The effect of metal remediation on the virulence and antimicrobial resistance of the opportunistic pathogen *Pseudomonas aeruginosa*

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Funding information
Natural Environment Research Council, Grant/Award Number: NE/N019717/1, NE/R011524/1, NE/T008083/1, NE/V012347/1 and NE/W006820/1; UK Research and Innovation, Grant/Award Number: MR/V022482/1

Abstract

Anthropogenic metal pollution can result in co-selection for antibiotic resistance and potentially select for increased virulence in bacterial pathogens. Metal-polluted environments can select for the increased production of siderophore molecules to detoxify non-ferrous metals. However, these same molecules also aid the uptake of ferric iron, a limiting factor for within-host pathogen growth, and are consequently a virulence factor. Anthropogenic methods to remediate environmental metal contamination commonly involve amendment with lime-containing materials. However, whether this reduces in situ co-selection for antibiotic resistance and siderophore-mediated virulence remains unknown. Here, using microcosms containing non-sterile metal-contaminated river water and sediment, we test whether liming reduces co-selection for these pathogenicity traits in the opportunistic pathogen *Pseudomonas aeruginosa*. To account for the effect of environmental structure, which is known to impact siderophore production, microcosms were incubated under either static or shaking conditions. Evolved *P. aeruginosa* populations had greater fitness in the presence of toxic concentrations of copper than the ancestral strain and showed increased resistance to the clinically relevant antibiotics apramycin, cefotaxime and trimethoprim, regardless of lime addition or environmental structure. Although we found virulence to be significantly associated with siderophore production, neither virulence nor siderophore production significantly differed between the four treatments. Furthermore, liming did not mitigate metal-imposed selection for antibiotic resistance or virulence in *P. aeruginosa*. Consequently, metal-contaminated environments may select for antibiotic resistance and virulence traits even when treated with lime.

Keywords
antibiotic resistance, liming, metal pollution, opportunistic pathogen, *Pseudomonas aeruginosa*, siderophores
1 | INTRODUCTION

Metals are ubiquitous in the Earth's crust and many are essential for cellular processes (Festa & Thiele, 2011; Palmer & Skaar, 2016; Sparks, 2005). However, agriculture and industry have resulted in toxic levels of metal contamination in many environments via pesticide use, sewage sludge application, atmospheric deposition and mining (Kibria et al., 2016; Sparks, 2005). Human exposure to metals via contaminated crops or water poses a serious health threat (Cui et al., 2005; Paul, 2017; Tóth et al., 2016; Wu et al., 2018; Yunus et al., 2016). Metal contamination can decrease microbial diversity, biomass and functionality, which in turn can affect ecosystem function (Giller et al., 1998; Nwuche & Ugoji, 2008; Wang et al., 2007) and potentially select for resistance and virulence in environmental pathogens (Ferrarese et al., 2020). Such are the potential health and economic costs that metal remediation is common practice (Palansooriya et al., 2020). Often, this involves adding lime-containing materials to acidic metal-contaminated environments (Derome & Saarsalmi, 1999; Haimi & Mätäsnemi, 2002; Rogora et al., 2016). By raising the pH, liming causes metal ions to precipitate and become less soluble and consequently less bioavailable (Chen et al., 2009; Giller et al., 1998; Hesse et al., 2019; Palansooriya et al., 2020; Ruttens et al., 2010). This decreases metal uptake by plants, which helps alleviate phytoxicity and prevent metals from entering the food chain (Bolan et al., 2003). Lime addition is the oldest and most widely used metal remediation method (Ruttens et al., 2010) and has been used to treat contamination in lakes (Gunn et al., 2016; Rogora et al., 2016), rivers (Egeberg & Håkedal, 1998) and soils (Derome & Saarsalmi, 1999; Haimi & Mätäsnemi, 2002; Hesse et al., 2019).

Reducing metal bioavailability by liming will inevitably affect the microbial community (Hesse et al., 2019; Kelly et al., 2003; Ramos et al., 1987). However, how such remediation could affect the ability of environmental pathogens to cause infection or to withstand antibiotic treatment remains largely unexplored. A key microbial trait likely to change after liming is the production of siderophore compounds (Hesse et al., 2019). Siderophores are extra-cellular compounds that can chelate toxic metal ions such as copper, zinc and nickel (Braud et al., 2010a). These toxic metal-siderophore complexes cannot re-enter the cell due to the selectivity of the outer-membrane transport proteins (Braud et al., 2010a; Rajkumar et al., 2010), and consequently, siderophore production can be selected for as a detoxifying method in the presence of bioavailable toxic metals (Braud et al., 2010a, 2010b; Hesse et al., 2018). However, alongside their role in detoxification, siderophores are also selected for to aid iron (Fe) sequestration from the extra-cellular environment (Hesse et al., 2018; O’Brien et al., 2014). Fe is vital for microbial growth as a cofactor for a number of essential enzymes (Cuív et al., 2006; Rajkumar et al., 2010), but is most commonly present as insoluble Fe$^{3+}$ and therefore is of limited bioavailability, especially at near-neutral pH (Braud et al., 2010a; Cassat & Skaar, 2013; Cuív et al., 2006; Kümmerli, Jiríny, et al., 2009; Rajkumar et al., 2010; Winkelmann, 2007). Siderophores are released by cells into the environment where they form extra-cellular complexes with Fe$^{3+}$ at a much greater association constant than for non-ferrous metals (Braud, Hoegy, et al., 2009). The siderophore-iron complexes can then be taken up by selective outer-membrane transport proteins before Fe$^{3+}$ is reduced to bioavailable Fe$^{2+}$ and the siderophore made available for reuse (Hider & Kong, 2010). Siderophores are important virulence factors as they allow pathogens to grow within hosts that actively withhold iron (Harrison et al., 2006; Skaar, 2010). Consequently, toxic metal concentrations can select for greater virulence by selecting for increased siderophore production (Lear et al., 2022). Lime remediation of metal-contaminated environments thus could potentially select either for the upregulation of siderophore production when it predominantly results in decreased bioavailability of Fe, or for the downregulation of siderophore production when it predominantly results in lower metal toxicity, with concomitant expected changes in virulence. Previous work has shown siderophore production to decrease as a consequence of liming at the level of whole microbial communities (Hesse et al., 2019), but whether this also occurs in environmental pathogens that rely on siderophore-mediated iron uptake remains untested.

It is well established that some mechanisms that bacteria use to resist metal contamination also confer resistance to antibiotics (Baker-Austin et al., 2006). This can occur through cross-resistance when a single mechanism provides resistance to both types of stressors (e.g. efflux pumps (Baker-Austin et al., 2006; Hamzehpour et al., 1995; Köhler et al., 1996; Perron et al., 2004; Sarma et al., 2010; Seiler & Berendonk, 2012; Teitzel & Parsek, 2003)), through co-resistance when metal and antibiotic resistance genes are located on the same genetic element (Ghosh et al., 2000; Gullberg et al., 2014), or through co-regulation when transcriptional and translational responses to both stressors are linked (Allen et al., 1977; Baker-Austin et al., 2006; Dickinson et al., 2019; Perron et al., 2004; Stepanauskas et al., 2006). However, to our knowledge, it remains untested whether metal remediation could decrease such co-selection for antibiotic resistance.

In this study, we use the opportunistic pathogen *Pseudomonas aeruginosa* to test whether liming alters virulence by influencing siderophore production and whether it decreases co-selection by metals for antibiotic resistance. We applied an experimental evolution approach, utilizing microcosms containing water and sediment and the resident microbial community from a river heavily contaminated with historical mine waste (Vos et al., 2020; Pirrie et al., 2003). We embedded *P. aeruginosa* within this natural microbial community and quantified antibiotic resistance, siderophore production and virulence in this focal species after 14 days. *P. aeruginosa* is responsible for a significant proportion of nosocomial infections, particularly those in intensive care units and immunocompromised patients (Rice, 2008). This species is of significant clinical importance as it is resistant to many treatments, both intrinsically and due to its ability to rapidly evolve resistance (Pachori et al., 2019). Outside of the clinical setting, *P. aeruginosa* is commonly found in soil and water (Rutherford et al., 2018), especially in areas closely linked with human activity and pollution (Crone et al., 2020), where it can pose a significant health risk (Mena & Gerba, 2009). Moreover, *P. aeruginosa* has...
been found to occur in 43% of samples taken from environments significantly impacted by human activity and in 19% of samples from more pristine environments (Crone et al., 2020). The production of siderophores by P. aeruginosa is well-studied as a virulence factor, metal resistance mechanism and public good (Braud et al., 2010a; Cuiv et al., 2006; Granato et al., 2016; Kümmerli, Jiricny, et al., 2009; O’Brien et al., 2014, 2018). Furthermore, the growing interest in its use along with other siderophore-producing species to assist phytoremediation of metals using plants (Braud, Jézéquel, et al., 2009; Rajkumar et al., 2010), makes it an ideal focal species for this study.

To quantify any changes in the virulence of P. aeruginosa, we used the insect infection model Galleria mellonella (Greater Wax Moth larvae), a low-cost and ethically expedient alternative for mammalian virulence screens (Guerrieri et al., 2019; Hernandez et al., 2019). We quantified total siderophore production using a Chrome Azurol S (CAS) assay (Schwyn & Neilands, 1987) and pyoverdine production—the main siderophore produced by P. aeruginosa (Dumas et al., 2013)—by measuring fluorescence; and tested whether these are correlated with virulence. Extra-cellular siderophore-metal complexes offer a fitness advantage not only to the producer but also to neighbouring cells, whether these are fellow producers or not (Buckling et al., 2007; Kümmerli, Griffin, et al., 2009; Ross-Gillespie et al., 2007). Non-siderophore-producing ‘cheats’ could gain a selective advantage as they benefit from siderophore production but do not carry the cost of production (Cordero et al., 2012; Kümmerli, Jiricny, et al., 2009; O’Brien et al., 2014; Winkelmann, 2007). Cheat fitness is increased in spatially unstructured environments because the greater mixing increases the opportunity to take up siderophore-iron complexes and benefit from siderophores detoxifying the area (Ross-Gillespie et al., 2007). To take into account any possible effect that spatial structure may have on siderophore production, and consequently virulence, we performed our experiments in both static and shaken microcosms. We tested whether the addition of lime or a change in spatial structure affects P. aeruginosa resistance to the antibiotics apramycin, cefotaxime and trimethoprim. Both apramycin and cefotaxime have been declared ‘critically important’ for human medicine and trimethoprim ‘highly important’ by the WHO (World Health Organization, 2019). Moreover, apramycin has been shown to be effective against highly drug-resistant strains of P. aeruginosa (Kang et al., 2017), including those isolated from cystic fibrosis patients (Di Bonaventura et al., 2022) and cefotaxime is used in aquaculture to treat P. aeruginosa infections (Ali et al., 2021).

2 | METHODS

2.1 | Collection of river samples and microcosm set-up

Water and sediment samples were collected from the metal-contaminated Carnon River in Cornwall, UK (50°13′54.6″N, 5°07′48.7″W; Vos et al., 2020). We chose this site as it is polluted by a host of toxic metals, including copper, manganese and zinc (Vos et al., 2020; Hesse et al., 2019), and liming has been shown to reduce total non-ferrous metal availability in samples collected in the near vicinity (Hesse et al., 2019). The water sample in reference (Vos et al., 2020) was taken at the same time as the samples used in this study, and contains high concentrations of non-ferrous metals. Sediment was collected using a sterile spatula and water was collected by filling a sterile 1000-mL duran bottle (Schott Duran). Sediment (3±0.1 g) and river water (6 mL) were added to each microcosm (25 mL, Kartell). The combined water and sediment pH was measured using a Jenway 3510 pH metre (Jenway).

2.2 | Experimental design

Two treatments—liming (lime amendment/no amendment) and spatial structure (shaken/unshaken)—were carried out in a full factorial design (Figure 1); six replicates were used per unique treatment combination resulting in a total of 24 microcosms. All microcosms were incubated at 20°C. To raise the pH from 5.8 to ~7.0 to represent a metal remediation scenario, 30 mg (±1.0 mg) of undissolved hydrated lime (Verve Garden lime (Hesse et al., 2019)) was added to each relevant microcosm, then left for 14 days to equilibrate (a pilot study showed this is how long it took the pH to become stable). To observe differences between structured and non-structured environments, microcosms were either kept static or were continuously shaken at 210rpm (Stuart orbital incubator S1600). Shaking began on day 14 when the pH was stable and ended on day 28 (Figure 1). Although shaking was primarily undertaken to influence the structure of the environment, it will also increase the oxygen content and may increase the dissolution rate of lime—and consequently the alkalinity—of the shaken flasks compared with the static flasks. To account for these effects, iron speciation and redox potential were monitored as indicators of sediment oxygenation (see below) and the pH was also monitored throughout. Finally, alongside the 24 microcosms inoculated with P. aeruginosa, 24 microcosms were set up without P. aeruginosa added to test for the virulence of the resident microbial community.

On day 14, 30 μL (7.3×10⁵ colony forming units: cfu) of Pseudomonas aeruginosa (PAO1lacZ: O’Brien et al., 2017) was added to each microcosm. It was added on day 14 when the pH was stable, so that the populations in the shaking treatment were never static. This lab strain is both lacZ marked and gentamicin resistant allowing it to be easily distinguished from the rest of the community on agar containing X-gal (5-bromo-4-chloro-3-indolyl-β-D-galactopyranoside; 100 μg/L; VWR Chemicals) and gentamicin (30 μg/mL Sigma). P. aeruginosa was grown overnight in shaking microcosms containing 6 mL of King’s medium B (KB; 10 g glycerol, 20 g proteose peptone no. 3, 1.5 g K₂HPO₄, 1.5 g MgSO₄ per litre). To stop residual nutrients being added with P. aeruginosa to the microcosms, 2 mL of culture was centrifuged at 1233 g for 30 min, after which supernatant was decanted, and the pellet resuspended in 1 mL of M9 salt buffer (221.1 mM KH₂PO₄, 42.3 mM Na₂HPO₄, 85.6 mM NaCl) followed by plating on KB agar to calculate the inoculation density. On day 28, all microcosms were destructively sampled by adding sterile glass beads and 12 mL of M9 buffer and vortexing for 1 min. Samples were then aliquoted and stored in glycerol (25% final volume) at −80°C.
Iron analysis (ferrozine assay)

To determine whether liming affected Fe speciation and therefore bioavailability, a ferrozine assay was used to measure relative concentrations of Fe$^{2+}$ and total bioavailable iron (Lovley & Phillips, 1986, 1987). The first step of this assay quantifies Fe$^{2+}$, which is easily obtainable by bacteria and so does not require siderophores. The second step quantifies both Fe$^{2+}$ and Fe$^{3+}$ and therefore gives a measure of total bioavailable iron including the fraction requiring scavenging mechanisms, such as siderophores. By dividing the first measurement by the second it is possible to estimate the proportion of iron in each treatment that is of relatively high bioavailability to *P. aeruginosa* (Lovley & Phillips, 1986, 1987). The first measurement is given by digesting 100 μL of fresh river water and sediment sample (*n* = 3) in 4.9 mL of 0.5 M hydrochloric acid for 1 h before 50 μL was mixed with 2.45 mL of ferrozine solution (1 g ferrozine, 11.96 g (4- (2-hydroxyethyl)-1-piperazineethanesulfonic acid/L); adjusted to pH 7) in a cuvette (*n* = 3 per replicate). This was left to stand for exactly 1 min before absorbance at 562 nm was measured using a spectrophotometer (Jenway 7315). To quantify total bioavailable Fe (step 2), 200 μL of 6.25 M hydroxylamine hydrochloride was added to the digested samples and left to stand for another hour. This was then added to ferrozine solution in cuvettes and measured as before. Standards of known concentrations of FeSO$_4$·7H$_2$O were measured to allow conversion of absorbance to Fe concentrations.

Copper growth assay

To test that metal concentrations in our river water and sediment samples were sufficiently high to select for metal resistance, and to test whether liming impacted this selection, we used a copper growth assay. Specifically, we added 20 μL of either the ancestral *P. aeruginosa* strain or defrosted samples of the evolved populations to a 96-well plate well containing 180 μL of plain Iso-Sensitest broth (Oxoid) and 20 μL to a well containing 180 μL of Iso-Sensitest broth at a concentration of 1 g/L of copper sulphate (CuSO$_4$; Alfa Aesar). The optical density OD$_{600}$ was then read every 10 min for 18 h using a Biokit Synergy 2 spectrophotometer, with a 5-s shake at 180 rpm before each read. We used 1 g/L of copper sulphate as this equates to a copper concentration (6.26 mM) previously found in highly polluted environments (Brun et al., 1998; UKWIR, 2018).

Siderophore (CAS) assay

Total siderophore production was quantified using the CAS assay (Schwyn & Neilands, 1987). Samples were plated onto tryptic soy agar (TSA: Oxoid) supplemented with nystatin (Sigma: 20 μg/mL) to suppress fungal growth and X-gal to allow visual identification of our focal species. After 48 h, *P. aeruginosa* colonies were counted to quantify density, before 24 colonies per replicate were randomly picked using sterile toothpicks. Selected colonies were resuspended in 1 mL of KB media in a deep 96-well plate and grown overnight at 28°C. These were mixed with glycerol (final concentration 25%) and frozen at −80°C. A scraping from each frozen monoculture was then grown in 2 mL of iron-limited casamino acid (CAA: Fisher) medium overnight. Iron limitation was caused by the addition of human apotransferrin (100 mg/mL; BBI Solutions) and sodium bicarbonate (20 mM; Acros Organics) to induce the production of siderophores. Cultures were centrifuged, and the supernatant was assayed using the liquid CAS assay to quantify total siderophores, whilst pyoverdine was quantified by measuring the fluorescence of each culture at 460 nm following excitation at 400 nm. By measuring the optical density of the precentrifuged cultures and quantifying absorbance of sterile media, siderophore production per clone was estimated.
using: \[ 1 - (A_i/A_{ref})/(OD_i) \], where \( OD_i \) = optical density 600 nm and \( A_i \) = absorbance at 630 nm of the assay mixture and \( A_{ref} \) = absorbance at 630 nm of reference mixture (CAA + CAS) (Harrison & Buckling, 2005; Hesse et al., 2019).

### 2.6 | Galleria mellonella virulence assay

The insect infection model *Galleria mellonella* was used to quantify *P. aeruginosa* virulence (Guerrieri et al., 2019; Hernandez et al., 2019). Defrosted freezer stocks containing the whole sample microbiome were diluted 100-fold using M9 salt buffer, before 10 μL was injected into 20 final instar larvae per replicate using a 50-μL syringe (Hamilton). Injected larvae were incubated at 37°C and mortality was monitored hourly after 13 h for 12 h with a final check at 42 h. Larvae were classed as dead when mechanical stimulation of the head caused no response (Hernandez et al., 2019). M9-injected and non-injected controls were used to confirm mortality was not due to injection trauma or background *G. mellonella* mortality; >10% control death was the threshold for re-injecting (no occurrences). Prior to assays on the samples containing *P. aeruginosa*, we confirmed the natural microbial community caused zero mortality by injecting replicates not inoculated with *P. aeruginosa* as described above, except with mortality checks being daily after the first 24 h and the final check being at 93 h.

### 2.7 | Antibiotic resistance assay

To test the evolved tolerance of *P. aeruginosa* to the clinically relevant antibiotics apramycin, cefotaxime and trimethoprim, we used the same *P. aeruginosa* colonies isolated for the siderophore analysis. We first determined the minimum inhibitory concentration of the three antibiotics for our ancestral strain, by growing the ancestral strain for 24 h (as described above) and plating it on TSA containing a range of concentrations of the antibiotics that increased in 10 μg/mL increments from 0 to 60 μg/mL. The minimum inhibitory concentrations were found to be 12, 30 and 40 μg/mL for apramycin, cefotaxime and trimethoprim, respectively. Next, the individual evolved clones were defrosted before 2 μL of each was plated onto either plain TSA, TSA containing apramycin (Sigma: 15 μg/mL) cefotaxime (Molekula: 50 μg/mL) or TSA containing trimethoprim (Sigma: 60 μg/mL). Strains were considered tolerant if colonies could be observed after 48 h. The ancestral strain was used as a negative control.

### 2.8 | Statistical analysis

The effect of liming and shaking, plus their interaction, on the final pH, density of *P. aeruginosa* \( (\log_{10}(cfu/mL)) \) and the proportion of total bioavailable iron \( (Fe^{2+}+Fe^{3+}) \) that was \( Fe^{2+} \) \( (Fe^{2+}/(Fe^{2+}+Fe^{3+})) \) was tested using linear models with liming and shaking as explanatory variables. In general, model reduction was carried out by sequentially removing terms from the full model and comparing model fits using F-tests; we report parameter estimates of the most parsimonious model. The effect of pH on the density of *P. aeruginosa* populations was tested using a linear model with density (cfu/mL) \( \log_{10} \) transformed.

To test whether evolved samples had greater resistance to copper than the ancestral strain, we first calculated the relative fitness, \( w \), of each population by dividing its maximum optical density after 18 h when grown with copper \( (OD_{max,W}) \) by its maximum optical density when grown without copper \( (OD_{max,WC}) \), that is, \( w = OD_{max,WC}/OD_{max,W} \). We then carried out a one-way ANOVA with \( w \) as the response variable and treatment (including ancestor) as the explanatory variable. Secondly, we carried out a Dunnett’s test, using the ‘DescTools’ R package (Signorell et al., 2019), to test whether each treatment differed from the ancestor. Finally, we tested the effect of liming and shaking on the metal resistance of the final populations in a linear model, with \( w \) as the response variable, and liming, shaking and their interaction as the explanatory variables. In all tests, \( w \) was log-transformed to normalize the residuals.

To test liming and shaking effects on total siderophore and pyoverdine production, linear mixed-effects models (LMM) were carried out using the ‘Ime4’ package (Douglas Bates et al., 2015) with liming and shaking as explanatory variables, and random intercepts fitted for each replicate to control for multiple clones being sampled from the same microcosm. For these LMMs, we used the ‘DHARMa’ package (Hartig, 2021) to check residual behaviour, after which the most parsimonious model was arrived at by comparing models with and without the liming-shaking interaction using \( \chi^2 \)-tests. Two samples had pyoverdine values much lower than the rest, so a Grubbs test (‘outliers’ package; (Komsta & Kostma, 2007)) was used to check whether they were significant outliers. They were and therefore were removed from this and all further models to improve model fit. To test the association between copper resistance and both total siderophore and pyoverdine production, we carried out two linear models with log(\( w \)) as the dependent variable and either mean total siderophore production per microcosm or mean pyoverdine production per microcosm as the explanatory variable.

Virulence was analysed in three separate models. First, we tested whether larvae that died before 42 h were injected with samples containing more siderophores and pyoverdine than those that remained alive after 42 h. This was done by carrying out two separate binomial generalized linear mixed models (GLMM) using the ‘Ime4’ package (Douglas Bates et al., 2015), with number of *G. mellonella* dead versus alive as the binomial response variable, and either the production of total siderophores or pyoverdine as the explanatory variable. In this model, pyoverdine production was \( \log_{10} \) transformed to normalize residuals. Secondly, we tested whether the mean time it took deceased larvae (20 per replicate) to die was associated with total siderophore and pyoverdine production (both values taken from the mean of 24 clones) using a linear model. Finally, we tested whether virulence differed between treatments.
To do this, survival curves were fitted using Bayesian regression in the R package ‘rstanarm’ (Brilleman et al., 2020) and the package ‘tidybayes’ (Kay, 2019) was used to estimate parameters. A proportional hazards model with an M-splines baseline hazard was fitted, with liming, shaking plus their interaction as fixed effects. We additionally included random intercepts for each sample to control for multiple (Rutten et al., 2010) G. mellonella being inoculated with the same sample. Models used three chains with uninformative priors and were run for 3000 iterations. Model convergence was assessed using Rhat values (all values were 1) before we manually checked chain mixing.

3.1 | Liming and shaking affected pH but not the relative abundance of Fe$^{2+}$

Here, we tested whether liming of metal-contaminated aquatic environments decreases co-selection for virulence and antibiotic resistance in the opportunistic pathogen *P. aeruginosa*. To do this, we evolved *P. aeruginosa* with or without lime in microcosms containing a mixture of metal-contaminated river water and sediment in the presence of the natural microbial community. We employed both shaking and static microcosms to represent turbulent and stagnant aquatic environments, in order to test whether liming effects were dependent on environmental structure (Figure 1).

As expected, liming significantly decreased the acidity of sediment and water from the initial pH of 5.8. However, the extent of this effect was significantly greater in the shaking treatments (liming-shaking interaction: $F_{1,20} = 23.1, p < 0.001$; Figure 2), likely due to the increased mixing of lime and oxygen throughout the microcosms. The shaken-limed treatment reached a pH of 7.2 (±0.11 SD) whereas the static-limed treatment reached a pH of 6.7 (±0.25 SD). Both non-limed treatments had a final pH of 5.7 (±0.19 SD). As pH is often a good predictor of iron speciation (Gotoh & Patrick Jr, 1974), we tested how the treatments affected the relative proportions of Fe$^{2+}$ and Fe$^{3+}$. We found the proportion of more bioavailable Fe$^{2+}$ to not significantly differ as a result of liming, shaking, nor their interaction (lime main effect: $F_{1,9} = 3.47, p = 0.10$; shaking main effect: $F_{1,9} = 3.00, p = 0.12$; lime-shaking interaction $F_{1,8} = 0.73, p = 0.42$; Figure 2), with Fe$^{2+}$ making up 82% of the total iron available on average across the treatments. Given that iron speciation remained similar in all treatments, this indicates that the redox potential within the microcosms did not change to become more anaerobic under static conditions (Gotoh & Patrick Jr, 1974). Hence iron bioavailability was not significantly influenced by the different experimental conditions and therefore iron limitation was unlikely to represent a significant driver for siderophore production.

3.2 | Evolved *P. aeruginosa* populations had greater tolerance to copper than the ancestor

In order to test whether our river water and sediment samples selected for greater metal tolerance, we incubated the ancestral *P. aeruginosa* strain and final populations in a medium containing a high concentration of copper (1 g/mL of copper sulphate). We then compared the maximum optical density of each culture relative to that of cultures grown without copper ($w$). The ancestral strain had a lower relative fitness ($w$) when grown with copper than all final populations (Dunnett’s test: $p = 0.017$ for all contrasts; Figure 3), consistent with the presence of toxic metals. However, when comparing the effect of the different treatments on $w$, we found populations from the non-limed treatments to not have greater relative fitness in a toxic copper environment than those from the limed treatments (liming main effect: $F_{1,20} = 3.42, p = 0.079$; Figure 3), and shaking to not have an effect on relative fitness (shaking main effect: $F_{1,20} = 2.75, p = 0.11$; liming-shaking interaction: $F_{1,19} = 0.18, p = 0.68$). Growth curves are presented in Figure S1. The optical density of one replicate in the shaken, limed treatment fluctuated after 16 h (likely due to the presence of air bubbles) and consequently this single replicate was removed from the analysis (Figure S2). These results suggest that our river microcosms imposed selection on *P. aeruginosa* to evolve metal tolerance.
3.3 Neither liming nor shaking affected P. aeruginosa density or siderophore production

Next, we tested the treatment effects on P. aeruginosa density and siderophore production. The final density of P. aeruginosa after 2 weeks of evolution varied substantially between samples (1.1 x 10^8 ± 1.6 x 10^6 SD cfu/mL) but was not significantly affected by liming, shaking, nor their interaction (liming main effect: F_{1,21} = 1.96, p = 0.18; shaking main effect: F_{1,21} = 2.77, p = 0.11; liming-shaking interaction: F_{1,21} = 0.70, p = 0.41). There was also no significant effect of pH on P. aeruginosa density (F_{1,22} = 0.97, p = 0.36). Although pH can affect bacterial density (Dennis et al., 2009), our finding of no effect is consistent with previous results demonstrating that P. aeruginosa densities are similar across an equivalent pH range as used here (Moriarty et al., 2007).

To test whether liming and shaking affected siderophore production, both total siderophore production and the production of pyoverdine—the primary siderophore produced by P. aeruginosa (Granato et al., 2016)—were measured for 24 clones per replicate (24 x 24 clones). Quantifying pyoverdine production in addition to total siderophores is important, as it is a key virulence factor in P. aeruginosa, but its production does not necessarily correlate with that of other siderophores, such as pyochelin (Dumas et al., 2013).

We found neither liming, shaking nor their interaction significantly affected mean total siderophore production (liming main effect: χ^2 = 1.49, df = 1, p = 0.22; shaking main effect: χ^2 = 0.49, df = 1, p = 0.48; liming-shaking interaction: χ^2 = 0.08, df = 1, p = 0.78) or pyoverdine production (liming main effect: χ^2 = 0.56, df = 1, p = 0.46; shaking main effect: χ^2 = 2.3, df = 1, p = 0.13; liming-shaking interaction: χ^2 = 1.14, df = 1, p = 0.29). However, we note that there was a large variation in production between the 24 clones used to represent each microcosm (mean production: total siderophores = 4.23; pyoverdine = 766; σ replicate: total siderophores = 1.94; pyoverdine = 69.3), and that two pyoverdine values were significant outliers and consequently were removed from all further analysis in order for model assumptions to be met [these were one from the non-limed shaken treatment (pyoverdine production = 26.9, p < 0.001) and one from the limed-static treatment (pyoverdine production = 174, p < 0.001), which were lower than the pre-removed mean pyoverdine production of 710.6 and median of 789.8]. As we found no significant difference in mean siderophore production between the treatments, nor could we visually identify any cheats, we did not test for differences in the frequency of cheats in our samples. That siderophore production, which is regulated by iron availability and the presence of toxic metals, did not significantly differ between treatments concurs with the non-significant differences in Fe^2+ availability and copper tolerance between treatments. Next, we tested whether either total siderophore or pyoverdine production was associated with copper tolerance, and found neither of them to be (Total siderophores: F_{1,20} = 0.013, p = 0.91; Pyoverdine: F_{1,20} = 0.294, p = 0.59). This suggests other metal resistance mechanisms, such as decreased outer-membrane permeability and increased induction of ATPase efflux transporters, could be responsible for the increased copper tolerance of evolved populations (Teitzel et al., 2006). Our finding of no significant differences in siderophore production contrasts with that of Hesse et al. (2019), who found that the addition of lime to soils collected in the near vicinity of our locality significantly reduced community-wide siderophore production. This difference is most likely due to shifts in siderophore production driven by changes in community composition with liming selecting for non-producing isolates (Hesse et al., 2019), whereas here we solely focussed on siderophore production by P. aeruginosa. This suggests that although liming reduces community-wide siderophore production in metal-contaminated acidic soils, this effect may not be seen in specific species. Interestingly, P. aeruginosa has been proposed as a suitable siderophore-producing bacterium for use in phytoremediation, which relies on the combined use of microorganisms and plants to aid toxic metal remediation (Rajkumar et al., 2012; Sinha & Mukherjee, 2008). It has been proposed that liming, by reducing siderophore production, may hinder phytoremediation (Hesse et al., 2019) as metal uptake by plants is often increased when metals are bound to bacterial siderophores. Given that no significant effect of liming on siderophore production by P. aeruginosa was observed, we suggest that liming and P. aeruginosa-assisted phytoremediation could be used simultaneously without compromise. Future work is needed to test whether other, less-pathogenic (and thus potentially less problematic), pseudomonads such as P. lurida (Kumar et al., 2021), may be more appropriate for bioremediation purposes.
3.4 | Virulence did not differ between treatments but was positively associated with siderophore production

As we found a large variation in siderophore production, which is a known virulence factor in *P. aeruginosa* (Buckling et al., 2007), we tested whether virulence, quantified using the *G. mellonella* infection assay, differed as a consequence of pyoverdine production, total siderophore production or treatment. We note that all injected samples also contained the natural microbial community, but that prior testing with samples not inoculated with *P. aeruginosa* showed that this was of very low virulence (mean death across all treatments after 93 h = 11.5%; Figure S3). Firstly, we tested whether *G. mellonella* larvae alive at the final time check (42 h) had been injected with populations producing less total siderophores and pyoverdine compared with larvae that died before this point and found that they were (total siderophores: $\chi^2 = 6.11$, df = 1, $p = 0.013$; pyoverdine: $\chi^2 = 6.98$, df = 1, $p = 0.004$). Next, we tested whether increased siderophore and pyoverdine production resulted in increased virulence. We found a significant positive association between virulence (mean time to death per population) and both total siderophore and pyoverdine production (total siderophores: $F_{1,22} = 8.9$, $p = 0.007$; Figure 4a; pyoverdine: $F_{1,20} = 10.3$, $p = 0.004$; Figure 4b), with every one unit increase in total siderophores associated with an increase of 8.9 h in the mean time to death.

**FIGURE 4** Mean virulence of *Pseudomonas aeruginosa* evolved in metal-contaminated aquatic communities as a function of (a) mean total siderophore production and (b) mean pyoverdine production. Virulence was quantified using the *Galleria mellonella* infection model ($n = 20$ per replicate) and given as the mean time to death. Pyoverdine and total siderophore production were measured in standardized fluorescence units per OD$_{600}$. Individual circles show the mean production by 24 clones from each replicate. Colours and shapes represent different treatments: grey and □ = static, no lime, blue and + = static, limed, black and △ = shaken, no lime and red and ✶ = shaken, limed. Panel (c) shows the change in the survival probability of larvae over time within each treatment. These do not significantly differ from one another. Shaded areas represent 95% confidence intervals.
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siderophore production decreasing mean time to death by 1.42 h (SE = 0.4745), and every one unit increase in pyoverdine production decreasing the mean time to death by 0.032 h (SE = 0.0098). This positive association between siderophore production and virulence is supportive of previous work both in G. mellonella (Lear et al., 2022) and murine models (Kang et al., 2019). In addition to directly aiding iron uptake within a host, siderophores can increase virulence by triggering other virulence factors (Lamont et al., 2002; Lopez-Medina et al., 2015). Finally, virulence was compared between treatments using survival curves (Figure 4c). Virulence did not significantly differ as a function of treatment, with the credible intervals for liming, shaking and their interaction all crossing 1. No significant treatment effect on virulence is concurrent with the finding that the treatments did not significantly affect siderophore production. Finding virulence to not be significantly different between structured (static) and unstructured (shaking) environments contrasts with findings by Granato et al. (2018), who found that pyoverdine-mediated virulence in P. aeruginosa was greater when grown in solid media than in liquid. The lack of changes detected in siderophore production and virulence between the experimental treatments might be due to the more subtle (and arguably more realistic) conditions under which spatial structure was varied in our study, as well as the presence of a resident microbial community.

3.5 | Antibiotic resistance evolution

As metal pollution has been shown to co-select for antimicrobial resistance (Baker-Austin et al., 2006), we tested whether lime addition altered P. aeruginosa tolerance to the clinically relevant antibiotics apramycin (15 μg/mL), cefotaxime (50 μg/mL) and trimethoprim (60 μg/mL) after evolution in metal-contaminated river water and sediment whilst embedded in the resident microbial community. The concentrations of each of the antibiotics used are greater than the MIC of the ancestral strain, which was determined prior to this assay. Circles show individual replicates; those with a red outline are from the same sample, which is the least resistant to all three antibiotics.

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FIGURE 5 The proportion of 24 Pseudomonas aeruginosa clones per replicate (n = 6) resistant to (a) apramycin (15 μg/mL), (b) cefotaxime (50 μg/mL) and (c) trimethoprim (60 μg/mL) antibiotics. Clones were tested after 2 weeks of evolution in microcosms containing metal-contaminated river water and sediment whilst embedded in the resident microbial community. The concentrations of each of the antibiotics used are greater than the MIC of the ancestral strain, which was determined prior to this assay. Circles show individual replicates; those with a red outline are from the same sample, which is the least resistant to all three antibiotics.
neither was the loss of spatial structure via shaking. A plausible reason for this is that liming reduces metal bioavailability by precipitating ions from solution into the solid phase. This would mean cells in the sediment (the vast majority of the population) would still be exposed to metals where, although at a lower bioavailability, they can still be a cause of co-selection (Dickinson et al., 2019). Although we did not determine the mechanistic basis of co-selection, we note that cross-resistance, co-resistance and co-regulation mechanisms have all been reported for pseudomonads, and the altering of cellular targets is a mechanism commonly used by P. aeruginosa to tolerate metal, trimethoprim and beta-lactam antibiotics such as cefotaxime (Conejo et al., 2003; Sarma et al., 2010). We are aware of a single study testing the effects of liming on antimicrobial resistance (Ramos et al., 1987). This study found liming decreased the susceptibility of Rhizobium species from soil to multiple antibiotics and hypothesized this was due to a greater production of natural antibiotics at near-neutral pH selecting for resistance. Although we note that increasing soil pH will generally decrease the bioavailability of any metals present, the authors (Ramos et al., 1987) stated that metal effects would not be operative in their study, suggesting no metal contamination was present.

4 | CONCLUSION

Pseudomonas aeruginosa populations evolved metal resistance after 2 weeks. However, liming and spatial structure (shaking) were observed to have little effect on P. aeruginosa pathogenic traits. Despite finding a positive association between siderophore production and virulence, neither siderophore production nor virulence systematically differed between treatments, suggesting that liming does not alter the effect of metals on siderophore-mediated virulence in P. aeruginosa. This finding also implies that concurrent use of liming and P. aeruginosa-assisted phytoremediation techniques is possible in scenarios where this bacterium can persist in a natural community. Moreover, we found P. aeruginosa rapidly evolved tolerance to three clinically relevant antibiotics regardless of treatment. We therefore show that a common metal remediation method did not reduce metal pollution-based co-selection for virulence or antibiotic resistance. Importantly, these findings further our understanding of how key determinants of pathogenicity evolve outside of the clinical setting and further demonstrate that metal-polluted environments can select for them.

ACKNOWLEDGMENTS

The acknowledgments, patient consent statement, permission to reproduce material from other sources and clinical trial registration sections are not applicable to this work.

FUNDING INFORMATION

LL would like to thank NERC FRESH GW4 award no. NE/R011524/1, EH: UKRI Future Leaders Fellowship award MR/V022482/1, LN: NERC award NE/W006820/1, WG: NERC award NE/N019717/1, AB: NERC award NE/V012347/1 and MV: NERC award NE/T008083/1.

CONFLICT OF INTEREST STATEMENT

The authors declare that there are no competing interests.

DATA AVAILABILITY STATEMENT

All data (https://doi.org/10.5281/zenodo.8074101) and code (https://doi.org/10.5281/zenodo.8075152) are publicly available on Zenodo.

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**How to cite this article:** Lear, L., Hesse, E., Newsome, L., Gaze, W., Buckling, A., & Vos, M. (2023). The effect of metal remediation on the virulence and antimicrobial resistance of the opportunistic pathogen *Pseudomonas aeruginosa*. *Evolutionary Applications*, 16, 1377-1389. [https://doi.org/10.1111/eva.13576](https://doi.org/10.1111/eva.13576)