Dying for Good: Virus-Bacterium Biofilm Co-evolution Enhances Environmental Fitness

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Abstract: Commonly used in biotechnology applications, filamentous M13 phage are non-lytic viruses that infect E. coli and other bacteria, with the potential to promote horizontal gene transfer in natural populations with synthetic biology implications for engineering community systems. Using the E. coli strain TG1, we have investigated how a selective pressure involving elevated levels of toxic chromate, mimicking that found in some superfund sites, alters population dynamics following infection with either wild-type M13 phage or an M13-phage encoding a chromate reductase (Gh-ChrR) capable of the reductive immobilization of chromate (ie, M13-phageGh-ChrR). In the absence of a selective pressure, M13-phage infection results in a reduction in bacterial growth rate; in comparison, in the presence of chromate there are substantial increases in both cellular killing and biomass formation following infection of E. coli strain TG1 with M13-phageGh-ChrR that is dependent on chromate-reductase activity. These results are discussed in terms of community structures that facilitate lateral gene transfer of beneficial traits that enhance phage replication, infectivity, and stability against environmental change.

Keywords: bioremediation, chromate reduction, community stability, population dynamics, selective pressure, synthetic biology, temperate phage
Introduction

Bacteriohage (phage), a bacterial virus, is about 1/40th the size of most bacteria and represent the simplest, most abundant organism on earth, thriving wherever bacteria grow—with an estimated $10^{30}$ viral particles in the Earth’s oceans alone.\(^1\) Metagenomic studies indicate that phage genes are widely present in bacterial genomes from groundwater samples, including those at major superfund sites (eg, Rifle, CO).\(^2\) Phage are suggested to play an important role in the biogeochemical cycle by controlling marine and other bacterial and phytoplankton communities, thereby influencing pathways of matter and energy transfer within global ecosystems.\(^6\)

At the population level, phage-mediated bacterial lysis results in boom-bust cycles of virus and bacterial host abundance increases, and imposes a well-understood co-evolutionary fitness in which hosts evolve novel phage adaptations to avoid infection, while viruses evade host defenses to retain their infectivity. Such measurements form the basis for theoretical models that support what has become known as “kill-the-winner” hypothesis,\(^7\) in which successful bacterial hosts that become abundant in the environment become targets of viral attack. This negative density-dependent selection leads to increased host diversity, and has been suggested to be critical to community stability. The co-evolutionary dynamics of the model are characteristic of the well known “Red Queen” effect,\(^8\) whereby both viruses and hosts show continual evolutionary adaptation while maintaining broad constancy in relative fitness. While such models are broadly consistent with a large number of ecological theories that describe population dynamics, these models typically do not take into account positive selective pressures whereby non-lytic phage might act to provide an ability for host to exhibit enhanced fitness through lateral gene transfer, and the potential of population dynamics to allow shifts in the metabolic capacities of populations that enhance their fitness against environmental change.

Prior measurements indicate that unlike lytic phages, which can dramatically disrupt microbial communities and the formation of biofilms through bacterial cell wall lysis,\(^9\) that M13 and other temperate (filamentous) phages can enhance growth and biofilm formation.\(^10,11\) Although the underlying mechanisms remain uncertain, it has been suggested that a major contribution to community stability involves the presence of extracellular DNA arising from cell death, which is thought to represent an important matrix element necessary for the formation of biofilms. Additional factors that enhance microbial growth may be related to the ability of many nonlytic phage, including M13, to promote horizontal gene transfer and the rapid acquisition of desired metabolic functionalities (eg, enzyme activities) that favor community stability.\(^12\) Examples include the ability of microbial populations to mobilize natural genetic variation in response to environmental change that enhance fitness. While these latter mechanisms are commonly suggested to involve the presence of naked DNA that arises through natural mechanisms of cellular death unrelated to mechanisms of lateral gene transfer, recent data suggests a coordination between DNA release and uptake within the population.\(^14\)

To better understand the possible role of phage in promoting lateral gene transfer, and their relevance to possible applications involving bioremediation, we have investigated the community dynamics associated with the phage M13 and an $E. \text{coli}$ host. In these measurements, a phagemid vector was constructed that encodes a chromate reductase (Gh-ChrR) (ie, M13-phage$_{\text{Gh-ChrR}}$), where Gh-ChrR has previously been demonstrated to efficiently reduce toxic chromate \((\text{Cr(VI)})\) in the presence of extracellular reductants naturally present within soils as humic compounds (such as quinones).\(^15\) Infection only occurs in the presence of a helper phage, which is essential for phage maturation and assembly, permitting an understanding of possible differences between initial infection and subsequent transfer of Gh-ChrR within the bacterial population. Using these constructs, we have assessed how the presence of toxic chromate modifies $E. \text{coli}$ growth and viability, focusing on the possible role of phage infection in promoting community stability. We find that in comparison to wild-type M13 phage (not expressing Gh-ChrR), that M13-phage$_{\text{Gh-ChrR}}$ infection of $E. \text{coli}$ results in substantial increases in the population dynamics, resulting in enhanced bacterial growth, cell death, and total biofilm formation of $E. \text{coli}$ strain TG1.

Results

Expression of dimeric chromate reductase on the surface of M13 phage

The DNA sequence of the Gh-ChrR gene from $\text{Gluconacetobacter hansenii}$ ATCC 23769 (ZP_06834583) was codon optimized for expression in $E. \text{coli}$,
synthesized, and inserted into the expression vector pJexpress411. As chromate reductase (ChrR) from *Gluconacetobacter hansenii* functions as a dimeric enzyme,\(^\text{18}\) we have created a fusion protein containing two copies of the gene encoding Gh-ChrR. The fusion construct was engineered to allow in-frame expression of ChrR with the heterodimeric subunits Fos and Jus, whose high-affinity enhances the likelihood that Gh-ChrR will form a stable protein complex on the cell surface\(^\text{20}\) (Fig. 1A). The sequence of the engineered construct was confirmed by DNA sequencing. To ensure that Gh-ChrR forms a functional protein of ChrR with the heterodimeric subunits Fos and Jus, whose high-affinity enhances the likelihood that Gh-ChrR will form a stable protein complex on the cell surface\(^\text{20}\) (Fig. 1A). The sequence of the engineered construct was confirmed by DNA sequencing. To ensure that Gh-ChrR forms a functional protein

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**Figure 1.** Functionally active chromate reductase is expressed on the surface of purified M13-phage \(_{\text{Gh-ChrR}}\). (Panel A) Map of engineered pCDisplay-4 phagmid vector encoding Gh-ChrR (green arrows) expressed as a fusion protein with Fos-Gh-ChrR and Jun-Gh-ChrR to facilitate dimerization following phage display. (Panel B) Kinetic reduction of chromate (monitored at 370 nm) by purified M13-phage \(_{\text{Gh-ChrR}}\) (red squares) in comparison to m13 helper phage (blue triangles). Notes: Measurements involved purified phage (5 \(\times\) 10\(^{10}\) pfu/mL) in 50 mM Tris-HCl (pH 7.4), 100 mM NaCl, 0.5 mM Cr\(_2\)O\(_4\), and 0.1 mM NADH. Chromate reduction rates for M13-phage \(_{\text{Gh-ChrR}}\) (\(\Delta\text{OD}_{370nm} = 0.006/min\)) are significantly increased in comparison to that observed for helper phage alone (\(\Delta\text{OD}_{370nm} = 0.004/min\)) \((P = 0.00015)\).
on the surface of M13-phage\textsubscript{Gh-ChrR}, we isolated individual phage following overnight infection of \textit{E. coli} strain TG1 host in the supernatant following cell pelleting. Phage concentrations were determined following infection of \textit{E. coli} TG1 host with dilutions of phage stock based on counts of ampicillin-resistant plaques visible on an \textit{E. coli} lawn. The ability of purified M13-phage\textsubscript{Gh-ChrR} ($5 \times 10^{10}$ pfu/mL) to reduce chromate, as previously observed for the purified protein,\textsuperscript{18} was measured at $37$ °C in triplicate under aerobic conditions (Fig. 1B). In comparison to wild-type phage (or M13 helper phage not expressing Gh-ChrR) there is a substantial increase in the rate of chromate (Cr(VI)) reduction, consistent with the presence of a functional dimer of the Gh-ChrR enzyme ($P$-value 0.00015). In comparison with the functional dimeric complex, expression of a monomeric ChrR protein on the M13 phage surface does not promote enhanced chromate reductase activity. This latter result is consistent with prior measurements of ChrR structure, which suggest that the catalytic center is near a dimeric interface.\textsuperscript{21–24} The requirement that the Gh-ChrR dimeric complex be stabilized as a fusion protein through complexation of the well understood Jun/Fos interaction suggests that the Gh-ChrR dimer is not sufficiently stable upon expression on the cell surface to maintain the native oligomeric state formed within the crowded cellular milieu. Our results are consistent with prior suggestions that functional chromate reductase activity requires a dimeric association.

Enhanced biomass formation upon infection of \textit{E. coli} with M13-phage\textsubscript{Gh-ChrR}

Using high concentrations of toxic chromate as a selective pressure, we examined the influence of infection with M13-phage\textsubscript{Gh-ChrR} on \textit{E. coli} viability and growth (measured as total biomass). In these experiments, static biofilms of \textit{E. coli} strain TG1 (with and without early infection of M13-phage\textsubscript{ChrR}) were formed in 96-well polystyrene plates. In comparison to wild-type cells (or those infected with either phagmid–Gh-ChrR or helper phage alone), there are substantial increases in total biomass of \textit{E. coli} strain TG1 infected with M13-phage\textsubscript{Gh-ChrR} (Fig. 2). Corresponding increases in chromate reduction are observed in the presence of M13-phage\textsubscript{Gh-ChrR} that are apparent approximately 34 hours following infection; increases in chromate reduction rates require the presence of both phagmid–Gh-ChrR and helper phage (ie, propagation of phage encoding Gh-ChrR). These results indicate that the enhanced reduction of chromate and corresponding tolerance of toxic chromate represent a selectable pressure that can promote bacterial growth and enhanced remediation.

Further examination of biofilms formed using the \textit{E. coli} strain TG1 involved their visualization using confocal microscopy. In these experiments, parallel measurements were made using an 8-well chamber slide using equal densities of dilute \textit{E. coli} TG1 bacteria in LB media at $37$ °C, permitting side-by-side comparisons (in duplicate) of biofilms comprised of TG1 alone, following tranformation with either phagmid-Gh-ChrR or helper phage alone, or the simultaneous transformation with both phagmid-Gh-ChrR and helper phage necessary for the maturation and propagation of M13-phage\textsubscript{Gh-ChrR}. Following phage exposure (16 hr), bacterial biofilms were washed twice, incubated with both BacLight Live/Dead stain, fixed, and analyzed using confocal microscopy. Irrespective of whether \textit{E. coli} biofilms were exposed to wild-type phage, phagmid-ChrR alone, or helper phage alone, we observe that the majority of cells are viable (ie, green), with minimal cell death (visualized by staining by propidium iodide)(PI) (ie, red) (Fig. 2, top panels). In comparison, upon co-infection with both phagmid-Gh-ChrR and helper phage it is apparent that there is a substantial amount of cell death, despite the large increase in biofilm thickness. These latter results suggest that phage infectivity promotes enhanced growth despite concurrent increases in cell killing.

Discussion

We have demonstrated that the need for host tolerance to high concentrations of toxic chromate limits the ability of hosts to evade viral attacks and creates an evolutionary trade-off between growth rate maximization and defense that enhances microbial community co-evolution. In our experiments, the host \textit{E. coli} bacterium challenged with infective phage encoding a functional chromate reductase experiences substantial amounts of both cellular killing and enhanced growth, leading to increases in biomass production and biofilm formation. These results indicate that phage infectivity can stabilize community structures, leading to more bacteria proliferation and initiation.
Co-evolution of community interactions for bioremediation

of a beneficial boom-bust cycle to produce more phages. Our measurements build on prior biosign principles that indicate the ability of M13 and other temperate phages to enhance biofilm formation.\textsuperscript{10,11} Central to this capability is the functional capacity of the M13-phage to express a function (ie, chromate reductase activity) that enhances fitness to a defined selective pressure (ie, chromate toxicity). Under these latter conditions, there are substantial increases in total biomass and biofilm thickness, despite substantial increases in phage-mediated cellular killing. These results indicate that the enhanced bioremediation of chromate and corresponding tolerance of toxic chromate by microbial communities represent a selectable pressure that can be engineered to promote community stabilization and enhanced remediation.

In nature bacteria commonly form biofilms, which represent natural microbial communities that form on virtually any surface exposed to water. Such biofilms are a common target of antimicrobials that seek to minimize contamination in medical, industrial, and food processing. However, the establishment of beneficial microbial communities is now recognized to have considerable importance for human health, and in natural environments may enhance stability to environmental change. In this respect, biofilm formation has been suggested to be critical for the environmental radionuclide waste bioremediation process, as observed following acetate injection experiments from US

Figure 2. Phage-dependent lateral gene transfer of chromate reductase by M13-phage results in enhanced biomass formation. Confocal images of live (Cyto9; green) or dead (PI; red) cells (top panels), rates of chromate reduction (middle panels), and measured biomass (bottom panels) for mature biofilms comprised of E. coli strain TG1 alone (left panels), in the presence of either phagmid Gh-ChrR or helper phage alone (center panels), or following infection (right panels). Chromate (5 mM) reduction was monitored by measuring the absorbance at 370 nm wavelength, and is normalized following correction for the absorbance of the LB medium. Total biomass was estimated by crystal violet staining and measured absorbance at 595 nm, and is normalized to that of TG1 alone (left panel).

Note: Significant differences (*) in comparison to TG1 alone (P-value < 0.01) were based on a Student’s t-test (n = 3).
Department of Energy environmental subsurface bioremediation projects. Indeed, subsurface environmental bacterial biofilms contribute to the long-term reductive immobilization of uranium (U(IV)) or chromium (Cr(III)) through formation of precipitates on sediment grain surfaces. As a result, strategies that stabilize bacterial biofilms, such as those described here, are expected to improve the bioremediation efficiency in the subsurface environment. Our results suggest that bacteriophage may play an important role in the biogeochemical cycle by controlling bacterial and phytoplankton communities. As a result, strategies that enhance formation of bacterial biofilms are expected to improve the bioremediation efficiency in the subsurface environment.

Bioremediation of chromate and uranyl toxic metals in contaminated soils remains a major technological challenge. Current solutions involve a range of biological approaches that include the targeted feeding of specific anaerobic microbes (e.g., *Geobacter sulfurreducens*) whose specialized metabolism permits the biosorption, biosequestration, and reductive immobilization of extracellular chromate and uranyl metals, which can serve as terminal electron acceptors of cellular respiration. Likewise, *Shewanella oneidensis* MR-1 contains specialized metal reductases located on the outer membrane that selectively associate with metal oxides to mediate their reduction. The recent structural determination of these metal reductases and the understanding of how cellular machinery regulates the targeted assembly of a metal reductase complex on the outer membrane, all suggest possible synthetic biology approaches to enhance these pathways and promote more effective bioremediation of contaminated sites. Likewise, identification of soluble enzymes capable of reductive immobilization of chromate and uranyl under both anaerobic and aerobic conditions offers a means to re-engineer microbes to enhance bioremediation. In this latter respect, bacteriophage present at natural sites offer a potential means to serve as gene delivery vectors for these synthetic biology applications. Additional mechanisms of bioremediation may take advantage of the ability to display catalytic protein moieties on the surface of bacteriophage, such as we describe, where the reducing potential of available humic compounds act as electron shuttles to allow transfer of reducing equivalents to extracellular catalysts. In this respect, it is necessary to further develop bioengineering strategies to retain viable phage in microbial populations responsive to environmental contaminates, permitting long-term immobilization of toxic metals such as chromate (Cr(VI)).

**Materials and Experimental Procedures**

**Materials**

Restriction endonuclease enzymes were from New England Biolabs (Ipswich, MA). The expression vector pJexpress411 (DNA 2.0 Inc., Menlo Park, CA, USA). Phagemid pCDisplay-4 and *E. Coli* strain TG1 were from Creative Biolab Inc. (Shirley, NY). Isolation of chromosomal DNA, plasmids, and the purification of polymerase chain reaction (PCR) products involved kits obtained from Qiagen (Valencia, CA). BacLight Live/Dead stain was carried out with Live/Dead Bacterial Viability Kit (Molecular Probes, Inc., Eugene, OR) using SYTO9 for green fluorescent nucleic acid stain and PI (propidium iodide) for the red fluorescent nucleic acid stain.

**Construction of dimer chromate reductase on M13 phage**

In-frame fusion mutants of Gh-ChrR-Fos and Gh-ChrR-Jun were designed and synthesized using overhang PCR methods, and then they were cloned into the phagemid of pCDisplay-4 (Creative Biolab Inc., Shirley, NY). The first subunit Gh-ChrR was cloned into pCDisplay-4 with restriction enzyme sites SphI/SpeI, and the second Gh-ChrR fragment was ligated into pFos-Jun-Gh-ChrR-I with SacI/XbaI sites. The recombinant phagemid vector pFos-Jun-Gh-ChrR-II was confirmed using DNA sequencing. The schematic map of pFos-Jun-Gh-ChrR-II is depicted in Figure 1A.

**Amplification and isolation of recombinant M13 phages**

Luria broth (LB) medium (500 mL) was inoculated with host *E. coli* strain TG1 transformed with the phagemid pFos-Jun-Gh-ChrR-II in the presence of ampicillin (100 µg/mL) and incubated at 37 °C (250 rpm) until the optical density at 600 nm reaches 0.8–0.9. To promote phage infection, M13 KO7 helper phage (Life Technologies, Grand Island, NY) was
added \((5 \times 10^9 \text{ pfu/mL})\) in the presence of kanamycin \((50 \mu g/mL)\); cultures were not shaken for 30 min to facilitate infection, followed by gentle agitation \((200 \text{ rpm})\) for 30 min prior to cellular recovery \((2200 \times g, 15 \text{ min})\). The pellet was resuspended in LB \((500 \text{ mL})\) and grown overnight at 30 °C \((300 \text{ rpm})\) in the presence of ampicillin \((100 \mu g/mL)\) and kanamycin \((50 \mu g/mL)\). Following cell pelleting \((7,000 \times g\) for 15 min at 4 °C), the supernatant containing the phage were separated into prechilled 1 L bottles and precipitated upon addition of PEG8000 \((20 g)\) for 1 h on ice. Phage were isolated by centrifugation \((7,000 \times g\) for 15 min in the same bottle at 4 °C) and resuspended in 8 mL of phosphate buffered saline (PBS). Debris and cellular contamination was removed by centrifugation \((12,000 \times g\) for 10 min), phage were isolated in the supernatant, and phage titer was determined by infecting TG1 cells with dilutions of phage stock and following incubation and the counting of the numbers of ampicillin resistant plaques visible. Phage were stored as aliquots at 4 °C.

**Chromate reduction assays**

The ability of Gh-ChrR expressed on phage to reduce Cr(VI) was assayed at 37 °C in a 100 μL assay buffer \((50 \text{ mM Tris-HCl, } 100 \pm 13 \text{ mM NaCl, pH 7.4})\) containing 100 μM NADH, 0.5 mM chromate \((K_2Cr_2O_7)\) and purified phage \((5 \times 10^{10} \text{ pfu/mL})\). The chromate reduction rates were measured by monitoring the absorbance of A370. Data were measured on a SpectraMax 384Plus microplate reader (Molecular Devices, Sunnyvale, CA). Values for the chromate reduction rate from wild type phage and phage expressing Gh-ChrR were estimated by the chromate reduction (based A370) in 2 hours. Both fit well with the linear equation, but the phage expressing Gh-ChrR has a significantly stronger reduction rate. All measurements were conducted in triplicate under aerobic conditions (Fig. 1B). The ability of Gh-ChrR expressed on phage in biofilms to reduce Cr(VI) was assayed directly by measuring the A370 from 100 μL of an aliquot from the culture in 1 mL deep well 96-well microplate with 5 mM chromate (Fig. 2).

**Biofilm formation**

Overnight cultures of *E. coli* strain TG1 were diluted 1:30 in LB medium and grown in polystyrene microtitre plates \((170 \mu l \text{ per well})\) at 37 °C for approximately 3 hours in order to reach mid-log phase. In some cases, added crystal violet \((0.5% \text{ (w/v)})\) retained following washing was measured at 595 nm to assess total biofilm formation, as previously described. Alternative experiments measured living and dead cells using BacLight Live/Dead stain added to each well prior to fixation. Biofilms were visualized using a Zeiss 710 laser scanning confocal microscope with a 20 × NA 1.0 water-dipping objective, simultaneously exciting green (Cyto9) and red (PI) fluorophores respectively at 488-nm and 633-nm. Fluorescence emission was collected using broad-band emission filters with bandwidths allowing 493–628 nm emission to be collected for the green channel and 638–759 nm emitted light for the red channel. Images of the 3-D biofilm projection used the software ImageJ (http://rsb.info.nih.gov/ij/).

**Author Contributions**

Conceived and designed the experiments: HJ. Analyzed the data: HJ, TCS. Wrote the first draft of the manuscript: HJ. Contributed to the writing of the manuscript: HJ, TCS, PEL. All authors reviewed and approved of the final manuscript.

**Funding**

This research is supported by the US Department of Energy Pacific Northwest National Laboratory (PNNL) Laboratory Directed Research and Development Funds (LDRD #90001). Part of the research was conducted at the W.R. Wiley Environmental Molecular Sciences Laboratory, a national scientific user facility sponsored by the US Department of Energy’s Office of Biological located at PNNL. PEL and HJ were also partly supported by BER’s Integrated Field Research Challenge Site (IFRC) at Rifle, CO, a multidisciplinary, multi-institutional project managed by PNNL and LBNL. Battelle operates PNNL for the US Department of Energy under Contract DE-AC05-76RL01830. University of California operates LBNL for the US Department of Energy.

**Acknowledgments**

We thank Dr. Dehong Hu for the technique support of the confocal microscope studies.

**Competing Interests**

Author(s) disclose no potential conflicts of interest.
Disclosures and Ethics
As a requirement of publication, author(s) have provided to the publisher signed confirmation of compliance with legal and ethical obligations including but not limited to the following: authorship and contributorship, conflicts of interest, privacy and confidentiality and (where applicable) protection of human and animal research subjects. The authors have read and confirmed their agreement with the ICMJE authorship and conflict of interest criteria. The authors have also confirmed that this article is unique and not under consideration or published in any other publication, and that they have permission from rights holders to reproduce any copyrighted material. Any disclosures are made in this section. The external blind peer reviewers report no conflicts of interest.

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