Large Preferred Region for Packaging of Bacterial DNA by phiC725A, a Novel Pseudomonas aeruginosa F116-Like Bacteriophage

Christine Pourcel *, Cédric Midoux, Yolande Hauck, Gilles Vergnaud, Libera Latino

Institute for Integrative Biology of the Cell (I2BC), CEA, CNRS, Univ. Paris-Sud, Université Paris-Saclay, Gif-sur-Yvette, France

* christine.pourcel@gmail.com

Abstract

Bacteriophage vB_PaeP_PAO1_phiC725A (short name phiC725A) was isolated following mitomycin C induction of C7-25, a clinical Pseudomonas aeruginosa strain carrying phiC725A as a prophage. The phiC725A genome sequence shows similarity to F116, a P. aeruginosa podovirus capable of generalized transduction. Likewise, phiC725A is a podovirus with long tail fibers. PhiC725A was able to lysogenize two additional P. aeruginosa strains in which it was maintained both as a prophage and in an episomal state. Investigation by deep sequencing showed that bacterial DNA carried inside phage particles originated predominantly from a 700-800kb region, immediately flanking the attL prophage insertion site, whether the phages were induced from a lysogen or recovered after infection. This indicates that during productive replication, recombination of phage genomes with the bacterial chromosome at the att site occurs occasionally, allowing packaging of adjacent bacterial DNA.

Introduction

Some bacteriophages are capable of transduction, as shown by their capacity to transfer a DNA fragment from one bacterium to another, resulting in acquisition of new genetic information. As opposed to specialized transduction in which transduction is limited to host DNA adjacent to prophage insertion site [1], generalized transduction (GT) is defined as a process in which any host gene can be transferred to another bacterium. Mu-like phages perform transduction following transposition at multiple sites in the bacterial chromosome, in the course of their replicative cycle [2, 3]. Other transducing phages perform rolling-circle replication and use packaging initiation (pac) sites for packaging their genome via a headful mechanism [4]. It has been suggested that cryptic pac site on the host chromosome may lead to packaging of bacterial DNA by such phages [5–7].

The most studied phages performing GT are Escherichia coli phage P1 [8] and Salmonella enterica phage P22 [9]. Phage F116, also capable of GT, was isolated from a Pseudomonas...
**Funding:** This study was supported by the French ‘direction générale de l’armement’ (DGA) through Agence Nationale de la Recherche (ANR, France) “Resisphage” ANR-13-ASTRID-0011-01 (http://www.agence-nationale-recherche.fr/), and a joint DGA-IDEX Université Paris-Saclay doctoral fellowship to Dr. Libera Latino. The Transposon Mutant Collection holds grant #NIH P30DK089507. The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

**Competing Interests:** The authors have declared that no competing interests exist.

*aeruginosa* clinical strain [10]. It was reported to maintain its DNA as a plasmid [11] and it was not clear whether it could integrate into the bacterial genome [12]. F116 binds to the type IV pili of its host [13] and can digest alginate [14]. Upon sequencing of its genome (65,195bp, genome accession number AY625898), it was shown that 2.6% of the packaged DNA was of bacterial origin, predominantly derived from a particular region of the chromosome [15]. Two F116-like *P. aeruginosa* temperate phages have been sequenced, H66 (65,270bp, genome accession number KC262634) and LKA5 (64,746bp, genome accession number KC900378). In addition, several *P. aeruginosa* strains were shown to possess a prophage with up to 98% DNA sequence similarity with F116 [16]. Based on genome and protein comparison these phages are recognized as a homology group [17].

Here we describe a novel F116-like phage induced from a clinical *P. aeruginosa* strain which, together with previously described phages and prophages, contribute to the forming of a new genus sharing common characteristics. We demonstrate that the phage preferentially packages bacterial DNA localized on one side of its insertion site, not only when recovered by induction of the prophage, but also after an infection cycle.

**Materials and Methods**

**Ethics statement**

The present project is in compliance with the Helsinki Declaration (Ethical Principles for Medical Research Involving Human Subjects). Bacterial strains were previously collected as part of the patients’ usual care, without any additional sampling for the present investigation [18–20]. The ethic committee “Comité Consultatif pour la Protection des Personnes dans la Recherche Biomédicale (CCPPRB) Ile-de-France”, who was consulted, specifically approved the study, and declared that patient informed consent was not needed.

**Strains and media**

Strain C7-25 was isolated from a cystic fibrosis patient and was previously investigated as part of a project on antibiotic resistance [20] and on susceptibility to bacteriophages [21]. Its genome has been totally sequenced and is presently under analysis (Pourcel et al. unpublished). PcyII-10 and 26 additional clinical strains were from Percy Hospital, Clamart, France. Phages vB_PaeP_PA01_Ab05 (Ab05), vB_PaeM_PA01_Ab17 (Ab17) and vB_PaeM_PA01_Ab27 (Ab27) were described in [22]. PAO1 LPS and type IV pilus transposon mutants were obtained from “The *P. aeruginosa* Transposon Mutant Library”, UW Genome Sciences, USA. Luria broth (LB) medium supplemented with 2 mM CaCl$_2$ was used for bacterial growth and phage titration. Saline magnesium (SM) buffer (50 mM Tris-HCl pH7.5, 100 mM NaCl, 8.1 mM MgSO$_4$, 0.01% gelatin) was used to preserve purified phages at 4˚C.

**Phage production**

To induce bacteriophages from C7-25, 20 ml LB medium were inoculated at 1/100 with an overnight C7-25 culture and shaken at 37˚C until the culture reached an $A_{600}$ of 0.6. Mitomycin C was added to a final concentration of three μg ml$^{-1}$ and the incubation was continued until lysis occurred. Culture supernatant was tested for the presence of bacteriophages by plating five μl of different dilutions after infection of a PAO1 strain culture. A single plaque was recovered and purified by three successive platings. The newly isolated phage phiC725A was amplified for 8 hours on solid medium, by mixing 10$^8$ cfu of PAO1 with 10$^6$ pfu of phiC725A per plate, and collected as previously described [23]. Bacteria and debris were pelleted by centrifugation, the supernatant was recovered and phages were precipitated with 10%...
polyethylene glycol (PEG) 8000 overnight at 4 °C. After centrifugation at 15,000xg for 20 min, the pellet was suspended in one ml of phosphate-buffered saline (PBS) and treated for two hours with DNase I (50 μg ml⁻¹) at 37 °C. Three chloroform extractions were performed, prior to filtration through a 0.45μM filter and centrifugation for two hours at 260,000xg. The pellet was suspended in 50 μl of PBS. For electron microscopy (EM) visualization, five μl of phage suspension were stained with 2% potassium phosphotungstate (pH 7.0) as previously described [21].

DNA extraction and PCR

Samples were lysed in lysis buffer (Tris 10 mM, pH 7.8, EDTA 10 mM, NaCl 10 mM, SDS 0.5%wt/vol), treated with proteinase K at 50 μg ml⁻¹ for 2 hours at 50 °C, followed by one phenol and one chloroform extraction, and ethanol precipitation.

Phage insertion was detected by PCR performed on purified DNA. Amplification using primer pairs C725-Ins-F 5’ TGCCGACGTCCGGCTTCA3’ and C725-Ins-R 5’ CGATGGTTT TACCGLAAGTCG3’ resulted in an amplification product if a prophage was inserted at the bacterial att site (attB). Amplification using primer pair phi725-Reg2-F 5’ GAATTTGAGCGG AAACAACTA3’ and phi725-Reg2-R 5’ CCAATTGGCGTGCTTCTGCGG3’ present inside the phage genome on both sides of the phage attachment site (attP) was possible if an intact attP site was present, as in non-integrated phage genomes.

Whole genome sequencing

Two μg of purified phage or bacterial DNA were sent for draft whole genome sequencing to the IMAGIF MiSeq Illumina platform (CNRS, Gif sur Yvette, France). Libraries were made from sheared fragments of DNA (average size 900bp), and 250bp or 300 bp paired-end reads were produced. Phage genome assemblies, sequencing reads mapping to genome reference, prophage insertion sites and other sequence analyses were done using tools in Geneious R9 (Biomatters, New Zealand).

Nucleotide sequence accession numbers

The DNA sequence of phage phiC725A has been deposited in the EMBL-EBI database under accession number LT603684 within project PRJEB14922. Sequencing reads from the phage induction and infection experiments as well as sequencing reads from the PAO1 or lysogen for phiC725A were also deposited in project PRJEB14922 and are available from the European Nucleotide Archives (ENA) browser at http://www.ebi.ac.uk/ena/data/view/PRJEB14922.

Results

Isolation of phiC725A

We observed that strain C7-25 spontaneously released phages, as evidenced by the presence of small plaques on a lawn of bacteria after several days of growth on solid LB medium. Upon treatment of planktonic cells with mitomycin C, bacterial growth was rapidly stopped and massive lysis was observed. A bacteriophage forming small clear plaques with a dark center was isolated from the culture supernatant and amplified on a reference strain PAO1 representative called PAO1 or (the genome sequence of the PAO1 representative used in our laboratory in Orsay (Or) was previously reported [24]). Upon examination by EM, the phages displayed a large head (78nm), and what seemed to be a short thin tail (66nm) (Fig 1). No evidence of tail flexibility or contractility could be seen from EM observation of numerous phage particles. A similar morphology was previously reported for phage F116 and it was recently proposed to
classify F116 as a podovirus with tightly packed long fibers [15, 25, 26]. The new phage was called vB_PaeP_PAO1_phiC725A (short name phiC725A), for prophage A of strain C7-25. The phage host range was tested by infecting 26 clinical strains, including the reference strains PAO1 (PAO1\textsuperscript{Or}) and PA14. Two were susceptible (PAO1 and PcyII-10) and six showed a reduced efficiency of plating. The rest were totally resistant. In order to identify the receptor for phage phiC725A, different amounts of phages (from $10^4$ to $10^7$ pfu) were spotted onto PAO1 transposon mutants affected in type IV pilus genes (\textit{pilY1}, \textit{pilR}, \textit{pilA} and \textit{pilQ}) and LPS synthesis genes (\textit{wzy}, \textit{algC}). No growth was observed on mutants lacking pili whereas normal growth was seen on LPS mutants.

**The phiC725A phage genome**

PhiC725A was amplified on PAO1\textsuperscript{Or}, then phage particles were purified and treated with DNase I before extracting the DNA for sequencing. To be able to characterize the phage genome but also the bacterial DNA packaged within the phage head, deep sequencing of the phage genome was performed so that an average coverage of almost 15,000X, was achieved. A total of 1,284,508 reads were obtained, of which 0.6% (7,844) mapped to PAO1\textsuperscript{Or}. The phage reads were assembled resulting in a single 65,149bp long molecule with a 63.5% GC content. BLAST analysis on Genbank (release 214) identified F116 (genome accession number AY625898.1) as the closest phage genome with a 86% mean similarity over the whole sequence. PhiC725A showed 84% and 76.5% similarity with phage LK5 (KC900378.1) and H66 (KC262634.1) respectively. Similarities were also found along the genomes of prophages of \textit{P. aeruginosa} strains N01-01092 (81.8%, CP012901.1), DHS01 (76.5%, CP013993.1) and H27930 (72.7%, CP008860.1). In keeping with the reference phage F116 genetic map, the first
nucleotide was assigned as the first nucleotide downstream the int gene (Fig 2A). Annotation led to the identification of 62 putative coding sequences (CDSs) (S1 Table) globally matching those of F116, H66 and LKA5 with some remarkable differences which are most likely the result of recombination events, as shown by alignment of the four phage genomes (S1 Fig). The region encompassing phiC725A_01 to phiC725A_09 showed the highest level of dissimilarity (S1 Fig). PhiC725A_54, a very large protein had a homologue in H66, LKA5 and the three prophages and corresponded to F116p59 and F116p60. In phage H66 this protein is described as a structural protein. By homology search it was shown to possess a domain matching within the internal virion protein D (IVP-D) of Pseudoalteromonas phage Prio-1, corresponding to the C-terminal domain of IVP N4 gp50, a very large multisubunits enzyme [17]. N4 gp50 is an RNA polymerase which central part holds the transcriptionally active domain [27]. Such proteins, injected into the host together with the viral DNA, transcribe early genes [28, 29]. In the central part of phiC725A_54 lies a domain matching to the S-adenosyl methionine binding site of DNA methylases. PhiC725A and LKA5 lacked a short CDS (F116p41) present in the F116 genome in between the portal gene (phiC725A_37) and a putative capsid gene (phiC725A_38). PhiC725A, like F116, possesses genes that are common to many podoviruses [17]. Some are characteristic of T7-like podoviruses with tubular tails, but the tail fibers of F116-like phages, as visualized by electron microscopy seem to be longer and suppler.
A 17bp attachment site (attP: TGGTGCCGGACGGAGAGA) was identified at position 62,750 upstream of the integrase of phiC725A and was identical to a sequence in between tRNA-His and tRNA-Leu in PAO1 (attB in position 1,947,646–1,947,662 of PAO1 accession number AE004091, and 3,568,180 to 3,568,196 of PAO1 Or accession number LN871187.1). It corresponded to the site of the prophage insertion in strain C7-25 (position 3,680,043–3,744,208 in the C7-25 genome, Pourcel et al. unpublished). The same insertion site was observed for F116-like prophages in P. aeruginosa strains N01-01092 and H27930. In contrast, in strain DHS01 (accession number CP013993.1) the prophage was present at position 2,550,071–2,615,626 corresponding to 2,363,267–2,363,274 of PAO1 Or (accession number LN871187.1).

The insertion was flanked by a 11 bp direct repeat sequence TCCATCATCGG. F116 has been described as a non-integrative phage [11], but it possesses a putative integrase, identical to that of phiC725A (F116p70) and the TGGTGCCGGACGGAGAGA attP site, suggesting that this phage could in theory lysogenize strain PAO1. The 17 bp attB site sequence identified here was present in 93 out of the 95 P. aeruginosa complete genome sequences which could be queried in Genbank (the site was absent in strains ATCC 27853 and Cu1510).

When the full dataset of phage reads was mapped back on the phiC725A sequence (Fig 2B), no high peak of reads characteristic of fixed genome ends was observed [22]. This is as expected if the phage genome is terminally redundant (TR) and circularly permuted (CP), as previously demonstrated for F116 [30]. However, we noticed a maximum in the coverage starting at position 23,500 (near the small and large terminase subunit genes) and decreasing over the whole genome (Fig 2B). This behavior might be related to the phage replication or recombination, which leads to the generation of free ends. It could also indicate the position of the packaging initiation of the phage genome, with no clear-cut starting point, i.e. a lack of specificity in cleavage initiation. Indeed there are numerous examples of pac sites residing in or near the terminase genes in different phage genomes [31, 32]. Because of the terminal redundancy, a portion of the genome starting from the packaging position will be overrepresented.

**Bacterial DNA packaged into phiC725A virions**

We then examined the 0.6% of host genome reads obtained from sequencing purified bacteriophages. Surprisingly, we found that 73% (5717 among 7844 bacterial reads) mapped over a region corresponding to the attB site, with a maximum coverage at the attB position and extending over 700–800 kb on one side only of the attB site (S2 Fig). The remaining reads of bacterial origin were homogeneously distributed along the whole genome. We detected six hybrid reads containing a attP/B site, suggesting that during phage replication some genomes inserted within the bacterial chromosome at attB site, leading to packaging of adjacent DNA.

We consequently decided to analyse the phages directly purified following mitomycin C induction of strain C7-25. Bacteriophages were treated with DNAse I to eliminate contaminating bacterial DNA, before lysis and extraction of DNA. Sequencing of 1,834,389 DNA fragments with an average fragment size of 680 bp, resulted in the production of 3,668,778 sequence reads. Most reads corresponded to phage sequence, achieving an average coverage of the phage genome of more than 25,000X. However, 81,578 reads, or 2.2% of the total amount or reads, corresponded to the bacterial genome. 55,000 reads of these (67%) mapped over the 700–800 kb region on the left side of the prophage insertion att site with a regular decay, similarly to what was observed during infection (Fig 3A). The average coverage was 60–70X in the vicinity of the attL site and was down to 1.8X, 790 kb away. The remaining 26,578 reads of bacterial origin were distributed along the whole genome achieving a mean coverage of 1.3X. On
close examination of the reads distribution (Fig 3B), it appeared that it formed waves of about 65-75kb, slightly larger than the size of the phage genome.

More than 25,000 sequence reads contained the att sequence in agreement with the phage genome coverage. In the vast majority of cases, it was the attP from bona fide encapsidated phage genomes, but fifty-six reads contained an attP/B site. Only hybrids corresponding to the left side of the attB site (attL) were recovered, showing that bacterial DNA packaging at this location was unidirectional and initiated within a copy of the phage genome integrated at the attB site (Fig 3B).

Consequently, the majority of transduced bacterial DNA appears to be associated with the encapsidation of phage linked with the attB site in both the infection and induction experiments. The only detectable difference is a three-fold higher ratio of packaged host DNA in the induction of strain C7-25 (2.2% +/- 0.015 (P<0.05) versus 0.6% +/- 0.014 (P<0.05) for infection of strain PAO1. These differences are highly significant (X² test p-value < 2.2e-16).
Production of lysogens using phage phiC725A

PhiC725A was able to lysogenize both PAO1ₕₕ (PAO1ₕₕ phiC725A) and PcyII-10 (PcyII-10ₕₕ phiC725A) as shown by the stable presence of the phage DNA tested by PCR, continuous production of phages (10⁸ pfu ml⁻¹ in the culture supernatant), and exclusion of super-infection by phage phiC725A. In order to test for heteroimmunity, the lysogens were challenged with three different phages which grew on the parental PAO1ₕₕ and PcyII-10 strains. Ab05, a phi-KMV-like phage which uses type IV pili as receptor, produced plaques on the PAO1ₕₕ phiC725A lysogen, although of smaller size, and grew poorly on the PcyII-10ₕₕ phiC725A lysogen. Ab17 and Ab27, which use LPS as receptors, grew normally. This suggested that phiC725A lysogens were not heteroimmune but restricted infection by phages using pili as a receptor.

To check whether the phage was inserted into the bacterial genome, we performed a PCR reaction for attP/B with primer pair C725-Ins containing one primer flanking the att site on one side in the bacterial genome and one primer on the other side inside the phage genome. A signal was obtained with the C7-25 strain used as positive control and with both PAO1ₕₕ phiC725A and PcyII-10ₕₕ phiC725A indicating that the prophage integrated at the same attB site (data not shown). The (control) parent strains PAO1ₕₕ and PcyII-10 were negative. A PCR reaction with phage primers localized on both sides of attP, aiming at detecting non-integrated genomes (primer pair Phi725-Reg2) was also positive with all three lysogens, indicating that free viral genomes were present in the cells (data not shown).

Discussion

We describe phiC725A, a novel F116-like virus of P. aeruginosa, and provide evidence that the phage can integrate its genome at a specific site in a tRNA genes locus, and also remains as an episome. Lysogens continuously produce phages. In both prophage induction and infection, we show that phiC725A packages bacterial host DNA originating preferentially from one side of the attB site, and expanding over several hundreds of kilobases. This portion of the genome does not correspond to a region of high sequence diversity as described by Spencer at al. [33], but is one of the region of genomic plasticity (RGP) described by Mathee et al. [34]. Indeed at this position a phage was found inserted in P. aeruginosa strain PA2192. It is known that tRNA genes often serve as insertion sites for phages and other integrative elements. Genes present in the 800kb region preferentially packaged within phages are supposedly involved in different metabolic functions and some might play a role in virulence. Immediately flanking the attB site is the his-tRNA, then the parRS operon, encoding the two-component regulatory system ParR-ParS, involved in adaptive resistance to polymyxin B and colistin [35]. A gene encoding a membrane-bound lytic murein transglycosylase (mltD), a virulence-related factor that may participate in beta-lactam resistance is located a few kilobases away [36].

Importantly, we also show that packaging of this region is associated with the presence of a phage genome inserted at the att site as demonstrated by the existence of hybrid reads. It appears that packaging of bacterial DNA at the attL site takes place during phage replication but is three times less frequent during infection as compared to activation of a prophage. A similar and fully compatible observation of a biased origin of packaged host DNA was made during sequencing of F116, but the extent of bacterial chromosome and the distribution of reads could not be mapped with such precision [15]. In particular, the link with the att site and the evidence for the occasional integration of phage genome could not be detected with the 4X phage genome coverage achieved at that time, and the authors instead hypothesized that the bias might result from the presence of one or more cryptic pac sites in this region. Massively parallel sequencing used here allowed to achieve the much higher coverage necessary for the finding of hybrid phage-bacteria sequences, demonstrating the role of integrations in the att
site. Its close similarity to F116 suggests that phiC725A is likely to replicate by rolling circle amplification of circular phage genome. It is expected that virions encapsidate phage DNA from these concatemers using a headful packaging strategy, therefore producing circularly permuted genomes with a terminal redundancy [37, 38]. A pac site inside the phage provides the first cut, and headful DNA packaging is then performed. In phages P22 and SPP1, the small terminase subunit TerS binds to the pac site and recruits TerL that cleaves the DNA, initiating packaging [4, 32]. Packaging of host DNA flanking a P22 excision-defective prophage on one side was reported by Youderian et al. [39, 40]. In several temperate phages found in Staphylococcus aureus strain RF122, Moon et al. showed that TerS and TerL proteins may bind to the pac site of the prophage and initiate packaging of the phage DNA and adjacent DNA [41]. This resulted in the mobilization of a genomic island by the temperate bacteriophage. Huang et al. inserted pac sites in the E. coli chromosome and showed that P1 packaging of bacterial genes to one side of and near to a pac site was increased more than 10-fold. This effect diminished with distance but packaging could be detected over 30% of the length of the chromosome [5].

The sequence data presented here indicates that occasionally, during replication and expression of late genes, phage genome recombines with the bacterial host genome at the attB site thus providing a pac site. This happens following activation of a prophage in a lysogen, as well as during the lytic cycle in an infected bacteria, although at a lower frequency. The pac recognition event will produce one virion containing part of the phage genome together with bacterial genome flanking the att site, followed by up to ten-twelve virions (800 kb equivalent taking into account 10% terminal redundancy) containing bacterial DNA (Fig 3B). The absence of clear-cut boundaries suggests that there is no specific cut during packaging. The decay profile of bacterial DNA coverage further indicates that, on average, one event of packaging from an integrated phage will result in three virions containing bacterial DNA. It is also possible that a concatemer of phage DNA is formed at the attB site, or more likely sometimes recombines with the attB site after replication and prior to packaging. As a consequence, the population of hybrid virions, i.e. containing phage DNA and bacterial DNA, as well as the extent of encapsidation of bacterial genome, will be heterogeneous. If many phage genomes are attached downstream the packaging site, no or only few “bacterial” virions will be produced and vice-versa.

Conclusion
PhiC725A is a novel F116-like temperate phage capable of transporting bacterial DNA at a high frequency. We show for the first time that bacterial DNA is packaged preferentially from one side of the prophage integration site not only during prophage induction but also during infection, indicating that packaging can occasionally be initiated in a prophage-like configuration and will then extend to neighboring DNA. This implies that general transduction by phiC725A is highly biased which may have interesting consequences in terms of P. aeruginosa genome evolution.

Supporting Information
S1 Fig. Alignment of four F116-like phage genomes; phiC725A, F116, LKA5, H66 using the Geneious software.
(TIFF)

S2 Fig. Distribution of packaged bacterial reads from PAO1 or infected by phiC725A. Mapping of the bacterial reads along the PAO1 or genome. The blue, black and yellow represent the
minimum, average and maximum read coverage observed respectively. The read box represents the phage att site.

(TIFF)

S1 Table. Putative function of CDS observed in annotated phiC725A phage genome.

(DOCX)

Acknowledgments

This work has benefited from facilities and expertise of the high throughput sequencing platform of I2BC.

Author Contributions

Conceptualization: CP GV LL.
Data curation: CM.
Formal analysis: CM.
Funding acquisition: CP GV.
Investigation: CP LL YH.
Methodology: CM YH.
Project administration: CP GV.
Software: CM.
Supervision: CP.
Validation: CP GV.
Visualization: LL CM.
Writing – original draft: CP.
Writing – review & editing: CP GV LL.

References

1. Cavenagh MM, Miller RV. Specialized transduction of Pseudomonas aeruginosa PAO by bacteriophage D3. J Bacteriol. 1986; 165(2):448–52. Epub 1986/02/01. PMID: 3080405
2. Summer EJ, Gonzalez CF, Carlisle T, Mebane LM, Cass AM, Savva CG, et al. Burkholderia cenocepacia phage BcepMu and a family of Mu-like phages encoding potential pathogenesis factors. J Mol Biol. 2004; 340(1):49–65. Epub 2004/06/09. doi: 10.1016/j.jmb.2004.04.053 PMID: 15184022
3. Hulo C, Masson P, Le Mercier P, Toussaint A. A structured annotation frame for the transposable phages: a new proposed family "Saltoviridae" within the Caudovirales. Virology. 2015; 477:155–63. Epub 2014/12/17. doi: 10.1016/j.virology.2014.10.009 PMID: 25500185
4. Oliveira L, Tavares P, Alonso JC. Headful DNA packaging: bacteriophage SPP1 as a model system. Virus Res. 2013; 173(2):247–59. Epub 2013/02/20. doi: 10.1016/j.virusres.2013.01.021 PMID: 23419885
5. Huang H, Masters M. Bacteriophage P1 pac sites inserted into the chromosome greatly increase packaging and transduction of Escherichia coli genomic DNA. Virology. 2014; 468–470:274–82. Epub 2014/09/13. doi: 10.1016/j.virology.2014.07.029 PMID: 25219407
6. Chen J, Carpena N, Quiles-Puchalt N, Ram G, Novick RP, Penades JR. Intra- and inter-generic transfer of pathogenicity island-encoded virulence genes by cos phages. ISME J. 2015; 9(5):1260–3. Epub 2014/10/15. doi: 10.1038/ismej.2014.187 PMID: 25314321
7. Chen J, Ram G, Penades JR, Brown S, Novick RP. Pathogenicity island-directed transfer of unlinked chromosomal virulence genes. Mol Cell. 2015; 57(1):138–49. Epub 2014/12/17. doi: 10.1016/j.molcel.2014.11.011 PMID: 25498143

8. Lennox ES. Transduction of linked genetic characters of the host by bacteriophage P1. Virology. 1955; 1(2):190–206. Epub 1955/07/01. PMID: 13267987

9. Zinder ND, Lederberg J. Genetic exchange in Salmonella. J Bacteriol. 1952; 64(5):679–99. Epub 1952/11/01. PMID: 12999698

10. Holloway BW, Egan JB, Monk M. Lysogeny in Pseudomonas aeruginosa. Aust J Exp Biol Med Sci. 1960; 38:321–9. Epub 1960/08/01. PMID: 13715401

11. Miller RV, Pemberton JM, Clark AJ. Prophage F116: evidence for extrachromosomal location in Pseudomonas aeruginosa strain PAO. J Virol. 1977; 22(3):844–7. Epub 1977/06/01. PMID: 406425

12. Ceysens PJ, Lavigne R. Bacteriophages of Pseudomonas. Future Microbiol. 2010; 5(7):1041–55. Epub 2010/07/17. doi: 10.2217/fmb.10.66 PMID: 20632804

13. Pemberton JM. F116: a DNA bacteriophage specific for the pili of Pseudomonas aeruginosa strain PAO. Virology. 1973; 55(2):558–60. Epub 1973/10/01. PMID: 4126411

14. Hanlon GW, Denyer SP, Olliff CJ, Ibrahim LJ. Reduction in exopolysaccharide viscosity as an aid to bacteriophage penetration through Pseudomonas aeruginosa biofilms. Appl Environ Microbiol. 2001; 67(6):2746–53. doi: 10.1128/AEM.67.6.2746-2753.2001 PMID: 11375190

15. Byrne M, Kropinski AM. The genome of the Pseudomonas aeruginosa generalized transducing bacteriophage F116. Gene. 2005; 346:187–94. Epub 2005/02/18. doi: 10.1016/j.gene.2004.11.001 PMID: 15716012

16. Sanchez D, Gomila M, Bennasar A, Lalucat J, Garcia-Valdes E. Genome analysis of environmental and clinical P. aeruginosa isolates from sequence type-1146. PLoS One. 2014; 9(10):e107754. Epub 2014/10/21. doi: 10.1371/journal.pone.0107754 PMID: 25329302

17. Hardies SC, Thomas JA, Black L, Weintraub ST, Hwang CY, Cho BC. Identification of structural and morphogenesis genes of Pseuodalteromonas phage phiRIO-1 and placement within the evolutionary history of Podoviridae. Virology. 2016; 489:116–27. Epub 2016/01/10. doi: 10.1016/j.virol.2015.12.005 PMID: 26748333

18. Vu-Thien H, Corbineau G, Hormigos K, Fauroux B, Corvol H, Clement A, et al. Multiple-locus variable-number tandem-repeat analysis for longitudinal survey of sources of Pseudomonas aeruginosa infection in cystic fibrosis patients. J Clin Microbiol. 2007; 45(10):3175–83. doi: 10.1128/JCM.00070-07 PMID: 17699654

19. Sobral D, Mariani-Kurkdjian P, Bingen E, Vu-Thien H, Hormigos K, Lebeau B, et al. A new highly discriminatory multiplex capillary-based MLVA assay as a tool for the epidemiological survey of Pseudomonas aeruginosa in cystic fibrosis patients. Eur J Clin Microbiol Infect Dis. 2012; 31(9):2247–56. Epub 2012/02/14. doi: 10.1007/s10096-012-1562-5 PMID: 22327344

20. Llanes C, Pourcel C, Richardot C, Plesiat P, Fichant G, Cavallo JD, et al. Diversity of beta-lactam resistance mechanisms in cystic fibrosis isolates of Pseudomonas aeruginosa: a French multicentre study. J Antimicrob Chemother. 2013; 68(8):1763–71. Epub 2013/05/01. doi: 10.1093/jac/dkt115 PMID: 23629014

21. Essoh C, Blouin Y, Loukou G, Cablanmian A, Lathro S, Kutter E, et al. The Susceptibility of Pseudomonas aeruginosa Strains from Cystic Fibrosis Patients to Bacteriophages. PLoS One. 2013; 8(4):e60575. Epub 2013/05/03. doi: 10.1371/journal.pone.0060575 PMID: 23637754

22. Essoh C, Latino L, Mideux C, Blouin Y, Loukou G, Nguedia Gs, et al. Investigation of a Large Collection of Pseudomonas aeruginosa Bacteriophages Collected from a Single Environmental Source in Abidjan, Cote d’Ivoire. PLoS One. 2015; 10(6):e0130548. Epub 2015/06/27. doi: 10.1371/journal.pone.0130548 PMID: 26115051

23. Latino L, Essoh C, Blouin Y, Vu Thien H, Pourcel C. A novel Pseudomonas aeruginosa Bacteriophage, Ab31, a Chimera Formed from Temperate Phage PAJU2 and P. putida Lytic Phage AF: Characteristics and Mechanism of Bacterial Resistance. PLoS One. 2014; 9(4):e93777. Epub 2014/04/05. doi: 10.1371/journal.pone.0093777 PMID: 24699529

24. Latino L, Mideux C, Hauck Y, Vergnaud G, Pourcel C. Pseudolysogeny and sequential mutations build multiresistance to virulent bacteriophages in Pseudomonas aeruginosa. Microbiology. 2016; 162(5):748–63. Epub 2016/02/28. doi: 10.1099/mic.0.0000263 PMID: 26921273

25. Miller RV, Pemberton JM, Richards KE. F116, D3 and G101: temperate bacteriophages of Pseudomonas aeruginosa. Virology. 1974; 59(2):566–9. Epub 1974/06/01. PMID: 420889

26. Krylov VN, Tolmachova TO, Akhverdian VZ. DNA homology in species of bacteriophages active on Pseudomonas aeruginosa. Arch Virol. 1993; 131(1–2):141–51. Epub 1993/01/01. PMID: 8328909
27. Kazmierczak KM, Davydova EK, Mustaev AA, Rothman-Denes LB. The phage N4 virion RNA polymerase catalytic domain is related to single-subunit RNA polymerase s. EMBO J. 2002; 21(21):5815–23. Epub 2002/11/02. doi: 10.1093/emboj/cdf584 PMID: 12411499

28. Rothman-Denes LB, Schito GC. Novel transcribing activities in N4-infected Escherichia coli. Virology. 1974; 60(1):65–72. Epub 1974/07/01. PMID: 4601458

29. Falco SC, Laan KV, Rothman-Denes LB. Virion-associated RNA polymerase required for bacteriophage N4 development. Proc Natl Acad Sci USA. 1977; 74(2):520–3. Epub 1977/02/01. PMID: 322130

30. Caruso M, Shapiro JA. Interactions of Tn7 and temperate phage F116L of Pseudomonas aeruginosa. Mol Gen Genet. 1982; 188(2):292–8. Epub 1982/01/01. PMID: 6296632

31. Schwudke D, Ergin A, Michael K, Volkmar S, Appel B, Knabner D, et al. Broad-host-range Yersinia phage PY100: genome sequence, proteome analysis of virions, and DNA packaging strategy. J Bacteriol. 2008; 190(1):332–42. Epub 2007/10/30. doi: 10.1128/JB.01402-07 PMID: 17965162

32. Wu H, Sampson L, Parr R, Casjens S. The DNA site utilized by bacteriophage P22 for initiation of DNA packaging. Mol Microbiol. 2002; 45(6):1631–46. Epub 2002/10/02. PMID: 12562802

33. Spencer DH, Kas A, Smith EE, Raymond CK, Sims EH, Hastings M, et al. Whole-genome sequence variation among multiple isolates of Pseudomonas aeruginosa. J Bacteriol. 2003; 185(4):1316–25. Epub 2003/02/04. doi: 10.1128/JB.185.4.1316-1325.2003 PMID: 12562802

34. Mathee K, Narasimhan G, Valdes C, Iu X, Matewish JM, Koehrsen M, et al. Dynamics of Pseudomonas aeruginosa genome evolution. Proc Natl Acad Sci USA. 2008; 105(8):3100–5. Epub 2008/02/22. doi: 10.1073/pnas.0711982105 PMID: 18287045

35. Fernandez L, Gooderham WJ, Bains M, McPhee JB, Wiegand I, Hancock RE. Adaptive resistance to the "last hope" antibiotics polymyxin B and colistin in Pseudomonas aeruginosa is mediated by the novel two-component regulatory system ParR-ParS. Antimicrob Agents Chemother. 2010; 54(8):3372–82. Epub 2010/06/16. doi: 10.1128/AAC.00242-10 PMID: 20547815

36. Xu Z, Wang Y, Han Y, Chen J, Zhang XH. Mutation of a novel virulence-related gene mlpD in Vibrio anguillarum enhances lethality in zebra fish. Res Microbiol. 2011; 162(2):144–50. Epub 2010/11/13. doi: 10.1016/j.resmic.2010.08.003 PMID: 21070855

37. Black LW. DNA packaging in dsDNA bacteriophages. Annu Rev Microbiol. 1989; 43:267–92. Epub 1989/01/01. doi: 10.1146/annurev.mi.43.100189.001411 PMID: 2679356

38. Tavares P, Lurz R, Stiege A, Ruckert B, Trautner TA. Sequential headful packaging and fate of the cleaved DNA ends in bacteriophage SPP1. J Mol Biol. 1996; 264(5):954–67. Epub 1996/12/20. doi: 10.1006/jmib.1996.0689 PMID: 9000623

39. Youderian P, Sugiono P, Brewer KL, Higgins NP, Elliott T. Packaging specific segments of the Salmonella chromosome with locked-in Mud-P22 prophages. Genetics. 1988; 118(4):581–92. Epub 1988/04/01. PMID: 2835289

40. Sternberg N, Weisberg R. Packaging of prophage and host DNA by coliphage lambda. Nature. 1975; 256(5513):97–103. Epub 1975/07/10. PMID: 1097937

41. Moon BY, Park JY, Robinson DA, Thomas JC, Park YH, Thornton JA, et al. Mobilization of Genomic Islands of Staphylococcus aureus by Temperate Bacteriophage. PLoS One. 2016; 11(3):e0151409. Epub 2016/03/10. doi: 10.1371/journal.pone.0151409 PMID: 26953931