s (the extension step was increased by 1 s for every new cycle); and a final step of 10 min at 75 °C.

# Processing, comparison and reproducibility of RAPD patterns

DNA band patterns were obtained after gel electrophoresis (0.8% agarose gel) of the RAPD-PCR reaction products ( $15 \,\mu$ L). Gels were run for about 55 min at 100 V and stained

in ethidium bromide  $(0.5 \,\mu g \,m L^{-1})$  for 30 min. DNA molecular weight marker ('500 bp molecular ladder', Bio-Rad) was used as a standard. Gel images were processed using the software FINGERPRINTING II (Bio-Rad). The similarity matrix was calculated on the basis of the Pearson product-moment correlation coefficient, and its corresponding dendrogram was deduced using the unweighted pair group method with arithmetic averages [Struelens & the Members of the European Study Group on Epidemiological Markers (ESGEM), of the European Society for Clinical Microbiology and Infectious Diseases (ESCMID), 1996]. Reproducibility was assessed by cluster analysis of triplicate reactions.

## **Results and discussion**

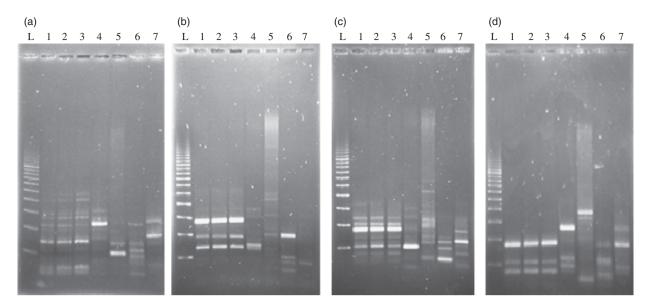
#### **RAPD-PCR** analysis of purified phage DNA

RAPD-based methods do not require sequence information for PCR primer design. However, they are extremely dependent on laboratory conditions such as template DNA concentration, PCR and electrophoretic settings, among others (Ellsworth *et al.*, 1993). To establish a quick and useful RAPD-PCR protocol to type phages, phages infecting strains belonging to the same species (four *Staphylococcus epidermidis* phages), or different species within the same genus (two *Staphylococcus aureus* phages) or a different genus (one *Lactococcus lactis* phage) were selected to test several experimental conditions in order to generate reproducible RAPD patterns and gain a preliminary insight into the discrimination power of this approach.

The selected S. epidermidis phages belonged to the Siphoviridae family (morphotype B1) and their genome sequences were unknown. However, previous DNA restriction analysis revealed distinct patterns for the temperate phages ØSepi-IPLA6 and ØSepi-IPLA7, while the DNA restriction patterns of the lytic phages ØSepi-IPLA4 and ΦSepi-IPLA5 (presumably virulent derivatives of ΦSepi-IPLA6) were very similar to each other (Gutiérrez et al., 2010; and our unpublished data). The two phages infecting S. aureus ΦH5 and vB\_SauS-phiIPLA35 (Φ35) belonged to morphotype B1 and morphotype B2, respectively, and their complete genome sequence was available (García et al., 2007, 2009). Finally, the lytic L. lactis phage  $\Phi$ C2 belonging to the morphotype B2 (Lubbers et al., 1995) was chosen as representative of phages infecting a different genus within Gram-positive bacteria.

Initially, pure phage DNA (10 ng) was used as a template. Because RAPD-PCR reactions are considerably influenced by primers and their concentration (Johansson *et al.*, 1995), four primers (OPL5, RAPD5, P1 and P2) at three different concentrations (1, 4 and 8  $\mu$ M) were tested. Furthermore, we tested whether the presence of magnesium oxalacetate and DMSO resulted in better defined band patterns. It has





**Fig. 1.** RAPD band patterns obtained from seven different bacteriophages using pure DNA as a template and primers OPL5 (a), RAPD5 (b), P1 (c) and P2 (d) at 8  $\mu$ M. Lane 1, vB\_SepiS-philPLA4; lane 2, vB\_SepiS-philPLA5; lane 3, vB\_SepiS-philPLA6; lane 4, vB\_SepiS-philPLA7; lane 5, vB\_SauS-philPLA35; lane 6,  $\Phi$ H5; and lane 7,  $\Phi$ C2; lane L, 500 bp molecular ladder. Reactions contained magnesium oxalacetate (3  $\mu$ M) and DMSO (5% v/v).

been described that  $Mg^{2+}$  ions form complexes with dNTPs, primers and template DNA, stimulating the action of DNA polymerase, and DMSO improves the DNA double-strand denaturalization and reduces secondary structures (Pomp & Medrano, 1991). Optimal results were obtained by the addition of 3 mM magnesium oxalacetate, 5% v/v DMSO and 8  $\mu$ M primer concentration (Fig. 1). Lower primer concentrations produced less defined bands for primers OPL5 and RAPD5, and no amplification for primers P1 and P2 (data not shown). Similar observations were reported previously when typing *Lactobacillus plantarum* strains by RAPD-PCR in which the optimal primer concentration was also 8  $\mu$ M (Johansson *et al.*, 1995).

As shown in Fig. 1, each primer generated distinct band patterns with amplicons ranging in size from approximately 500 bp to 12 kb. A total of 18 bands were observed for primer OPL5 (Fig. 1a), showing a greater discrimination among phages than the other primers that generated fewer (11–16) different bands (Fig. 1).

With the exception of *S. epidermidis* phages vB\_SepiSphiIPLA4, vB\_SepiS-phiIPLA5 and vB\_SepiS-phiIPLA6, which had shown a closely related DNA restriction pattern, the RAPD-PCR band profiles were unique for each phage (Fig. 1). It is worth noting that *L. lactis* phage  $\Phi$ C2 generated a small number of bands with all the primers assayed (Fig. 1, lane 7). Its lower genome size (22 163 bp) could explain this result (see Table 1).

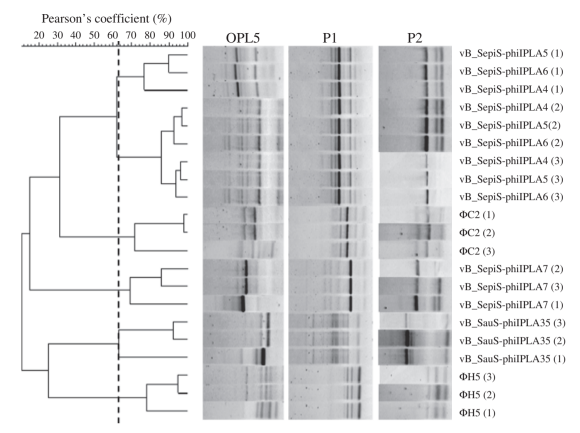
The genomic fingerprints resulting from the amplification of phage DNA samples performed on three separate days were compared to determine the RAPD-PCR reproducibility (Table 2). Each phage showed an identical band 
 Table 2. Reproducibility, indicated as Pearson's product correlation coefficient, of the RAPD-PCR reactions performed with different primers and templates

	Primer				
Template	OPL5 (%)	RAPD5 (%)	P1 (%)	P2 (%)	
Purified phage DNA Phage suspensions (liquid propagation) Phage suspensions (solid propagation)	95 < 20 > 90	58 < 20 25	28 < 20 > 90	92 < 20 60	

profile regardless of the assay date. Primers OLP5 and P2 provided high reproducibility values for genomic fingerprints and performed better than RAPD5 and P1. The low reproducibility of the later primers could be explained by the low number of amplification products obtained from phage  $\Phi$ C2 with RAPD5 (see Fig. 1). Moreover, differences in the band intensity on phage  $\Phi$ H5 DNA may have accounted for the low reproducibility of P1 (data not shown). No reproducible band intensities were likely due to nonspecific annealing between the primer and the DNA template as reported previously (Pérez *et al.*, 1998).

#### **RAPD-PCR** analysis of phage suspensions

Phage suspensions were evaluated as source of DNA template to avoid the phage DNA purification step. Phage propagation in liquid and solid culture media yielded a titer of  $10^7-10^8$  and  $> 10^9$  PFUmL<sup>-1</sup>, respectively, for all



**Fig. 2.** Dendrogram obtained after the analysis of RAPD band patterns generated with different sources of DNA and combining the primers OPL5, P1 and P2. The DNA source used as a template is indicated in each lane: (1) DNA isolated from purified phage suspensions, (2) phage suspensions from a liquid medium, (3) phage suspensions from a solid medium. The similarity between samples was calculated on the basis of the Pearson product–moment correlation coefficient and its corresponding dendrogram was constructed using the unweighted pair group algorithm method with arithmetic averages. The identity level for genotypes discrimination is represented by a dashed line.

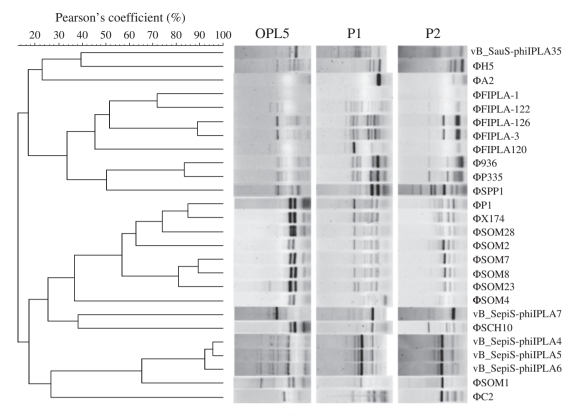
selected phages. To discard amplification from bacterial DNA, noninfected host bacterial cultures were processed under the same conditions as the phage lysates and used as a template in RAPD-PCR reactions. No amplification from host DNA was observed under the assay conditions (data not shown). Moreover, genomic fingerprints obtained using both phage lysates (from liquid and solid medium propagation) as a template were apparently similar to each other and to those obtained using pure DNA as a template (see Fig. 2).

The reproducibility of the assay using each template source with each single primer is shown in Table 2. In general, the RAPD profiles of phage suspensions from liquid propagation were poorly reproducible (< 20%) regardless of the primer used. By contrast, higher reproducibility values from phage suspensions obtained in a solid medium were recorded. Reproducibility seemed to be related to phage titer because suspensions from liquid propagation had 10–100 times less phages than those obtained from solid propagation ( $\geq 10^9 \text{ PFU mL}^{-1}$ ). We presume that the lower the phage titer, the lower DNA template is available for the PCR reaction, a factor that considerably influences the

performance of the RAPD-PCR reaction (Ellsworth *et al.*, 1993). Therefore, the low reproducibility of phage suspensions from liquid propagation is likely linked to variations in the initial phage titer. Moreover, a phage titer higher than  $10^9 \text{ PFU mL}^{-1}$  seems to be required to obtain a suitable reproducibility when using phage suspensions as a DNA source.

# Reproducibility analysis of RAPD-PCR combining type of template and primers OPL5, P1 and P2

A more detailed analysis was carried out comparing the genomic fingerprints generated from the three phage DNA sources with all three OPL5, P1 and P2 primers. RAPD5 was discarded due to the low reproducibility values obtained in the different assays. As shown in Fig. 2, the band patterns obtained from the different DNA templates clustered each phage together. As anticipated, the sensitivity of the RAPD-PCR assay was not enough to resolve the very close related *S. epidermidis* phages vB\_SepiS-phiIPLA4, vB\_SepiS-phiIPLA5 and vB\_SepiS-phiIPLA6. Still, our results support the



**Fig. 3.** Cluster analysis of the RAPD band patterns obtained from 26 different bacteriophages using the primers OPL5, P1 and P2. The similarity between samples was calculated on the basis of the Pearson product–moment correlation coefficient and its corresponding dendrogram was constructed using the unweighted pair group algorithm method with arithmetic averages.

use of sequence-specific 10-mer primers to reproducibly produce an adequate number of bands for the analysis of small genomes such as viruses. This is in accordance with previous reports showing that nondegenerate and degenerate 10-mer primers can produce robust band patterns for RAPD fingerprinting analysis (Comeau *et al.*, 2004; Winget & Wommack, 2008). In addition, pooling RAPD band patterns resulting from, at least, two different primers allows greater sensitivity.

### Validation of RAPD-PCR to type genetically diverse bacteriophages

According to our results, phage suspensions are also suitable to generate reproducible RAPD profiles, bypassing the need for isolating DNA. Consequently, RAPD-PCR could be a cost-effective and time-saving technique to assess the genetic diversity among phages. To validate its discriminatory power further, the RAPD-PCR assay was performed on a wide group of 26 phages infecting both Gram-positive and Gram-negative bacteria ranging from 33% to 50% in their G+C content. These phages belong to four different families (*Siphoviridae, Podoviridae, Myoviridae* and *Microviridae*). Phages infecting *L. lactis, Streptococcus thermophilus, Lacto*- *bacillus casei, Bacillus subtilis* and *E. coli* were used in the validation assay (Table 1). Genomic fingerprints were generated from phage suspensions after solid medium propagation using primers OPL5, P1 and P2 and the combined patterns were analyzed (Fig. 3).

The RAPD profiles were distinct for each phage and revealed the existence of four main clusters. These clusters matched largely with the bacterial species and most of the phages infecting the same bacterial species were clustered together, with few exceptions. Phages infecting *S. thermophilus* showed closed, but distinguishable patterns and slightly related to  $\Phi$ 936,  $\Phi$ P335 and  $\Phi$ SPP1. *Escherichia coli* phages also clustered together, except  $\Phi$ SOM1. Finally, *S. epidermidis* phages were also grouped, vB\_SepiS-phiIPLA7 being the exception.

This clustering was not surprising because of the phylogenetic relations among phages. As it has been described previously, phages infecting distantly related bacterial hosts typically share little or no nucleotide sequence similarity, while phages infecting a specific bacterial host are more similar (Hatfull, 2008). Moreover, module exchanging could be the reason why phages vB\_SepiS-phiIPLA7,  $\Phi$ C2 and  $\Phi$ SOM1 were grouped into a different cluster than the other phages infecting the same bacterial host. Phage morphology did not correlate with the RAPD-PCR clustering as phages belonging to different morphological families were grouped together. This is the case of  $\Phi$ X174 (*Microviridae*),  $\Phi$ P1 (*Podoviridae*),  $\Phi$ SOM8 and  $\Phi$ SOM2 (*Myoviridae*), which were clustered with the rest of the phages belonging to the *Siphoviridae* family. The classification in families is mostly based on virion morphology and nucleic acid type, and bacteriophages belonging to different families may have similar DNA sequences (Ackermann, 2003). Thereby, similar RAPD-PCR profiles can be found among families. A similar discrepancy has already been reported when using fRFLP for bacteriophage typing (Merabishvili *et al.*, 2007).

It remains to be confirmed whether RAPD typing using phage lysates is also a feasible technique when using phages infecting high G+C bacterial hosts as those were not included in this study. However, based on the use of DMSO in the reaction buffer and the availability of enhanced DNA polymerases and buffers active on high G+C DNA templates, it is reasonable to speculate that this approach may also be useful.

# Conclusions

RAPD-PCR on phage suspensions is a suitable approach to quickly assess the genetic diversity among newly isolated bacteriophages infecting the same species while circumventing the need for DNA extraction and purification. Using this assay, genomic fingerprints from different phages infecting *Staphylococcus*, *Bacillus*, *E. coli*, *Lactococcus* and *Streptococcus* were distinct and showed variations in the number of bands, fragment size and intensity.

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# Authors' contribution

P.G. and B.M. contributed equally to this work.

# References

- Ackermann HW (2003) Bacteriophage observations and evolution. *Res Microbiol* **154**: 245–251.
- Ackermann HW (2007) 5500 Phages examined in the electron microscope. *Arch Virol* **152**: 227–243.

- Braun V, Hertwig S, Neve H, Geis A & Teuber M (1989) Taxonomic differentiation of bacteriophages of *Lactococcus lactis* by electron microscopy, DNA–DNA hybridization, and protein profiles. *J Gen Microbiol* **135**: 2551–2560.
- Brüssow H & Kutter E (2005) Phage ecology. *Bacteriophages: Biology and Application* (Kutter E & Sulakvelidze A, eds), pp. 129–164. CRC Press, Boca Raton, FL.
- Comeau AM, Short S & Suttle CA (2004) The use of degenerateprimed random amplification of polymorphic DNA (DP-RAPD) for strain-typing and inferring the genetic similarity among closely related viruses. J Virol Methods 118: 95–100.
- Comeau AM, Chan AM & Suttle CA (2006) Genetic richness of vibriophages isolated in a coastal environment. *Environ Microbiol* **8**: 1164–1176.
- Dini C & de Urraza PJ (2010) Isolation and selection of coliphages as potential biocontrol agents of enterohemorrhagic and Shiga toxin-producing *E. coli* (EHEC and STEC) in cattle. *J Appl Microbiol* **109**: 873–887.
- Ellsworth DL, Rittenhouse D & Honeycutt RL (1993) Artifactual variation in randomly amplified polymorphic DNA banding patterns. *Biotechniques* 14: 214–217.
- Franklin RB, Taylor DR & Mills AL (1999) Characterization of microbial communities using randomly amplified polymorphic DNA (RAPD). J Microbiol Meth 35: 225–235.
- García P, Ladero V & Suárez JE (2003) Analysis of the morphogenetic cluster and genome of the temperate *Lactobacillus casei* bacteriophage A2. *Arch Virol* **148**: 1–20.
- García P, Madera C, Martínez B & Rodríguez A (2007) Biocontrol of *Staphylococcus aureus* in curd manufacturing processes using bacteriophages. *Int Dairy J* **17**: 1232–1239.
- García P, Obeso JM, Martínez B, Lavigne R, Lurz R & Rodríguez A (2009) Functional genomic analysis of *Staphylococcus aureus* phages isolated from the dairy environment. *Appl Environ Microb* 75: 7663–7673.
- Gill JJ, Svircev AM, Smith R & Castle AJ (2003) Bacteriophages of *Erwinia amylovora. Appl Environ Microb* **69**: 2133–2138.
- Guglielmotti DM, Reinheimer JA, Binetti AG, Giraffa G, Carminati D & Quiberoni A (2006) Characterization of spontaneous phage-resistant derivatives of *Lactobacillus delbrueckii* commercial strains. *Int J Food Microbiol* 111: 126–133.
- Gutiérrez D, Martínez B, Rodríguez A & García P (2010) Isolation and characterization of bacteriophages infecting *Staphylococcus epidermidis. Curr Microbiol* **61**: 601–608.
- Hatfull GF (2008) Bacteriophage genomics. *Curr Opin Microbiol* **11**: 447–453.
- Hendrix RW, Smith MC, Burns RN, Ford ME & Hatfull GF (1999) Evolutionary relationships among diverse bacteriophages and prophages: all the world's a phage. *P Natl Acad Sci USA* **96**: 2192–2197.
- Herrero M, de los Reyes-Gavilán CG, Caso JL & Suárez JE (1994) Characterization of φ393-A2, a bacteriophage that infects *Lactobacillus casei. Microbiology* **140**: 2585–2590.

Jarvis AW, Fitzgerald GF, Mata M, Mercenier A, Neve H, Powell IB, Ronda C, Saxelin M & Teuber M (1991) Species and type phages of lactococcal bacteriophages. *Intervirology* **32**: 2–9.

Johansson ML, Quednau M, Molin G & Ahrné S (1995) Randomly amplified polymorphic DNA (RAPD) for rapid typing of *Lactobacillus plantarum* strains. *Lett Appl Microbiol* **21**: 155–159.

Lavigne R, Seto D, Mahadevan P, Ackermann HW & Kropinski AM (2008) Unifying classical and molecular taxonomic classification: analysis of the *Podoviridae* using BLASTP-based tools. *Res Microbiol* 159: 406–414.

Lavigne R, Darius P, Summer EJ, Seto D, Mahadevan P, Nilsson AS, Ackermann HW & Kropinski AM (2009) Classification of *Myoviridae* bacteriophages using protein sequence similarity. *BMC Microbiol* 9: 224.

Li L, Yang H, Lin S & Jia S (2010) Classification of 17 newly isolated virulent bacteriophages of *Pseudomonas aeruginosa*. *Can J Microbiol* **56**: 925–933.

Lubbers MW, Waterfield NR, Beresford TP, Le Page RW & Jarvis AW (1995) Sequencing and analysis of the prolate-headed lactococcal bacteriophage c2 genome and identification of the structural genes. *Appl Environ Microb* **61**: 4348–4356.

Magadán AH (2007) Isolation, characterization and detection of *Streptococcus thermophilus* bacteriophages in the dairy industry. PhD Thesis, University of Oviedo, Spain.

Maiti B, Shekar M, Khushiramani R, Karunasagar I & Karunasagar I (2009) Evaluation of RAPD-PCR and protein profile analysis to differentiate *Vibrio harveyi* strains prevalent along the southwest coast of India. *J Genet* **88**: 273–279.

Merabishvili M, Verhelst R, Glonti T, Chanishvili N, Krylov V, Cuvelier C, Tediashvili M & Vaneechoutte M (2007) Digitized fluorescent RFLP analysis (fRFLP) as a universal method for comparing genomes of culturable dsDNA viruses: application to bacteriophages. *Res Microbiol* **158**: 572–581.

Muniesa M, Lucena F & Jofre J (1999) Study of the potential relationship between the morphology of infectious somatic

coliphages and their persistence in the environment. *J Appl Microbiol* **87**: 402–409.

Pérez T, Albornoz J & Domínguez A (1998) An evaluation of RAPD fragment reproducibility and nature. *Mol Ecol* 7: 1347–1357.

Pomp D & Medrano JF (1991) Organic solvents as facilitators of polymerase chain reaction. *Biotechniques* **10**: 58–59.

Riva S, Polsinelli M & Falaschi A (1968) A new phage of *Bacillus subtilis* with infectious DNA having separable strands. *J Mol Biol* **35**: 347–356.

Sambrook J, Maniatis T & Fritsch EF (1989) *Molecular Cloning: A Laboratory Manual*, 2nd edn. Cold Spring Harbor Laboratory Press, Cold Spring Harbor, New York.

Shivu MM, Rajeeva BC, Girisha SK, Karunasagar I, Krohne G & Karunasagar I (2007) Molecular characterization of Vibrio harveyi bacteriophages isolated from aquaculture environments along the coast of India. Environ Microbiol 9: 322–331.

Struelens MJ & the Members of the European Study Group on epidemiological Markers (ESGEM), of the European Society for Clinical Microbiology and Infectious Diseases (ESCMID) (1996) Consensus guidelines for appropriate use and evaluation of microbial epidemiologic typing systems. *Clin Microbiol Infec* **2**: 2–11.

Sulakvelidze A, Alavidze Z & Morris JG Jr (2001) Bacteriophage therapy. *Antimicrob Agents Ch* **45**: 649–659.

Thurber RV (2009) Current insights into phage biodiversity and biogeography. *Curr Opin Microbiol* **12**: 582–587.

Winget DM & Wommack KE (2008) Randomly amplified polymorphic DNA PCR as a tool for assessment of marine viral richness. *Appl Environ Microb* **74**: 2612–2618.

Yang YH, Yao J, Hu S & Qi Y (2000) Effects of agricultural chemicals on DNA sequence diversity of soil microbial community: a study with RAPD marker. *Microb Ecol* **39**: 72–79.