ORIGINAL ARTICLE

Comparative transcriptomics analyses reveal the conservation of an ancestral infectious strategy in two bacteriophage genera

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Although the evolution of tailed bacteriophages has increasingly been better understood through comparisons of their DNA sequences, the functional consequences of this evolution on phage infectious strategies have remained unresolved. In this study, we comprehensively compared the transcriptional strategies of two related myoviruses, PAK_P3 and PAK_P4, infecting the same Pseudomonas aeruginosa host strain. Outside of the conservation of their structural clusters, their highly syntenic genomes display only limited DNA similarity. Despite this apparent divergence, we found that both viruses follow a similar infection scheme, relying on a temporal regulation of their gene expression, likely involving the use of antisense transcripts, as well as a rapid degradation of 90% of the host non-ribosomal mRNA, as previously reported for PAK_P3. However, the kinetics of the mRNA degradation is remarkably faster during PAK_P4 infection. Moreover, we found that each virus has evolved specific adaptations, as exemplified by the distinct patterns of their core genes expression as well as the specific manipulation of the expression of iron-related host genes by PAK_P4. This study enhances our understanding of the evolutionary process of virulent phages, which relies on adjusting globally conserved ancestral infection mechanisms.

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Introduction

Viruses that infect bacteria, also known as bacteriophages (phages), have crucial roles in a variety of environments including the renewal of bacterial populations, recycling biochemical resources and horizontal gene transfer, as reflected by their abundance and diversity. Multiple comparative genomic analyses have suggested that related phage genomes can be clustered based on their sequence similarity (Kwan et al., 2005; Kropinski et al., 2007; Marinelli et al., 2012; Grose and Casjens, 2014). However, several studies have shown that bacteria and their phages rapidly coevolve, promoting genetic diversity in both populations (Marston et al., 2012; Williams, 2013; Koskella and Brockhurst, 2014). This genetic diversity is further stimulated by horizontal gene transfer (HGT) events, which in some cases can make it difficult to reconstruct evolutionary history of phages. Consequently, many studies have aimed at understanding how different phage clusters are related to each other or how phage populations are structured (based on comparative (meta)genomics) (Hatfull et al., 2006; Kwan et al., 2006; Grose and Casjens, 2014; Pope et al., 2015; Roux et al., 2015).

Apart from host-range and defense systems evolution, no mechanistic studies have yet experimentally addressed the evolution of phage infection strategies themselves, leaving unresolved questions on the consequences of divergent evolution of phage genomic sequences on the functions of those sequences. In particular, it has not yet been directly investigated how conserved the infection strategy is during the evolution of virulent phages, which can only be weakly predicted using bioinformatics tools. Following the isolation and genomic characterization of several phages infecting the same Pseudomonas aeruginosa host strain, an opportunistic pathogen widely present in the environment, we previously found that, although their morphologies were nearly...
identical, their genomic contents were divergent enough to support their classification within two new genera among Myoviridae, namely *Kpp10virus* and *Pakpunavirus* (Henry et al., 2015; Krupovic et al., 2016). Moreover, an additional study analyzing phages belonging to these genera pointed out that their genomes are typically composed of strongly conserved regions (0.1% single-nucleotide polymorphism (SNP) frequency) alternating with heterogeneous regions (up to 20% SNP frequency, scars of HGT and recombination events; Essoh et al., 2015). Overall, our previous comparative genomic analysis revealed that phages belonging to these distinct genera differ by their mean genome length, their number of predicted open reading frames (ORFs) and tRNA, and their GC content, but they also share homologous proteins (about 25%), most of which encoding structural components (Henry et al., 2015). Despite this significant genomic divergence, they display a strong synteny and the reconstruction of their phylogeny, based on four conserved proteins, revealed that they likely share a relatively recent common ancestor (Henry et al., 2015).

By selecting PAK_P3 and PAK_P4 as representatives of *Kpp10virus* and *Pakpunavirus* genera respectively, we aimed to investigate whether the transcriptional strategy is conserved during evolution of phages by performing a transcriptomic-based comparative analysis (RNA-sequencing). Our investigation was driven by the following questions. Do phages that have a common origin and subsequently diverge, still share common mechanisms to infect their host? How similar are the host transcriptional responses to infections by two different phages?

We found that the global infection process is conserved (temporal regulation of gene expression, production of antisense transcripts, rapid degradation of host mRNA) with few phage specific tweaks (specific manipulation of host gene expression, custom patterns of phage core gene expression). Altogether, these results suggest that, despite genomic divergence, phages are bound to their ancestral mode of action.

### Materials and methods

**Strains and growth conditions**

*P. aeruginosa* strain PAK (Takeya and Amako, 1966) was grown in LB medium supplemented with 10 mM CaCl$_2$ at 37 °C unless stated otherwise. For RNA-seq experiments, cells were infected with bacteriophage PAK_P4 (Henry et al., 2013) using a multiplicity of infection (MOI) of 25 to ensure the synchronicity of the infection (95% of the bacterial population killed after 5 min phage-introduction). Bacteriophage PAK_P3 was isolated and described by Morello et al. (2011). Updated phage genome sequences are available in GenBank and can be accessed via accession numbers NC_022970 (PAK_P3), NC_022986 (PAK_P4). Updated annotations of strain PAK genome are available via GenBank accession number CP020659.

**Adsorption assay and one-step growth experiment**

PAK_P4 adsorption assays (MOI $10^{-3}$) and one-step growth experiments (MOI $10^{-1}$) were performed as described in Chevallereau et al. (2016). Briefly, a liquid culture of strain PAK was infected at indicated MOI and incubated at 37 °C during 5 min with agitation to allow phage particles to adsorb. The culture was diluted (1/1000) and two 100 μl samples were collected every 2 min and either kept on ice until titration, or mixed with CHCl$_3$. For each sample, free phage titer (samples with CHCl$_3$) as well as the number of free phages and infective centers (samples without CHCl$_3$) were recorded to calculate eclipse and latency periods, respectively.

**Dependence to host RNA polymerase assay**

The experiment was performed according to the methods described by Ceyssens et al. (2014). The strain PAK was grown in LB medium until it reached OD$_{600nm}$ ~ 0.3, then the culture was split with one half supplemented with 100 μg ml$^{-1}$ rifampicin (Rif) and second half with the same volume of water. Ten minutes later, phages were added at MOI of 1. Samples were taken immediately (t=0), 15, 60 and 150 min post infection and phages were titrated.

**Whole transcriptome sequencing and analysis**

RNA-sequencing analysis was performed as described in Chevallereau et al. (2016). Over 12 million 75 bp reads mapping to non-ribosomal regions were obtained from each library with the exception of one early sample (2.4 million reads). After trimming, sequencing reads were aligned separately to both the phage and host genomes with the CLC Genomics workbench v7.5.1 (QIAGEN Bioinformatics, Aarhus, Denmark) These alignments were then summarized into count tables of Total Gene Reads that map to phage or host gene features respectively. Each statistical comparison presented was performed using the DESeq2 (Love et al., 2014) R/Bioconductor package to normalize host transcript populations to host transcript populations, or phage to phage, before testing for differential expression.

RNA-seq data have been deposited in NCBI-GEO with accession no. GSE86022.

RNA-seq coverage visualization is available through the COV2HTML software at https://mmonot.eu/COV2HTML/visualisation.php?str_id = -36 for a comparison of the host (0 min/13 min) and https://mmonot.eu/COV2HTML/visualisation.php?str_id = -38 for a comparison of the phage (3.5 min/13 min).

**Results**

### PAK_P3 and PAK_P4 share common life-history traits

To compare the molecular mechanisms of infection used by virulent phages that recently diverged, we chose two phages infecting the same *P. aeruginosa* host, strain PAK. These two myoviruses, PAK_P3 and
PAK_P4, respectively belong to the newly defined genera Kpp10virus and Pakpunavirus (Henry et al., 2015) and display distinct host ranges (Saussereau et al., 2014). PAK_P4 shares an average 76% (±9%) identity over 13% of PAK_P3 genome, in other words, only 9% of the nucleotide sequence is identical between the two phages (Supplementary Table 1; Figure 1). Sixty-nine homologous coding sequences, hereafter referred as core genes, have been defined (that is, ~40% of the proteomic content of the two phages, according to CoreGenes analysis with default parameters (Mahadevan et al., 2009)). The sequence identity for each pair of homologous predicted proteins ranges from 24 to 85% (with a mean of 49±14%). Most of these homologous proteins belong to the structural (49%) and nucleic acid metabolism (32%) modules, while 19% of them are encoded by early-expressed ORFs (defined in the third paragraph) which are generally considered to be the least conserved phage proteins, often responsible for the subversion of host metabolism (Calendar, 2006) (Supplementary Table 1; Figure 1). Sixty-nine homologous coding sequences, hereafter referred as core genes, have been defined (that is, ~40% of the proteomic content of the two phages, according to CoreGenes analysis with default parameters (Mahadevan et al., 2009)). The sequence identity for each pair of homologous predicted proteins ranges from 24 to 85% (with a mean of 49±14%). Most of these homologous proteins belong to the structural (49%) and nucleic acid metabolism (32%) modules, while 19% of them are encoded by early-expressed ORFs (defined in the third paragraph) which are generally considered to be the least conserved phage proteins, often responsible for the subversion of host metabolism (Calendar, 2006) (Supplementary Table 1; Figure 1). It has to be noted that among the coding sequences that do not meet the criterions required to be considered as homologous (that is, blastp score <75), some appear to have the same predicted functions. For example, the putative ribonucleoside-reductase beta subunits (PAK_P3_gp069 and PAK_P4_gp078) share only ~30% of similarity (~17% of identity) in their protein sequences. Nevertheless, most of the predicted ORFs could not be associated with a putative function.

Altogether, these observations raise the question of whether the infectious mechanisms and strategies used by PAK_P3 and PAK_P4 are conserved, given their divergent evolution. To answer this, we first compared their growth parameters. PAK_P4 adsorbs onto the strain PAK within 4.8 ± 1.7 min (k_a = 2.4 × 10^{-9} ± 1.4 × 10^{-9} ml min^{-1}) and rapidly assembles new functional virions (eclipse period: 13.3 ± 2.0 min), although they are not immediately released (latency period: 18.2 ± 2.9 min). PAK_P4 infected cells eventually produce an average of 13 ± 5 progeny phages within a mean infection cycle duration as short as 21.4 ± 1.8 min (Supplementary Figure 1). In comparison with PAK_P3, PAK_P4 displays a smaller burst size and a longer latency period (Chevallereau et al., 2016). These results hint that, if both phages have largely similar infection strategies (as shown below), they may have evolved unique processes for enacting them that may be variably effective. To investigate this further, we studied the transcriptomic dynamics of both phages.

PAK_P4 and PAK_P3 rapidly overwhelm the transcriptional environment of the host by efficiently hijacking bacterial RNA polymerase

In a previous study, we investigated the dynamics of host strain PAK transcriptome throughout PAK_P3 infection focusing on early (3.5 min) and late (13 min) stages of infection (Chevallereau et al., 2016). As PAK_P4 has a comparable eclipse period as PAK_P3, we performed a transcriptomic analysis using the same time points. While PAK_P3 induces a progressive depletion of host transcripts that eventually represent only 13% of non-ribosomal RNAs in the cell 13 min post infection, PAK_P4 is much more rapid as only 20% of the total transcripts matches the host genome as soon as 3.5 min post infection (Figure 2). Both phages shut down host cell transcription but following different kinetics. We hypothesize that they may have a different control of the machineries dedicated to (i) degrade host mRNAs and/or (ii) catalyze the transcription of viral genes.

The rapid and overwhelming replacement of host transcripts with viral transcripts could indicate that both phages rely on their own transcription apparatus even though they are not predicted to encode a viral RNA polymerase (RNAP) based on sequence homology searches. Therefore, we experimentally assessed the dependency of PAK_P3 and PAK_P4 infections on host RNAP using rifampicin (Rif), which is known to inactivate this enzyme. We found that addition of Rif abolished PAK_P3 and PAK_P4 amplification while it did not affect phage PhiKZ transcription, as previously shown (Supplementary Figure 2; Ceyssens et al., 2014). This indicates that both phages appear to
efficiently hijack the host transcription machinery to complete their infectious cycle.

The transcriptional strategy of PAK_P4 is broadly similar to PAK_P3

PAK_P4 temporally regulates its transcription over the course of infection, as previously found for PAK_P3 (Chevallereau et al., 2016). Most of the gene features encoding unique hypothetical proteins (that is, with low or no homology to any other amino acid sequence in databases) are expressed during the early stage of infection (3.5 min). This ‘early region’ encompasses gene features encoding gp080 through gp124, while the ‘late region’ encompasses genes encoding gp001 to gp034. These gene features, which are massively expressed 13 min post infection, primarily encode the predicted phage structural proteins (Supplementary Table 2; Figure 3). PAK_P4 also produces antisense RNAs (asRNAs) that appear to be acting as cis-encoded asRNAs since most of them (15 out of 17) are encoded within structural genes (Supplementary Table 2). These asRNAs are significantly more expressed (2 to 3-fold change) 3.5 min after infection compared with late infection stage (13 min). We hypothesize that the transcription of these non-coding RNAs act as repressors to attenuate the expression of the structural genes during the early stage of infection. Since both phage strategies display similar characteristics, our data support an overall conservation of regulatory mechanisms to control viral gene expression, but does not exclude specific differences (see below).

Core genes are similarly expressed in both infections but display extensive phage specific adjustment

To determine more precisely how the regulation of phage gene expression is adjusted, we compared the differential expression of the set of 69 core genes. While most conserved gene features still undergo the same temporal regulation, the strength of this differential expression appears to have changed between the two viruses (Figure 4; Supplementary Table 3). For instance, PAK_P3 tends to differentially express gp002-gp006 more drastically than PAK_P4, while PAK_P4 favors the differential expression of gp007-gp010. Strikingly, one gene is strongly upregulated during PAK_P4 late infection while it is not temporally regulated in PAK_P3. The corresponding hypothetical protein (gp014 in PAK_P4) possesses a wide range of homologs in other unrelated phages and in some bacteria. We also found that genes encoding DNA-metabolism related proteins as well as early genes are overall more strongly temporally regulated in PAK_P3 than they are in PAK_P4.

This comparative analysis of gene expression highlights that evolution has readily selected for small modifications of the mechanisms regulating a variety of specific systems, potentially having a major impact on the way the virus interacts with its host. To address this, we compare how PAK_P3 and PAK_P4 manipulate host transcription and investigate whether the host similarly responds to both phages.

PAK_P4 specifically evolved mechanisms to modulate the expression of genes involved in iron uptake

Phage PAK_P4 strongly upregulates the expression of three host operons predicted to be related to siderophore synthesis and transport, which are not upregulated during PAK_P3 infection (Figure 5; Supplementary Table 4). Indeed, PAK_4094-4104 (mean Log$_2$(FC) = 6.1 ± 1.1) and PAK_4106-09 (mean Log$_2$(FC) = 6.3 ± 0.4) are respectively homologous to the pchEFG and pchABCD operons of P. aeruginosa PAO1 (Winsor et al., 2016), which are involved in pyochelin biosynthesis, one of the major siderophores of P. aeruginosa while PAK_4090-93 (mean Log$_2$(FC) = 5.4 ± 0.5) is homologous to fpt system, encoding the corresponding siderophore receptor. Interestingly, another upregulated operon, PAK_1971-78 (mean Log$_2$(FC) = 3.0 ± 0.3) is homologous
to the pvdIJD operon (Winsor et al., 2016), encoding non-ribosomal peptide synthetases involved in the biosynthesis of the other major \textit{P. aeruginosa} siderophore pyoverdin. Moreover, \textit{PAK\_4645-51} and \textit{PAK\_4652}, respectively homologous to the \textit{phuS-TUVW} operon and \textit{phuR} receptor gene (Winsor et al., 2016), are also upregulated, although to a lesser extent (mean Log$_2$(FC) = 2.3 ± 1.0). These genes encode an ABC transporter involved in heme uptake (Ochsner et al., 2000). Finally, 14 other genes putatively involved in iron acquisition (mostly TonB-dependent receptors) are upregulated upon infection (mean Log$_2$(FC) = 2.0 ± 0.4) while \textit{PAK\_4711}, a Fur-like protein (ferric uptake regulator), which represses the expression of genes involved in iron uptake, storage and metabolism in the presence

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**Figure 3** PAK\_P4 Transcription map. Transcript densities for early and late infections, normalized by the Total Count of reads that align to the phage genome for each sample. Read density and coding sequence annotations on the Watson and Crick strands are highlighted in red and green respectively while antisense RNA, tRNA and other non-coding RNA annotations are depicted in light blue, dark blue and yellow.

**Figure 4** Comparison of PAK\_P3 and PAK\_P4 homologous genes expression. Phage homologous genes are listed and their differential expression (late versus early) during PAK\_P3 and PAK\_P4 infections are indicated as Log$_2$(fold change) values in light and dark gray, respectively.
of sufficient iron (Hantke, 2001), is downregulated (Figure 5; Supplementary Table 4).

As this upregulation of iron uptake-related genes may be the downstream consequence of a general activation of stress responses, we investigated the level of expression of alternative sigma factors which are acknowledged to orchestrate such responses. The only sigma factor that is meaningfully transcriptionally upregulated is \( \text{algU} \), which may reflect an increase of AlgU-associated bacterial stress response. We subsequently looked at the expression of \( \text{algU} \) regulon that typically includes genes involved in alginate, peptidoglycan and LPS biosynthesis, and found that none of them were upregulated upon PAK_P4 infection (Supplementary Table 4). Overall, the differential expression of iron-related genes does not appear to be part of a more general stress response.

Altogether, these results indicate that, unlike PAK_P3, which only triggers the specific and meaningful upregulation of a single operon (Chevallereau et al., 2016), PAK_P4 has specifically evolved processes leading to the complex manipulation of iron homeostasis in the host cell. We next investigate whether we could detect common host transcripts variations to both phage infections which would reflect a conserved host response against this group of viruses.

A common host response is elicited during infection by PAK_P3 and PAK_P4

We identified 55 host coding sequences upregulated in response to both PAK_P3 and PAK_P4 infection while 445 were downregulated (Supplementary Table 5). While the latter may mainly be the result of host mRNA degradation, the former could more likely represent a transcriptional general host response to phage infection, suggesting that some mechanisms for manipulating host transcription may have been conserved. Specifically, infections by both PAK_P3 and PAK_P4 prompt the marked transcriptional upregulation of a predicted P2-like prophage (Figure 6; Supplementary Table 5). However, like PAK_P3, PAK_P4 transcripts globally replace all host mRNAs faster than the increased prophage specific transcription can compete with (Chevallereau et al., 2016). Moreover, one operon linked to RNA processing (PAK_4493-4499), which was previously found to be specifically upregulated during PAK_P3 infection, is also upregulated during PAK_P4 infection, albeit more weakly (mean Log2(FC) = 6.4 vs mean Log2(FC) = 2.5, respectively; Figure 6; Supplementary Table 5). The level of upregulation of this operon between both phages also inversely correlates with the extent and speed of RNA degradation. Additionally, we found that both infections also specifically deplete Hfq transcripts, similar to PhiKZ infected PAO1 cells (Ceyssens et al., 2014; Figure 6 and Supplementary Table 5). Future experiments will address the biological consequences of these transcriptional modifications.

Discussion

While in cellular organisms, the increasing amount of genomic information is strengthening comparative
analyses delineating evolutionary relationships between cellular lineages, the lack of universal genes has hampered such analyses in viruses. In addition, the diversity of phage sequences is such that most genomes contain numerous ORFs displaying no homology to any other sequences in databanks. Therefore, reconstruction of viral evolution cannot be based solely on genomic sequences and requires further information such as elucidation of mechanistic processes (Forterre et al., 2014). In this context, we have assessed whether two phages (PAK_P3 and PAK_P4), with divergent proteomic contents and infecting the same host strain, conserved the infective strategy of their common ancestor during their evolution.

First, we observed that both phages share similar infection parameters although PAK_P4 displays a smaller burst size and a longer latent period than PAK_P3. This would not be predicted by the optimal lysis timing model which suggests that longer latency periods result in larger burst sizes (Wang et al., 1996), and requires an alternative explanation. Comparison of expressions of putative endolysin-coding genes (PAK_P4gp29 vs PAK_P3gp28), as well as the four downstream ORFs (holins and/or spanins candidates), revealed that these genes are more efficiently transcribed in PAK_P4, which also appears contradictory with the longer latency period. However, PAK_P4 encodes an asRNA (PAK_P4as15), not present in PAK_P3, matching the endolysin gene that could act as a translational repressor, preventing a rapid accumulation of endolysins and delaying lysis. With a supplementary level of lysis control, it was anticipated that PAK_P4 should be able to produce higher number of progeny than PAK_P3. However, as it is not the case, we hypothesized that both phages may rely on distinct infection mechanisms, consistent with their largely divergent protein content, that could explain the observed differences in latency period and burst size. The consequences of these adjustments in the efficiency of each phage may seem of little importance but actually appear to have much broader consequences outside the test tube. This is exemplified by the distinct ability of PAK_P3 to more efficiently treat murine pneumonia compared with PAK_P4 (Henry et al., 2013).

Therefore, we further explored phage molecular lifecycles and found that they both massively degrade host RNA with little manipulation of host gene expression, while relying on the host RNAP which is hijacked to efficiently transcribe phage gene features throughout infection. However: (i) the kinetics of RNA degradation are different and (ii) each phage co-opts the host transcription in a unique way. Remarkably, among host gene features differentially expressed in common between the two phage infections, the PAK_4493-4499 operon encoding an RtcAB system was previously proposed to be upregulated by PAK_P3 to use the RNA 3'-phosphate cyclase RtcA to prime host transcripts for adenyla- tion as part of a degradation pathway (Chevallereau et al., 2016). However, upon PAK_P4 infection, the upregulation of this operon is much weaker while the degradation of host transcripts is more extensive. Thus, the upregulation of this operon could instead be a reflection of the host attempting to respond to

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**Figure 6** Host strain specifically upregulates the expression of one operon involved in RNA processing in response to both phage infections. Host genes significantly ($P<0.05$) differentially expressed (|Log$_2$(FC)| > 1.3) between $t=0$ and $t=13$ min upon both PAK_P3 and PAK_P4 infections were listed and their Log$_2$(FC) values upon each infection are plotted.

Compared transcriptomes of two related phages

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the extensive RNA damage imposed by both phages, which is more weakly active and thus more weakly protective in PAK_P4 infected cells, which may explain the different kinetics of RNA degradation upon both infections.

As a result, we anticipated that co-option of nucleotide metabolism must be central in PAK_P4 infection process, as it has been shown for other phages, including PAK_P3 (Chevallereau et al., 2016; De Smet et al., 2016). The strong and extensive increased transcription of iron acquisition genes upon PAK_P4 infection supports this hypothesis as increased concentrations of intracellular iron would supply cofactors for viral auxiliary metabolic enzymes, eventually favoring nucleotide turnover.

Alternatively, this increased expression of iron uptake genes could potentially be explained by the recent ‘Ferrojan Horse hypothesis’, proposing that phage tail fibers are bound to iron ions which allow them to use siderophores as receptors and consequently take advantage of bacterial iron uptake mechanisms (Bonnain et al., 2016). However, no HxH motifs, known to facilitate iron binding, have been found in PAK_P4 putative tail fibers and, more generally, PAK_P4 does not possess more HxH motifs throughout its proteome than PAK_P3, which does not up-regulate iron-related genes (14 motifs versus 10, respectively).

In addition to studying direct phage-host interaction, we compared how these phages regulate their own gene expression, which appears to be broadly conserved and thus inherited from their common ancestor. Notably, recent work by Doron et al. has elegantly determined that the Syn9 phage produces a nearly identical transcriptional program when infecting three evolutionarily divergent hosts. In contrast, host response to phage infection was found to be host-specific (Doron et al., 2016). Similar results were obtained by Howard-Varona et al. using the φ38:1 phage on two Cellulophaga baltica hosts (Howard-Varona et al., 2016) indicating that neither of these two phages adjust their transcriptional program to different hosts. We have now added another piece to the puzzle by conversely studying the infection processes of two divergent phages on one common host.

We found that both phages display similar temporal regulatory schemes of their syntenic gene sets. In addition, this regulation appears to partially rely on the production of antisense transcripts, indicating that this mechanism probably already existed in the common ancestor. However, beside the genes unique to each phage (expected to specifically modulate each phage infection), we noted that differences in the transcriptional program of conserved genes (core genes) also impact infection cycles, as exemplified by the endolysin. These observations highlight the extensive panel of solutions that phages have developed to orchestrate bacterial infections. In addition, such transcriptomics studies revealing infection strategies could help implementing phage taxonomy. Indeed, including genome transcription patterns add a novel layer of properties that could be considered to cluster and differentiate phage species and genera.

In summary, even though they have almost identical morphologies and are able to infect the same host, both phages display substantial variability of their proteomic content, especially within early proteins. Thus, it was unclear whether such related phages would share common mechanisms of action. We showed that the global subversion scheme underlying infection cycle is conserved between phages deriving from a common ancestor although with distinct evolutionary trajectories. We hypothesize that this common mechanism is most likely mediated by the set of the 13 homologous phage early proteins. More intuitive is the observation that both phages have evolved unique processes to infect one particular host, probably explaining why they display distinct killing efficiency. These processes may also indicate that both phages adapted to different ecological niches (for example, low iron environment for PAK_P4).

Conflict of Interest
The authors declare no conflict of interest.

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