Characterization and genomic analysis of two *Staphylococcus aureus* bacteriophages isolated from poultry/livestock farms

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*Staphylococcus aureus* is one of the most important pathogens, causing various diseases in humans and animals. As methicillin-resistant *S. aureus* (MRSA) has become increasingly prevalent, controlling this pathogen with standard antibiotic treatment has become challenging. Bacteriophages (phages) have attracted interest as alternative antibacterial agents to control MRSA. In this study, we isolated six *S. aureus* phages from soils of poultry/livestock farms. Based on the results of host range determination with 150 *S. aureus* strains and restriction enzyme treatment of phage DNA, two phages, designated SP5 and SP6, were selected for further characterization and genome sequencing. Both SP5 and SP6 were classified as members of the family *Siphoviridae*. The genome of SP5 comprises 43,305 bp and contains 63 ORFs, while the SP6 genome comprises 42,902 bp and contains 61 ORFs. Although they have different host spectra, the phage genomes exhibit high nucleotide similarity to each other. Adsorption assay results suggested that the host range determinants of the two phages are involved in both adsorption and infection. Comparative genomic analyses of the two phages provided evidence that the lysogenic/lytic control module and tail proteins may be important for host specificity.

**INTRODUCTION**

*Staphylococcus aureus* is usually commensal on human skin, but it is one of the most important pathogens, causing illnesses ranging from minor skin infections to life-threatening diseases such as pneumonia and bacteraemia (Mann, 2008). As well as humans, livestock animals are also infected with *S. aureus*, resulting in bovine mastitis and chicken arthritis, which threaten food safety and cause food poisoning (Vanderhaeghen et al., 2010). Notably, the increasing incidence of antibiotic-resistant *S. aureus* strains, such as methicillin-resistant *S. aureus* (MRSA) and vancomycin-resistant *S. aureus* (VRSA), has become a major concern, because *S. aureus* is one of the most common causes of nosocomial infections (DeLeo & Chambers, 2009; Dulon et al., 2011).

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The GenBank/EMBL/DDBJ accession numbers for the complete genome sequences of SP5 and SP6 are JX274646 and JX274647, respectively.

Four supplementary tables and two figures are available with the online version of this paper.

*S. aureus* bacteriophages have been investigated extensively. All sequenced *S. aureus* bacteriophages are grouped into three classes based on genome size: class I (~20 kb), class II (~40 kb) and class III (>125 kb) (Kwan et al., 2005). Bacteriophages belonging to class III have been characterized as members of the family *Myoviridae* and are obligate lytic phages with relatively wide host ranges (Kwan et al., 2005; Mann, 2008). Obligate lytic phages have received renewed interest as potential therapeutic agents to replace or supplement antibiotics in the treatment of MRSA or other antibiotic-resistant *S. aureus* strains. Clinical trials performed in Eastern Europe showed that bacteriophage therapy was effective in the treatment of many *S. aureus*-related diseases, including lung infection, sepsis and leg ulcers (Kutateladze & Adamia, 2010; Rhoads et al., 2009). Temperate phages, on the other hand, have been regarded to be pathogenically important because they often carry virulence genes in *S. aureus*. For example, phage 80x can mobilize a variety of superantigen-encoding pathogenicity islands (SaPIs) (Tallent et al., 2007), and another major group of virulence factors, the Panton–Valentine leukocidin...
genes, are encoded by prophages such as φPVL and φSLT (Narita et al., 2001; Zou et al., 2000).

In this paper, we report isolation of S. aureus bacteriophages from soil samples from a poultry farm and a cattle shed, and present the results of microbiological and molecular characterization of two selected phages that can be utilized in developing biocontrol agents against staphylococcal contamination.

RESULTS AND DISCUSSION

Isolation of bacteriophages infecting S. aureus

Bacteriophages infecting S. aureus are potential biological control agents that can prevent staphylococcal food poisoning. In order to apply isolated bacteriophages to control S. aureus infection by contaminated foods in human and livestock, we first isolated and characterized phages with strong lytic activity against S. aureus strains. Six S. aureus bacteriophages were isolated from six soil samples collected from poultry and livestock farms, and designated SP1–SP6. Based on host specificities and restriction enzyme digestion profiles, these six phages were categorized into two groups. SP5 and SP6 were selected as representatives of these two groups and characterized in depth.

Host specificity of six phages

The host specificities of the six isolated phages were determined using 10 S. aureus reference strains and 140 isolates originated from humans, livestock, vegetables and foods (Table 1). SP1, SP2, SP4 and SP5 (SP5-like phages) showed identical host specificities, lysing 29 isolates (20.7 %); SP3 and SP6 (SP6-like phages) shared different host specificities, lysing 33 of 140 isolates (23.6 %). Only 18 isolates were sensitive to both phages. None of the six phages could lyse other bacterial species, including Staphylococcus epidermidis [ATCC 35983, Culture Collection of Antimicrobial Resistant Microbes (CCARM) 3787 and 3789], Listeria monocytogenes (ATCC 19115), Bacillus subtilis (ATCC 23857), Bacillus cereus (ATCC 14579), Pseudomonas aeruginosa (ATCC 27853) and Escherichia coli (MG1655, ATCC 35150) (data not shown).

Restriction enzyme digestion profiles

For genotyping analysis between the isolated phages, genomic DNA was extracted from the six phages and treated with various restriction enzymes. Comparison of the restriction fragments by agarose gel electrophoresis showed that SP1, SP2, SP4 and SP5 shared the same profile, while SP3 and SP6 had an alternative profile (Fig. 1).

Phage morphologies

All the isolated phages were visualized by transmission electron microscopy (TEM) (Fig. 2). All phages exhibited hexagonal heads (diameter ~60 nm) and non-contractile, flexible tails (length ~200 nm), which are representative features of bacteriophages belonging to the family Siphoviridae (order Caudovirales). The tail of each phage contained a baseplate with spikes at the end.

Differences between SP5 and SP6 in adsorption to their hosts

Based on the results of host range determination and genotyping analyses, SP5 and SP6 were chosen for further studies below. The difference in host specificity between SP5 and SP6 may be attributable to bacterial host determinants such as receptors as well as phage features. Interestingly, the 11 bacterial isolates sensitive only to SP5-like phages were human- or livestock-derived, suggesting the possibility of the presence of common genetic features among these 11 isolates. To gain insight into the host range determination, multiple analytical tools including diagnostic PCR and pulsed-field gel electrophoresis (PFGE) were applied to S. aureus isolates to examine their genetic relationships and virulence factors. Strong genetic diversity was high among bacteria isolates; there was little correlation between host genetic diversity and phage host range determination (data not shown).

In order to find evidence to explain the difference in host specificity between SP5 and SP6, an adsorption assay was performed (Fig. 3). Both phages adsorbed efficiently to their common host, S. aureus ATCC 29213, as expected. For S. aureus isolate 112, which could be infected by SP5 only, the adsorption affinity of SP5 was higher than that of SP6, suggesting that SP6 phage might be lacking the ability to attach to the host surface. However, for isolate 96, which

Table 1. Lytic activity of isolated phages towards S. aureus strains

| Strains tested | SP1, SP2, SP4 and SP5 | SP3 and SP6 |
|---------------|----------------------|-------------|
| ATCC 29213, CCARM 3089 and 18 isolates | + | + |
| CCARM 3090 and 11 isolates | + | – |
| 15 isolates | – | + |
| ATCC 25904 (Newman), ATCC 12600, ATCC 25923, ATCC 6538, ATCC 23235, ATCC 27664, CCARM 3793 and 96 isolates | – | – |
is sensitive to SP6 only, the adsorption affinity of SP5 was comparable to that of SP6, suggesting that SP5 might fail in intracellular replication or escaping the host cell after invasion. These adsorption assay results indicate that host range determinants are associated not only with the adsorption stage (isolate 112) but also with the stage after adsorption (isolate 96) (Fig. 3).

**Genome overviews of SP5 and SP6 phages**

For a better understanding of the different host specificity among phages, genomic comparison between SP5 and SP6 was carried out. The genome of SP5 comprised 43,305 bp and contained 63 putative ORFs (Table S1, available in JGV Online); the genome of SP6 comprised 42,902 bp and contained 61 putative ORFs (Table S2). The G+C contents of SP5 and SP6 were 34.4 and 34.5 mol%, respectively, slightly higher than that of the sequenced *S. aureus* strains (32.8%). Neither phage seemed to contain tRNA-coding regions, according to tRNAscan-SE v.1.21 analysis. Neither known virulence factors, including staphylokinase, exfoliative toxin A, enterotoxin A, Panton–Valentine leukocidin, nor the innate immune modulators CHIPS (chemotaxis inhibitory protein of *S. aureus*) and SCIN (staphylococcal complement inhibitor), were found in either SP5 or SP6 (Betley & Mekalanos, 1985; Narita et al., 2001; van Wamel et al., 2006; Winkler et al., 1965; Yamaguchi et al., 2000).

The overall genome organizations of SP5 and SP6 are compared in Fig. 4. All of the ORFs, except for the first three, are transcribed rightward in both genomes. Both genomes are organized into three major gene clusters: (i) the 'lysogeny' control region; (ii) the 'early' region, encoding products involved in replication, recombination and modification of the phage DNA; and (iii) the 'late' region, encoding structural and assembly proteins and lysis proteins.

BLASTN analyses revealed that SP5 and SP6 had high nucleotide similarity with other known *S. aureus* phages including 80s (GenBank accession no. DQ517338), φ11 (GenBank accession no. AF424781), 53 (GenBank accession no. AY954952) and 69 (GenBank accession no. AY954951) phages. The query coverage was from 65 to 73% and homologous sequences were conserved at the latter parts of SP5 (>12 kb) and SP6 (>12.5 kb) genomes. Interestingly, both phages also exhibited high sequence similarity with the φ37-like prophage region of *S. aureus* 11819–97 (GenBank accession no. CP003194) with 89–91% query coverage (Stegger et al., 2012) (Table S3).

**Comparative genomics of SP5 and SP6**

SP5 and SP6 bacteriophages appeared to be temperate, rather than virulent, phages belonging to class II *S. aureus* phages (Kwan et al., 2005) based on genome analyses but also had strong lytic activities showing ~10^6-fold reduction in bacterial number for 3 h infection (data not shown). Genome sequencing of SP5 and SP6 revealed that these two phages shared the highest sequence similarity to each other (query coverage >90%), albeit there were also other phages or prophages with high nucleotide similarity. Among the 63 ORFs of SP5 and 61 ORFs of SP6, 53
**Fig. 3.** Adsorption of SP5 and SP6 to their host strains. *S. aureus* ATCC 29213 was sensitive to both phages, while human isolates 112 and 96 were only susceptible to SP5 and SP6, respectively. Data are presented as the means ± SD of triplicate assays.

**Fig. 4.** Genomic maps of phages SP5 and SP6. Predicted ORFs are denoted by arrows, and genes encoding proteins with at least 60% amino acid identity between the two genomes are indicated by shaded regions.
ORFs were highly conserved in both phages: 32 ORFs had 100% identities; 19 ORFs had >90% identities; and two ORFs had >70% amino acid identities (Fig. 4, Table S4). Comparative genomic analysis between the two phage genomes provided deeper insights into their characteristics including the different host spectra.

The ‘lysogenic’ control region

Comparing the phage genomes of SP5 and SP6, the most divergent part lay in the ‘lysogenic’ control region. Although the first three ORFs (integrase to CI protein) showed high identities (76–100%) between the two genomes, the subsequent ORFs (Cro-like protein to orf8 in SP5/orf9 in SP6) showed no such similarities to each other (Tables S3 and S4). The functions of many ORFs in this divergent region are unknown, but this region appears to be important for regulation of phage lysis/lysogeny because it includes proteins such as Cro-like repressor, phage antirepressor (orf6 in SP5 and SP6), and a helix–turn–helix domain-containing protein (orf5 in SP5). While CI is the regulatory protein that stabilizes the lysogenic state, Cro makes lytic growth and the lytic pathway possible by repressing CI (Little, 2007). An antirepressor protein (orf6) of SP5 and SP6 may be responsible for inactivation/bypass of the CI transcription repressor (García et al., 2009). In addition, the nucleotide sequences of the intergenic regions between cl and cro, which contain promoter regions for cl and cro, are totally different. This indicates that the mechanisms regulating the switch between the lysogenic and lytic states may differ between the two phages, and may be affected by different signals or regulator proteins.

Computer-aided promoter prediction (http://linux1.softberry.com/berry.phtml) of the intergenic space between the cl and cro genes suggested differential transcriptional regulation by a variety of regulators in the two phages (Fig. S1). Features included consensus RpoD-binding sites upstream of cl and cro in SP6 and cl in SP5, but not in the upstream region of cro gene in SP5, and an Lrp-binding site upstream of cro in SP5. This computational inference suggests the possibility that the transcription of cro in SP5 by σ70 may be repressed by Lrp, and that alternative regulatory factors may be required to induce the lytic pathway in the host. Lrp, a global regulator, interferes with open complex formation in σ70- or σS-containing RNA polymerase holoenzymes at promoters and inhibits the initiation of transcription (Colland et al., 2000). Thus, this difference in the lysogenic/lytic regulation module may be the determinant of the different host specificities between SP5 and SP6, and may explain why phage SP5 could not lyse S. aureus isolate 96, despite equivalent adsorption affinities for isolate 96 between the two phages, although this possibility needs to be tested (Fig. 3).

The ‘early’ region

The regions from orf9 to orf32 in SP5 and from orf10 to orf28 in SP6 appeared to be included in the ‘early’ region. This region seemed to contain genes related to DNA replication and metabolism. Included in this region are the DNA recombination protein RecT (orf15 in SP5 and orf16 in SP6), the ssDNA-binding protein Sb (orf17 in SP5 and orf18 in SP6), the crossover junction endoDNaSe RusA (orf20 in SP5 and orf21 in SP6) and dUTP diphosphatase (orf26 in SP5 and orf27 in SP6), all of which appeared to function during DNA recombination. All the ORFs belonging to this region were conserved in the genomes of SP5 and SP6, with extremely high amino acid identities. However, five ORFs (orf27 and orf29–orf32) were found only in the genome of SP5. These ORFs have high similarity to proteins belonging to S. aureus subsp. aureus Mu50. However, the functions of these five ORFs remain unknown.

The ‘late’ region

This region contains the structural and lysis modules. The structural module can be divided into packaging, head, head–tail connection and tail submodules. This region starts from rinB (orf33 in SP5 and orf29 in SP6) and ends at the termini of the genomes. As shown in Fig. 4, all ORFs in the ‘late’ region were conserved in both genomes, with high amino acid sequence identity (>96%), except the rightward accessory region (orf60 and orf61 in SP6), which was unique to the genome of SP6.

This region starts with the rinAB regulator genes. RinA is reported to be the major regulator of phage-mediated packaging, capsid assembly and cell lysis in phage 11, whereas RinB does not appear to control the expression of ‘late’ genes (Ferrer et al., 2011). Among ORFs belonging to the structural module, orf59 of SP5 and orf55 of SP6 were annotated as tail-associated cell-wall hydrolases. Tail-associated cell-wall hydrolase is localized at the tail tip and allows contact with the cell during adsorption/DNA injection in phage infection (Rashel et al., 2008). However, the tail-associated cell-wall hydrolases from SP5 and SP6 had 100% identity to each other, ruling out the possibility of tail tip hydrolase being responsible for host specificity.

To date, no report on determinants of host specificity of S. aureus phages has been published. However, with respect to other phages, the best-known determinant is the tail tip of phages, by which bacteriophages recognize receptors and inject DNA (Letellier et al., 2007; Tétart et al., 1996; Werts et al., 1994). Since the adsorption affinities of two phages were clearly distinguished by an SP5-specific host (isolate 112) (Fig. 3), we considered the possibility that tail fibre proteins may control host specificity. When the amino acid sequences of tail fibre proteins were compared among phages SP5 and SP6, and another 14 phages with tail fibre proteins of similar sizes (~390 aa) from Staphylococcus prophages and staphylococcal phages in the family Siphoviridae, moderate sequence diversities at four loci in the C-terminal region were found (each locus ~20 aa), whereas N-terminal portions (the N-terminal 240 aa) were highly conserved among 16 tail fibre proteins (Fig. S2). Host specificity among Staphylococcus phages may be

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attributable to the sequence diversity in the C-terminal portion of tail fibre proteins, which is implicated in binding to host receptors. Enterobacteria phage λ utilizes tail fibre protein J in binding to the LamB receptor on the outer membrane of E. coli and the C terminus of tail fibre protein J (20% distal C-terminal portion) was found to be important for the interaction of the phage with the host receptor protein in vitro and in vivo (Hofnung et al., 1976; Werts et al., 1994).

The lysis modes of both phages comprise the canonical holin–endolysin cassette. The amino acid sequences of the holin and endolysin proteins of SP5 (orf62 and orf63) and SP6 (orf58 and orf59) exhibited 96% identities. These endolysin proteins harbour a CHAP (cysteine-histidine-dependent amidohydrolases/peptidases) (PF05257, cysteine peptidase) domain and an amidase (PF01510) domain as catalytic domains at the N terminus, and an SH3_5 (PF08460) domain for substrate-binding at the C terminus.

**Conclusion**

In conclusion, our data demonstrated that two S. aureus siphoviruses isolated from a poultry farm and a cattle shed have high similarity in their genomic sequences but different host specificities. Comparative genomic analyses of these phages suggested the possibility that the difference in host specificities might be caused by different lysogenic/lytic control mechanisms and different tail fibre proteins between the two phages. Although further studies to support this conclusion are necessary, the results described here improve our understanding of the genomes of S. aureus phages.

**METHODS**

**Bacterial strains, growth conditions and bacteriophage methods.** The bacterial strains were acquired from the American Type Culture Collection (ATCC) and CCARM (Culture Collection of Antimicrobial Resistant Microbes, Seoul, Republic of Korea). S. aureus ATCC 29213 was routinely used to propagate isolated bacteriophages after purification and to determine the titres of phage stocks. Other strains used for host range determination are shown in Table 1 or are described in Results and Discussion. One hundred and forty S. aureus isolates were obtained from the collection of the Rural Development Administration (Suwon, Republic of Korea). These isolates originated from humans (37 isolates), livestock and their products (40 isolates), vegetables (54 isolates) and gimbap (Korean rice rolled in laver paper) (nine isolates).

All bacterial strains were routinely cultivated at 37 °C using tryptic soy broth (TSB) or tryptic soy agar (TSA) (Difco). CaCl₂ (10 mM) and MgCl₂ (10 mM) were added when bacteriophages were propagated. Spotting and overlay assays were routinely performed to form plaques and determine phage titres using TSA plates (0.5% agar for the top, 1.5% agar for the bottom) containing CaCl₂ (10 mM) and MgCl₂ (10 mM) according to methods described previously (Kronpinski et al., 2009; Mazzocco et al., 2009). SM buffer (50 mM Tris/HC1 (pH 7.5), 100 mM NaCl, 10 mM MgSO₄) was used to dilute phage stocks. Plaque formation was verified after incubation at 37 °C for 16 h.

**Isolation of bacteriophages.** Six soil samples were collected from a poultry farm and a cattle shed. Bacteriophages were isolated as described previously (Kim & Ryu, 2011). Briefly, samples (25 g) were homogenized in 25 ml homogenization buffer (0.25 M KH₂PO₄ adjusted to pH 7.2 with NaOH) for 90 s in a BagMixer 400 blender (Interscience Laboratory). Initial propagation was performed with the filtrate of each homogenized sample and a mixture of overnight cultures of 140 S. aureus isolates. Bacteriophages were isolated and purified by overlay assays (repeated three times) by picking plaques and elution in SM buffer.

**Propagation and purification of bacteriophages.** S. aureus ATCC 29213 was employed to propagate bacteriophages routinely. Bacteriophage particles were precipitated with polyethylene glycol 6000 (10%) and NaCl (1 M) at 4 °C overnight. After centrifugation (10 000 g, 20 min, 4 °C), precipitated bacteriophages were resuspended in SM buffer, and purified by CsCl density gradient ultracentrifugation (75 000 g, 2 h, 4 °C). Separated bacteriophages were dialysed against 1000 vols TM buffer (50 mM Tris/HC1 (pH 7.5), 20 mM MgSO₄) for 2 h with one buffer change. The titre of each purified bacteriophage was determined by overlay assay (Kronpinski et al., 2009). Phage stocks were stored in glass tubes at 4 °C.

**Host range determination.** The host range of each phage was determined by spot test. Lysis zone or plaque formation was monitored upon application of 10 µl of each phage lysate, adjusted to contain 1 x 10⁷ p.f.u. ml⁻¹, by spotting assay.

**Bacteriophage DNA extraction and restriction.** Phage DNA was extracted by an alkaline lysis method (Wilcox et al., 1996). Phage DNA (1 µg) was digested with 20 U restriction enzymes (Takara Biomedical) for 16 h according to the manufacturer’s instructions, and was resolved in a 1% agarose gel.

**TEM.** Purified bacteriophages were visualized by TEM. TEM was conducted as described previously (Kim & Ryu, 2011). Based on their morphology, phages were identified and classified according to the guidelines of the International Committee on Taxonomy of Viruses (Fauquet et al., 2005).

**Genome sequencing and in silico analysis.** Genomic DNA was isolated from phages SP5 and SP6 by the alkaline lysis method (Wilcox et al., 1996). A pyrosequencing approach, using a Genome Sequencer FLX System (Titanium series; Macrogen) was applied, and the high-quality filtered reads were assembled into a complete genome sequence using GS De novo Assembler (v.2.6). ORFs were predicted with Glimmer 3 (Delcher et al., 2007), GeneMarkS (Besemer et al., 2001) and FgenesB (http://linux1.softberry.com/). BBS finder (Suzek et al., 2001), which facilitates determination of the start codon of each ORF, was used to predict ribosome-binding sites. Annotation of predicted ORFs was performed by analysing the results of a similar protein search using BLASTP (Altschul et al., 1990) and a motif search using InterProScan (Quevillon et al., 2005). A search for tRNA-coding regions was performed using tRNAscan-SE v.1.21 (Schattner et al., 2005).

**Adsorption assay.** Adsorption of the phages was assayed as follows. Overnight cultures of the bacterial strains were diluted 1:100 in tryptic soy broth (TSB). When the OD of the reference strains at 600 nm reached 2.0, 1 ml of each culture was diluted 10-fold in fresh TSB. SP5 and SP6 phases were added at an m.o.i. of 0.01 to each diluted culture, which contained 10 mM CaCl₂ and 10 mM MgCl₂, mixed gently, and incubated at 37 °C for 15 min. An aliquot (100 µl) was removed immediately for determination of the initial phage titre. Incubation was continued for 15 min, and samples (100 µl) were
collected after 1, 5, 10 and 15 min and diluted immediately in 900 µl cooled SM buffer. The diluted samples were centrifuged at 14 000 g for 1 min at 4 °C and filtered using 0.22 µm filters. Finally, the titres of unabsorbed phages in the supernatant were determined after serial dilution.

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REFERENCES

Altschul, S. F., Gish, W., Miller, W., Myers, E. W. & Lipman, D. J. (1990). Basic local alignment search tool. J Mol Biol 215, 403–410.

Besemer, J., Lomsadze, A. & Borodovsky, M. (2001). GeneMarkS: a self-training method for prediction of gene starts in microbial genomes. Implications for finding sequence motifs in regulatory regions. Nucleic Acids Res 29, 2607–2618.

Betley, M. J. & Mekalanos, J. J. (1985). Staphylococcal enterotoxin A is encoded by phage. Science 229, 185–187.

Colland, F., Barth, M., Henge-Aronis, R. & Kolb, A. (2000). Sigma factor selectivity of Escherichia coli RNA polymerase: role for CRP, IHF and lrp transcription factors. EMBO J 19, 3028–3037.

Delcher, A. L., Bratke, K. A., Powers, E. C. & Salzberg, S. L. (2007). Identifying bacterial genes and endosymbiont DNA with Glimmer. Bioinformatics 23, 673–679.

DeLeo, F. R. & Chambers, H. F. (2009). Reemergence of antibiotic-resistant Staphylococcus aureus in the genomics era. J Clin Invest 119, 2464–2474.

Dulon, M., Haamann, F., Peters, C., Schablon, A. & Nienhaus, A. (2011). MRSA prevalence in European healthcare settings: a review. BMC Infect Dis 11, 138.

Fauquet, C., Mayo, M. A., Maniloff, J., Desselberger, U. & Bale, L. A. (editors) (2005). Virus Taxonomy: Classification and Nomenclature of Viruses: Eighth Report of the International Committee on the Taxonomy of Viruses. San Diego: Elsevier Academic Press.

Ferrer, M. D., Quiles-Puchalt, N., Harwich, M. D., Tormo-Más, M. À., Campoy, S., Barbé, J., Lasa, I., Novick, R. P., Christie, G. E. & Penadés, J. R. (2011). RinA controls phage-mediated packaging and transfer of virulence genes in Gram-positive bacteria. Nucleic Acids Res 39, 5866–5878.

Garcia, P., Martinez, B., Obeso, J. M., Lavigne, R., Lurz, R. & Rodriguez, A. (2009). Functional genomic analysis of two Staphylococcus aureus phages isolated from the dairy environment. Appl Environ Microbiol 75, 7663–7673.

Hofnung, M., Jezierska, A. & Braun-Breton, C. (1976). lamB mutations in E. coli K12: growth of lambda host range mutants and effect of nonsense suppressors. Mol Gen Genet 145, 207–213.

Kim, M. & Ryu, S. (2011). Characterization of a T5-like coliphage, SPC35, and differential development of resistance to SPC35 in Salmonella enterica serovar typhimurium and Escherichia coli. Appl Environ Microbiol 77, 2042–2050.

Kronpinski, A. M., Mazzocco, A., Waddell, T. E., Lingohr, E. & Johnson, R. P. (2009). Enumeration of bacteriophages by double agar overlay plaque assay. In Isolation and Characterization, and Interaction (Bacteriophages: Methods and Protocol vol 1), pp. 69–76. Edited by M. R. Clokie & A. M. Kropinski. New York: Humana Press.

Kutateladze, M. & Adamia, R. (2010). Bacteriophages as potential new therapeutics to replace or supplement antibiotics. Trends Biotechnol 28, 591–595.

Kwan, T., Liu, J., DuBow, M., Gros, P. & Pelletier, J. (2005). The complete genomes and proteomes of 27 Staphylococcus aureus bacteriophages. Proc Natl Acad Sci U S A 102, 5174–5179.

Letellier, L., Plancon, L. & Boulanger, P. (2007). Transfer of DNA from phage to host. In Bacteriophage: Genetics and Molecular Biology, pp. 208–209. Edited by S. McGrath & D. Van Sinderen. Norfolk, UK: Caister Academic Press.

Little, J. W. (2007). Prophage induction of phage. In Bacteriophage: Genetics and Molecular Biology, pp. 251–272. Edited by S. McGrath & D. Van Sinderen. Norfolk, UK: Caister Academic Press.

Mann, N. H. (2008). The potential of phages to prevent MRSA infections. Res Microbiol 159, 400–405.

Mazzocco, A., Waddell, T. E., Lingohr, E. & Johnson, R. P. (2009). Enumeration of bacteriophages using the small drop plaque assay system. In Isolation and Characterization, and Interaction (Bacteriophages: Methods and Protocol vol 1), pp. 81–85. Edited by M. R. Clokie & A. M. Kropinski. New York: Humana Press.

Narita, S., Kaneko, J., Chiba, J., Piémont, Y., Jarraud, S., Etienne, J. & Kamo, Y. (2001). Phage conversion of Panton–Valentine leukocidin in Staphylococcus aureus molecular analysis of a PVL-converting phage. phiSLT. Gene 268, 195–206.

Quevillon, E., Silventoinen, V., Pillai, S., Harte, N., Mulder, N., Apweiler, R. & Lopez, R. (2005). InterProScan: protein domains identifier. Nucleic Acids Res 33 (Web Server issue), W116–W120.

Rashel, M., Uchiyama, J., Takemura, I., Hoshiba, H., Ujihara, T., Takatsui, H., Honke, K. & Matsuizaki, S. (2008). Tail-associated structural protein gp61 of Staphylococcus aureus phage phiMR11 has bifunctional lytic activity. FEMS Microbiol Lett 284, 9–16.

Rhoads, D. D., Wolcott, R. D., Kuskowski, M. A., Wolcott, B. M., Ward, L. S. & Sulakvelidze, A. (2009). Bacteriophage therapy of venous leg ulcers in humans: results of a phase I safety trial. J Wound Care 18, 237–243, 248–240.

Schattner, P., Brooks, A. N. & Lowe, T. M. (2005). The tRNAscan-SE, snocan and snoGPS web servers for the detection of tRNAs and snoRNAs. Nucleic Acids Res 33 (Web Server issue), W686–W689.

Stegger, M., Price, L. B., Larsen, A. R., Gilleece, J. D., Waters, A. E., Skov, R. & Andersen, P. S. (2012). Genome sequence of Staphylococcus aureus strain 11819-97, an ST80-IV European community-acquired methicillin-resistant isolate. J Bacteriol 194, 1625–1626.

Suzek, B. E., Ermolaeva, M. D., Schreiber, M. & Salzberg, S. L. (2001). A probabilistic method for identifying start codons in bacterial genomes. Bioinformatics 17, 1123–1130.

Tallent, S. M., Langston, T. B., Moran, R. G. & Christie, G. E. (2007). Transducing particles of Staphylococcus aureus pathogenicity island SaP11 are comprised of helper phage-encoded proteins. J Bacteriol 189, 7520–7524.

Tétart, F., Repoila, F., Monad, C. & Krisch, H. M. (1996). Bacteriophage T4 host range is expanded by duplications of a small domain of the tail fiber adhesin. J Mol Biol 258, 726–731.

van Wamel, W. J., Rooijakkers, S. H., Ruyken, M., van Kessel, K. P. & van Strijp, J. A. (2006). The innate immune modulators staphylococcal complement inhibitor and chemotaxis inhibitory protein of

http://vir.sgmjournals.org
Staphylococcus aureus are located on beta-hemolysin-converting bacteriophages. *J Bacteriol* 188, 1310–1315.

Vanderhaeghen, W., Hermans, K., Haesebrouck, F. & Butaye, P. (2010). Methicillin-resistant *Staphylococcus aureus* (MRSA) in food production animals. *Epidemiol Infect* 138, 606–625.

Werts, C., Michel, V., Hofnung, M. & Charbit, A. (1994). Adsorption of bacteriophage lambda on the LamB protein of *Escherichia coli* K-12: point mutations in gene J of lambda responsible for extended host range. *J Bacteriol* 176, 941–947.

Wilcox, S. A., Toder, R. & Foster, J. W. (1996). Rapid isolation of recombinant lambda phage DNA for use in fluorescence *in situ* hybridization. *Chromosome Res* 4, 397–404.

Winkler, K. C., de Waart, J., Grootsen, C., Zegers, B. J. M., Tellier, N. F. & Vertregt, C. D. (1965). Lysogenic conversion of staphylococci to loss of beta-toxin. *J Gen Microbiol* 39, 321–333.

Yamaguchi, T., Hayashi, T., Takami, H., Nakasone, K., Ohnishi, M., Nakayama, K., Yamada, S., Komatsuzawa, H. & Sugai, M. (2000). Phage conversion of exfoliative toxin A production in *Staphylococcus aureus*. *Mol Microbiol* 38, 694–705.

Zou, D., Kaneko, J., Narita, S. & Kamio, Y. (2000). Prophage, phiPV83-pro, carrying Panton–Valentine leukocidin genes, on the *Staphylococcus aureus* P83 chromosome: comparative analysis of the genome structures of phiPV83-pro, phiPVL, phi11, and other phages. *Biosci Biotechnol Biochem* 64, 2631–2643.