Lytic activity by temperate phages of *Pseudomonas aeruginosa* in long-term cystic fibrosis chronic lung infections

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*Pseudomonas aeruginosa* is the most common bacterial pathogen infecting the lungs of cystic fibrosis (CF) patients. The transmissible Liverpool epidemic strain (LES) harbours multiple inducible prophages (LESu2; LESu3; LESu4; LESu5; and LESu6), some of which are known to confer a competitive advantage in an *in vivo* rat model of chronic lung infection. We used quantitative PCR (Q-PCR) to measure the density and dynamics of all five LES phages in the sputa of 10 LES-infected CF patients over a period of 2 years. In all patients, the densities of free-LES phages were positively correlated with the densities of *P. aeruginosa*, and total free-phage densities consistently exceeded bacterial host densities 10–100-fold. Further, we observed a negative correlation between the phage-to-bacterium ratio and bacterial density, suggesting a role for lysis by temperate phages in regulation of the bacterial population densities. In 9/10 patients, LESu2 and LESu4 were the most abundant free phages, which reflects the differential *in vitro* induction properties of the phages. These data indicate that temperate phages of *P. aeruginosa* retain lytic activity after prolonged periods of chronic infection in the CF lung, and suggest that temperate phage lysis may contribute to regulation of *P. aeruginosa* density *in vivo*.

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Introduction

Cystic fibrosis (CF) patients are subject to life-long chronic respiratory infections, most commonly with the bacterium *Pseudomonas aeruginosa*. Periodic exacerbation of symptoms occurs throughout the lifetime of CF patients leading to progressive deterioration of lung function. Phage particles have been detected in the sputa of CF patients (Ojeniyi et al., 1991; Fothergill et al., 2011). Metagenomic analysis of CF sputa has identified >450 viral genotypes, whereas most viruses were unknown, of those that could be identified as viruses the majority were infective against CF pathogens, including many *Pseudomonas* phages (Lim et al., 2014). More generally, phages outnumber eukaryotic viruses both in abundance and diversity in the human virome (Reyes et al., 2012) and are known to be present at various body sites including the gastrointestinal (Breitbart et al., 2003; Kim et al., 2011) and the respiratory tracts (Willner et al., 2009). However, the *in vivo* ecological dynamics of temperate bacteriophages and their role during bacterial infections remain largely unknown.

Upon infection of a bacterial cell, a temperate phage can either complete the lytic cycle or integrate into the bacterial chromosome as a prophage, which may subsequently be induced to enter the lytic cycle by a range of bacterial or environmental cues (Little, 2005; Ghosh et al., 2009). Because lysis is obviously detrimental to the individual host bacterium it is often assumed that integrated prophages will eventually lose their lytic activity, becoming cryptic. Whereas the selective forces and mechanisms driving this remain poorly understood, inactive prophage remnants have been detected in many bacterial species, and are thought to result from an ongoing process of phage decay (Brussow et al., 2004). This could be owing to the accumulation of mutations; for example, to inactivate phage N-antiterminator genes (Desiere et al., 2001) and portal protein genes (Lawrence et al., 2001), preventing completion of
the replicative cycle. By contrast, lytic activity may be retained if it enhances bacterial population fitness, for example, by acting as an anticompetitor strategy or through the release of virulence-related toxins upon lysis (Brussow et al., 2004; Brown et al., 2006; Willner et al., 2009). Over longer evolutionary timescales, the domestication of prophages is thought to be an important process in the evolution of bacteria, leading to the origin of a number of phage-derived traits (for example, bacteriocins, killer particles etc., (Bobay et al., 2014)).

The \textit{P. aeruginosa} Liverpool epidemic strain (LES) exhibits increased antibiotic resistance levels compared with other \textit{P. aeruginosa} isolates from CF patients (Ashish et al., 2012) and is widespread across the UK (Martin et al., 2013). Patients infected with this strain have been shown to suffer greater morbidity than those infected with other strains (Al-Aloul et al., 2004; Fothergill et al., 2012). The \textit{P. aeruginosa} LESB58 genome contains five inducible prophages and transposon mutagenesis of this isolate identified several mutations in prophages encoding LES\textsubscript{p}2, LES\textsubscript{p}3 and LES\textsubscript{p}5 that reduced bacterial competitiveness in a rat model of chronic lung infection (Winstanley et al., 2009), suggesting that the phages have a key role in the infection process. We have previously characterised the infection properties of several LES phages in vitro. Induction experiments demonstrated that free LES\textsubscript{p}2 was produced more rapidly and in higher numbers than LES\textsubscript{p}3 and LES\textsubscript{p}4 in response to norfloxacin. Each phage was shown to exhibit a different immunity profile and was able to infect a range of susceptible \textit{P. aeruginosa} hosts (James et al., 2012). Owing to a lack of suitable acceptor strains, we have thus far been unable to isolate and purify LES\textsubscript{p}5 and LES\textsubscript{p}6.

In this study, we used culture-independent quantitative PCR (Q-PCR) to follow the ecological dynamics of all 5 active LES phage populations in 188 expectorated sputum samples from 10 long-term LES-infected patients over a period of 28 months. To our knowledge, this represents the first longitudinal study of a bacterial pathogen and its temperate phages in a human chronic infection.

### Materials and methods

#### Patients and Samples

Sputum samples (\(n = 188\)) were collected from 10 LES-infected CF patients for diagnostic purposes over a period of >2 years (January 2009 to May 2011). The details of each patient and the sampling rationale have been described previously (Fothergill et al., 2010; Mowat et al., 2011). All patients had long-term LES infections (duration at beginning of the study ranged from >5 to >10 years). Table 1 and Supplementary Figure S1 describe the number and dates of acquired samples from each patient that were analysed for density of LES bacteria and LES phages. During routine visits 98 samples were collected when each patient was well (stable) and 90 samples were collected during periods of acute exacerbated symptoms of respiratory infection (acute). Sputa obtained during exacerbation periods included samples taken before and during aggressive intravenous antibiotic treatment (Table 1, Supplementary Figure S1). The criteria for diagnosing exacerbations were physician-based and have been described previously (Mowat et al., 2011). Briefly, patients were considered to be undergoing an exacerbation if they showed signs of reduced lung capacity, increased sputum production and discoloration, increased temperature, cough, dyspnoea and malaise (Goss and Burns, 2007). Where known, antibiotics administered to patients during the study period are listed in Table 1. However, detailed information of the antibiotics administered during each exacerbation is very incomplete.

| Patient | \(\phi\) Complement | Total | Stable | Exacerbation | Antibiotics |
|---------|---------------------|-------|--------|--------------|-------------|
| CF1     | 3, 4, 6             | 17    | 14     | 3            | TOB (N), AZT (O) |
| CF2     | 2, 3, 4, 6          | 11    | 9      | 2            | CEPH (O), CEF (IV), COL (IV) |
| CF3*    | 2, 3, 4, 6          | 7     | 1      | 6            | AZT (O), COL (N), CEF (IV), COL (IV) |
| CF4     | 2, 3, 4, 6          | 14    | 10     | 4            | CEF (IV), COL (IV) |
| CF5     | 2, 3, 4, 6          | 10    | 7      | 3            | AZT (O), COL (N) |
| CF6     | 2, 3, 4, 6          | 16    | 9      | 7            | AZT (O), COL (N), CEF (IV), COL (IV), MER (IV), FOS (IV) |
| CF7*    | 2, 3, 4, 5, 6       | 28    | 11     | 17           | CEPH (O), FOS (IV), MER (IV), CEF (IV) |
| CF8     | 2, 3, 4, 5, 6       | 33    | 8      | 25           | AZT (O), COL (N), TOB (N), MER (IV), COL (IV) |
| CF9     | 2, 3, 4, 5, 6       | 25    | 19     | 6            | MER (IV), COL (IV) |
| CF10    | 2, 3, 4, 5, 6       | 27    | 10     | 17           | MER (IV), COL (IV) |
| Total   |                     | 188   | 98     | 90           |             |

**Abbreviations:** AZT, azithromycin; CF, cystic fibrosis; CEPH, cephadrine; CEF, cefadiazime; COL, colomycin; FOS, fosfomycin; IV, intravenous (used during exacerbations only); LES, Liverpool epidemic strain; MER, meropenem; N, nebulised; O, oral; TOB, tobramycin.

CF patient sputum contained LES variants that harboured different phage complements. Stable samples were collected during periods of relative patient health. Exacerbation samples were collected during periods of reduced lung function and hospitalisation of patients, who underwent antibiotic treatment for which data are incomplete. Antibiotics used during the period of the study are shown (where known): Route of administration is indicated in brackets.

*Patients CF3 and CF7 died before completion of this study.*
Thus we were unable to fully assess the effect of different antibiotics on phage induction in vivo. This study was approved by the local Research Ethics Committee (REC reference 08/H1006/47).

Detection of viable phage particles by plaque assay
We were unable to accurately determine phage densities by culture-dependent techniques for several reasons: (a) we lacked acceptor strains for LESp5 and LESp6; (b) culturable phages were indistinguishable by plaque morphology; and (c) sputum samples were routinely frozen upon collection, which reduced phage viability. However, to confirm the presence of viable phage particles in sputum samples, we quantified the density of culturable phages in 10 sputum samples from 3 LES-infected patients (CF3, CF4 and CF7). Sputum samples (50 µl) were treated with sputasol (200 µl) and incubated at 37 °C for 1 h, with shaking at 200 r.p.m. Treated samples were diluted with sterile phosphate-buffered saline. A rifampicin-resistant mutant (PAO1-rif) was created by successive passage in increasing rifampicin concentrations (method described by James et al., 2001). This enabled enumeration of active phages capable of infecting PAO1 directly from unfiltered sputum. Briefly, mid-exponential phase PAO1-rif (OD600 0.5; 100 µl) was added as an indicator host to treated sputum samples (400 µl). Rifampin (300 mg ml⁻¹) was incorporated in the soft agar overlay (5 ml; 0.4% (w/v) Luria broth (LB) agar) to select for the indicator host and incubated overnight at 37 °C before the plaques were counted. This method only provides confirmation that active P. aeruginosa phages are present in the sputa. It does not accurately reflect abundance and does not discriminate between phages.

Real-time Q-PCR
To overcome the limitations of culture-dependent methods, we have developed and validated a simple Q-PCR protocol to measure the density of each individual LES phage (James et al., 2012). Each sputum sample was treated with an equal volume of Sputasol (Oxoid, Basingstoke, UK) and incubated at 37 °C for 30 min, with shaking at 200 r.p.m. DNA was prepared from each treated sputum sample (400 µl) using the ‘Qiasymphony Virus/Pathogen DNA extraction kit’ (Qiagen, Valencia, CA, USA) and the automated QIASymphony machine (QIAGEN; pathogen complex 200 protocol). The protocol yielded 0.3–0.9 µg µl⁻¹ DNA. Each sample was diluted 1:100 with sterile distilled H₂O. The number of DNA copies of each LES phage in sputum and bacterial culture samples was quantified from extracted DNA. The number of specific copies detected for each phage was compared with a concentration gradient of known standards (James et al., 2012). For each LES phage (LESp2–LESp6), two specific primer sets were used to quantify i) prophage and ii) total copies (10 primer sets in total). Differentiation between total phage and prophage copies, allowed free-phage densities to be calculated as the difference between these values as previously described (James et al., 2012). Bacterial host density was quantified using primers specific for P. aeruginosa (PS21-6F1/PS21-6R1 and gyrPA-F1/gyrPA-R1) (Fothergill et al., 2013). All primer sequences and targets are listed in Supplementary Table S1.

Q-PCR reactions (25 µl) contained 1 µM each primer pair and 1X Rotorgene-SYBR green supermix (Qiagen). All primer sets were used with the same cycling conditions: 95 °C for 10 min; followed by 40 cycles of 95 °C for 10 s, 60 °C for 15 s and 72 °C for 20 s. Phage DNA copy numbers were quantified from DNA samples (1 µl) in triplicate using a Rotorgene cycler (Qiagen). Q-PCR data were analysed using Rotorgene Q series software 1.7 (Qiagen).

Sputasol induction experiments
To test for any potential induction properties of sputasol, DNA was extracted from LESB58 cultures grown to mid-exponential phase in LB (James et al., 2012) treated (in triplicate) with an equal volume of sputasol or LB for 30 min. DNA was prepared from each culture using a DNA mini kit (Qiagen) and phage densities estimated by Q-PCR as described for the sputum samples. No effect of sputasol on phage induction was observed (Supplementary Figure S2).

Statistical analysis
To model phage densities (or phage-to-bacterium ratios), we fitted linear mixed effects models with maximum likelihood using the R package nlme (Pinheiro and Bates 2000) with and without temporal autocorrelated errors (an ARMA(1) model). Models with temporal autocorrelated errors were significant improvements over those without, and therefore we present only these models below. We included a random effect for patient ID and fixed effects for time, exacerbation, bacterial load and the interaction between bacterial load and exacerbation. We compared full models with and without temporally autocorrelated errors using a likelihood ratio test, and then used a backwards stepwise process to remove nonsignificant fixed effects until the minimum adequate model was identified. Models analysing normalised variables gave similar results to those analysing non-normalised data and are presented in the supplementary information (Supplementary Table S2a).

Results
Dynamics of total free-phage abundance
Plaque assays confirmed the presence of viable particles of the subset of culturable phages...
in all sputum samples that were tested (range \(1.25 \times 10^2\)–\(1.39 \times 10^4\) PFU ml\(^{-1}\)). Free-LES phage DNA was detected in all patient sputa (9.35 \(\times\) \(10^4\)–5.54 \(\times\) \(10^9\) copies ml\(^{-1}\)) and, in general, total free-phage density (that is, the sum of all the free-LES phages present in the sample) exceeded that of \(P.\) aeruginosa within each sample (mean range 11-fold to 90-fold) (Figure 1 and Supplementary Table S3). We observed a positive linear relationship between total free-phage density and bacterial density (Figure 2a and Supplementary Table S2b; bacterial coefficient \(0.607 \pm 0.054\), \(LRT = 85.9\), d.f. = 1, \(P < 0.001\)), consistent with ongoing lytic activity by the temperate phages, but no effect of time or exacerbations on total free-phage densities. Further, we observed a negative linear relationship between the phage-to-bacterium ratio and bacterial density (Figure 2b and Supplementary Table S2c bacterial coefficient \(-3.206 \pm 0.484\), \(LRT = 108.4\), d.f. = 1, \(P < 0.001\)), suggesting a role for phage lysis in regulation of bacterial densities. Time and exacerbations had no significant effect on phage-to-bacterium ratios. It is perhaps surprising that exacerbations were not associated with either a change in phage densities or a change in the phage-to-bacterium ratio (Figure 3), given that these episodes are associated with the administration of high-dose intravenous antibiotics. However, it should be noted that these patients were all subject to variable cocktails of antibiotics over several years irrespective of exacerbations (Table 1). Moreover, clinical data on antibiotic use in these patients was too incomplete to be used in analyses, and therefore effects of particular antibiotics on phage dynamics may have been missed.

Abundance hierarchy among individual phages within lungs

Next, we considered each phage individually, observing a general hierarchy of free-phage densities, though the precise patterns were clearly influenced by the fact that the LES populations for each patient did not all share the same prophage complement (Table 1). Figure 4 illustrates the free-phage densities of individual LES phages for each of the patients. In the majority of patients (CF2–CF9) similar free-phage dynamics were observed, in that the density of free LES\(_{\varphi}2\) was consistently higher than that of the other LES phages, closely followed by LES\(_{\varphi}4\) (Figure 4). A positive correlation was observed between LES\(_{\varphi}2\) and LES\(_{\varphi}4\) densities in these patients (Supplementary Table S4). The dynamics observed in samples from patient CF10 (Figure 3) exhibited a change in the hierarchy of free phage, with considerably higher free-LES\(_{\varphi}4\) densities observed. Despite consistent carriage of LES prophage 3, very little free LES\(_{\varphi}3\) was detected in patients CF2–CF10. However, higher levels of free LES\(_{\varphi}3\) (\(3.29 \times 10^7\) copies ml\(^{-1}\)) were observed in all sputa from patient CF1 (Figure 4), whose \(P.\) aeruginosa were the only populations not to carry prophage 2 (Table 1). We showed previously that LES populations exhibit genotypic diversity,

![Figure 1](image-url)
including variation in carriage of LES phages. In particular, the carriage of LES prophage 5 was not consistent in all individuals of a given LES population (Fothergill et al., 2010). In this study, prophage 5 was intermittently detectable in the sputum from patients 7 (up to $10^5$ copies µL$^{-1}$) and 10 ($10^2$–$10^4$ copies µL$^{-1}$). This explains the low density of free LESφ5 in these patients. Free copies of LESφ6 were not detected in the majority of sputum samples. Where free copies were detected, the density was lower than the host bacterial load ($6.7 \times 10^5$–$1 \times 10^7$ copies µL$^{-1}$).

**Discussion**

The levels of free-LES phages detected in all patients throughout this study suggest an active lytic cycle that may be promoted by the presence of H$_2$O$_2$, or DNA-damaging antibiotics in the CF lung (McGrath et al., 1999; Fothergill et al., 2011). Surprisingly, we observed no effect of patient exacerbation on total free-phage density, although this is consistent with previous studies showing that neither fluctuations in *P. aeruginosa* populations (Mowat et al., 2011), nor in the wider bacterial population (Fodor et al., 2012), show any relationship with the exacerbation period in chronically infected patients, despite the use of high-level intravenous antibiotic therapy. It is known that particular antibiotics can induce phage lysis and it is possible that different antibiotics regimes may have influenced differential induction of phages between patients. Indeed, we have shown previously that LES induction varies in response to different antibiotics (Fothergill et al., 2011). Unfortunately, because records of antibiotic treatments for these patients were very incomplete, we were unable to explicitly test for effects of particular antibiotics in this study. This would in any case be difficult because of the extensive and varied use of antibiotics in this group of patients (Table 1), which was not restricted to periods of exacerbation.

Our data do however suggest that ongoing phage lysis may have a role in regulating bacterial density in the CF lung. Treatments that induced the lytic cycle of temperate phages could therefore offer a promising alternative or addition to standard antibiotic therapies, which in themselves often do not successfully reduce *P. aeruginosa* densities in long-term chronically infected patients (Foweraker, 2009; Mowat et al., 2011). Several studies have demonstrated effective phage-antibiotic synergism in the reduction of bacterial numbers *in vitro* and *in vivo* (Hagens et al., 2006; Comeau et al., 2007; Knezevic et al., 2013). However, this strategy would need to be considered with caution. Antibiotic therapies that induce stx phages of Shiga-toxigenic *E. coli* have been shown to increase expression of Shiga toxin genes that are encoded in the late region of the phage genome and thus increase cytotoxic damage and exacerbate symptoms (Matsushiro et al., 1999). Although we have not identified any obvious virulence factors encoded in the late gene region of the LES phages (Winstanley et al., 2009; James et al., 2012), we cannot ignore the possibility that the lytic cycle might induce upregulation of virulence genes.

We demonstrate here that LESφ2 was the most abundant free phage in 9-out-of-10 LES-infected patients. The hierarchy of free-LES phage in patient sputa was also observed in our previous studies of LES phage induction in *in vitro* bacterial cultures (James et al., 2012). This suggests therefore that LESφ2 is generally more readily induced or exhibits a more efficient lytic cycle than the other phages both *in vitro* and *in vivo*. In the sputa of patient CF1, who was infected by a LES that lacked prophage 2, LESφ3 reached far higher abundances than observed in other patients, suggesting potential suppression of LESφ3 lysis by LESφ2 *in vivo*. In accordance with our previous *in vitro* observations of coinduction of lysis by phages, we observed a degree of synchronisation of free-phage dynamics *in vivo*. 
suggesting that the phages may be responding to shared signals, which could include a wide variety of human host, bacterial and environmental triggers (Little, 2005). It is exceptionally difficult to disentangle to drivers of microbial dynamics in vivo due to the complexity of host microenvironments; future
studies using laboratory models of the infection environment allowing the constituent drivers to be decomposed will be necessary to elucidate this (Wright et al., 2013; Fothergill et al., 2014).

The long-term maintenance of intact, active temperate phages in the LES genome despite substantial cell lysis suggests some selective or competitive advantage in vivo, consistent with previous work highlighting a loss of competitiveness observed following the introduction of mutations to some LES prophage regions (Winstanley et al., 2009). One possibility is that free-phage particles produced by a subpopulation of LES could kill competing bacteria (Brown et al., 2006). Indeed, LESp2, LESp3 and LESp4 are capable of infecting and lysing other clinical P. aeruginosa isolates (James et al., 2012). Thus frequent induction of the lytic cycle may enhance the competitive ability of LES by promoting superinfection, which has been observed clinically (McCallum et al., 2001), and preventing invasion of the lung by other strains of P. aeruginosa. Alternatively the prophages may contain accessory genes that contribute directly to LES fitness in the CF lung, which are only expressed during the lytic cycle, as observed for other pathogens (Wagner et al., 2001). Little is known about the consequences for the human host of the presence of large numbers of phage in the lung. However, high titre phage preparations have recently been found to interact with the immune system in vivo (Lektiewicz et al., 2010). It has also been suggested that, following adherence to mucus, some phages may act as a form of innate host immunity enhancing host defences against bacterial pathogens (Barr et al., 2013). Our findings of high free-phage abundances in CF lungs highlight the urgent need for research into the interaction of phages with host immunity, particularly in CF where dysfunctional immune responses contribute to pathological processes.

Conflict of Interest
The authors declare no conflict of interest.

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