


 Cite this: *RSC Adv.*, 2021, **11**, 8346

Regenerating heavily biofouled dissolved oxygen sensors using bacterial viruses†

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Bacterial biofilms are aggregates of bacterial cells embedded in a self-produced extracellular polymeric matrix. Biofilm formation has always been considered a major challenge for sensors used in underwater measurements, and is a primary source of measurement error, especially when it comes to long-term *in situ* monitoring. We demonstrate the utility of lytic bacteriophages (bacterial viruses) as a non-invasive strategy for removing bacterial biofilms formed on the gas permeable membrane of electrochemical dissolved oxygen sensors. Our results show that a 4 day *Pseudomonas aeruginosa* biofilm with a fully developed matrix significantly affected the sensor signal and response time, decreasing the signal by 32% and increasing the response time by 94%. In addition, measurements with the biofouled membrane had a very low signal to noise ratio compared to a clean sensor membrane. A single dose of overnight phage treatment effectively removed the biofilm (as indicated by scanning electron micrographs and fluorescence images of the membrane), without the need for repeated treatments. Furthermore, the sensor signal that had plummeted by 32% for a fully biofouled membrane, was returned to the original value ($7.96 \pm 0.27 \text{ mg L}^{-1}$) after phage treatment and the signal to noise ratio (calculated as the ratio of mean to standard deviation) increased 8 folds for a phage-treated membrane compared to a biofouled membrane. Our data indicate near complete regeneration and signal recovery for the dissolved oxygen sensor, making the biofouled sensor reusable without the use of harsh chemicals that could destroy the fragile sensor membrane.

 Received 2nd December 2020
 Accepted 11th February 2021

DOI: 10.1039/d0ra10156g

rsc.li/rsc-advances

Introduction

Biofouling, or biological fouling, is an unwanted deposition of microorganisms on biotic and abiotic surfaces.¹ Bacteria are major contributors to biofouling events, and they can readily form surface-bound, 3-dimensional communities. These communities are known as bacterial biofilms and are usually encased in a matrix of biopolymers, called the extracellular polymeric substance (EPS), which is secreted by the bacterial cells themselves.⁴ From the point of view of a bacterial cell, biofilms provide a means for survival and that is why bacterial biofilms are the most common form of bacterial existence in

nature. Bacteria form biofilms on infected catheters and implants,² on surfaces in the food industry⁵ and membranes in water treatment plants,⁶ on ship hulls,⁷ and on marine sensors.^{8,9} Biofouling has always been considered a major challenge for underwater measurements, especially when it comes to long-term *in situ* monitoring (*e.g.*, and environmental monitoring devices installed in lakes and rivers).¹⁰ Biofouling of sensor surfaces is a major concern in all physico-chemical sensors, including pH and redox sensors, optical sensors, and electrochemical sensors, and could result in unreliable measurements.^{11,12} Biofilms start to grow as soon as a sensor is immersed in water, and it has been reported that the quality of measurements decreases in less than a week.¹⁰ Although many potential solutions have been proposed to combat biofouling in underwater measurements and monitoring, there is no single universally accepted method applicable to underwater environmental sensors.

Various methods and technologies have been used to prevent biofouling or remove biofilms from biofouled surfaces, such as mechanical treatment (wipers, water jetting, ultrasonic or electrical surface excitation),^{11,13} chemical detergents and biocides,¹⁴ UV light exposure (which may kill bacteria but will not completely remove the biofilm),^{15,16} chlorination,⁸ chemical modifications of surfaces (*e.g.*, using zwitterions, carboxylation,

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† Electronic supplementary information (ESI) available. See DOI: 10.1039/d0ra10156g

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and charged hydrogels), and coating surfaces with antimicrobials¹⁷ or repellent coatings.^{18–20} For sensing surfaces, however, cleaning methods should be chosen with care. Cleaning methods that involve harsh physical or chemical treatments could cause structural damage and harm the sensitive and sometimes delicate sensing surfaces.⁵ In addition, improved intercellular interactions in biofilms leads to enhanced metabolic capacity of cells and antimicrobial resistance.²¹ As a result, biofilms have shown to be more resistant to chemical disinfectants and biocides compared to planktonic (free swimming) bacteria. Antimicrobial or repellent surface coatings are not as invasive as mechanical scrubbing or chemical chlorination, but require changing the sensing surface, which may interfere with measurements and sensor sensitivity due to altered bio-interface properties. Besides, once the coating is depleted, the sensor will start to foul and will thus have to be removed and changed.

Bacteriophages are bacterial viruses that were discovered over a hundred years ago, well before penicillin, and were almost immediately used to treat bacterial infections.^{22,23} Bacteriophages (or phages, for short) are natural bacteria devourers, and their job in nature is to keep bacterial populations in check. They have been proposed and explored as potential biological tools to control or mitigate biofilm formation in different settings.²⁴ These human-friendly viruses are capable of infecting bacteria very specifically, thus not harming the delicate microbial balance in the surrounding environment, a characteristic that has made them attractive for different applications as therapeutics, biocontrol agents, prophylactics and even sensing agents.^{25–27} After binding to bacterial host cell surface receptors, phages insert their genome into the host bacterium. A class of phages known as lytic or virulent phages immediately take control of the host cell machinery and turn the bacteria cell into a phage propagation factory. At the end of this replication process, known as the lytic cycle, the bacterial cell is lysed, and hundreds of new phages are released to infect other bacteria. Temperate or mild phages are another class of phage. Those integrate their genome with that of the host bacteria and lay dormant until activated by an external stimulus such as UV light, temperature change, or a sub-lethal concentration of a toxic chemical, which in turn activate the lytic cycle and ultimately lyse the bacterial cell.^{28,29} Lytic phages are the group of phages that are mostly explored for controlling bacterial populations, because they offer an immediate anti-bacterial action.

Phages are currently heavily investigated for their use as therapeutics and for biomedical applications^{24,30,31} but have remained obscure to most environmental engineers. With only a few reports published on mitigation of membrane fouling^{32,33} and no prior report on regenerating biofouled sensors, there is a gap in knowledge regarding the performance of phage for sensor regeneration, which calls for more investigations in this field. It is noteworthy that chemical antimicrobial agents in general have limited activity against biofilms, because they could be neutralized or diluted to sublethal concentrations by binding to the biofilm matrix before reaching all the individual bacterial cells within the biofilm.³⁴ In addition to propagating

on-site, lytic phages have enzymes that can degrade the biofilm matrix, which is a major advantage over chemical treatment.²⁴ Bacteriophages, however, are not the silver bullet for fighting bacterial biofilms. Even in demonstrated biomedical applications of phage against bacterial biofilms, bacteriophages have been demonstrated to work along with the host immune system to completely eradicate the biofilm, a phenomenon absent in environmental applications. Despite the imperfection, what makes bacteriophages ideal for biomedical applications over antibiotics (mild and non-destructive to non-target bacteria, and evolution of resistant mutants that are less virulent^{13,35}) also makes them important options of investigations for environmental applications. However, for regeneration of submerged sensors, it is important to adequately clean the sensing substrate so as to regenerate a sensing signal without damaging or disintegrating the surface, and so effectiveness of phage treatment is not immediately clear and must be investigated with a correct choice of phage. A wrong choice of phage could be completely ineffective or even result in increased biofilm formation.³⁶

In this work, we use a lytic bacteriophage cocktail to remove a fully developed biofilm of *Pseudomonas aeruginosa* PAO1 (PA) from the membrane of a dissolved oxygen (DO) sensor. The cocktail consists of vB_Pae-Kakheti25 and vB_Pae-TbilisiM32, which are well-characterized *Pseudomonas* phages that we have determined are capable in dispersing established biofilms of PA.³⁷ *Pseudomonas aeruginosa* is one of the most prevalent opportunistic microorganisms, inhabiting soil, seawater, freshwater, sewage, plants, and animals.³⁸ This microorganism is a strong biofilm former and is responsible for difficult to treat hospital acquired infections,³⁹ persistent food contamination,³⁹ and marine biofouling, due to its high capacity to colonize and form biofilms on surfaces submerged in aquatic environments.⁴⁰ It has been shown that other microorganisms are stimulated to settle and colonize the submerged surfaces in marine environments in the presence of the EPS secreted by biofilm forming species such as *Pseudomonas aeruginosa*.³ In addition, studying the biofilms formed on nanofiltration membranes in water treatment plants has confirmed that *P. aeruginosa* is one of the important species dominating the biofilm communities.⁹ As a result, in this proof of concept study, *P. aeruginosa* is used for biofilm formation on the surface of DO sensor membranes as a representative for biofilm forming species in aquatic environments. We demonstrate the effectiveness of a single dose of lytic phage treatment on biofouled sensor membranes in recovering DO sensor signals to the original level, increasing the signal to noise ratio, and decreasing the response time.

Methods

Materials

The ExStik II DO600 dissolved oxygen meter and DO603 membrane kits were purchased from Exttech Instruments (Nashua, USA). Phosphate buffered saline (PBS) (0.1 M, pH 7.4) was prepared using tablets from VWR (Mississauga, Canada). Clear, flat-bottom, sterilized, 96-well tissue culture plates and



crystal violet were also purchased from VWR. LB broth, Miller, sterile 50 mL conical centrifuge tubes, and glutaraldehyde (50%) were purchased from Fisher Scientific (Mississauga, Canada). Hoechst 33342 nucleic acid stain (ex/em 350/461 nm) was purchased from Thermo Fisher Scientific (Massachusetts, USA). Formaldehyde was purchased from Caledon Laboratories (Georgetown, Canada).

Bacterial strain, phage, and culture conditions

The bacterial strain used in this study was *Pseudomonas aeruginosa* PAO1 (PA) and the phages used were vB_Pae-Kakheti25 and vB_Pae-Tbilisi32 (DSMZ, Germany). Bacterial overnights of PA were prepared by inoculating LB broth with a frozen glycerol stock and incubating at 37 °C and 180 rpm for 16–18 h. Phage suspensions were propagated by diluting the overnight culture 1 : 1000 into 50 mL of LB media in a baffled Erlenmeyer flask and incubating at 37 °C and 180 rpm. When the optical density of the culture at 600 nm (OD_{600}) reached 0.5, 10 μ L of either phage vB_Pae-Kakheti25 or vB_Pae-Tbilisi32 was added and the culture and incubated for 6 h, or overnight. The resulting phage lysate was centrifuged for 10 min at 7000 \times g, then the supernatant was filtered through a 0.2 μ m syringe filter and stored in the fridge. The concentration of the phage suspension (number of plaque forming units per mL, PFU mL⁻¹) was determined using the agar overlay technique.⁴¹

Biofilm formation and quantification

Biofilms were formed on the membranes by placing the DO membrane cassette face down in a 50 mL conical tube containing 2 mL of an overnight PA culture, diluted 1 : 1000 in LB broth. The membranes were incubated at 37 °C and 80 rpm for 1–4 days. Biofilm was then quantified through fluorescence microscopy followed by image analysis. For biofilms formed in 96-well plates, we diluted an overnight culture 1 : 1000 in LB broth, added 200 μ L of diluted suspension to each well, and incubated the plate at 37 °C and 80 rpm for 1–4 days. Biofilms were then quantified using the crystal violet method, as described elsewhere.⁴² Briefly, biofilms were washed three times with PBS buffer, stained with crystal violet, and then quantified by dissolving the crystal violet in acetone and reading the absorbance at 590 nm with a BioTek plate reader.

Phage treatment

Membranes with biofilm were rinsed in PBS three times and then incubated in 2 mL of a 1 : 1 mixture of vB_Pae-Kakheti25 and vB_Pae-Tbilisi32 (10^9 PFU mL⁻¹ each) in a 50 mL conical tube at 37 °C and 80 rpm for 6 h or overnight. The control fouled membranes were treated the same way, but incubated in LB broth instead of a phage suspension.

Bleach treatment

Same as phage treatment, membranes with biofilm were rinsed in PBS three times, and then incubated in 2 mL of a 10% bleach solution in a 50 mL conical tube at 37 °C and 80 rpm overnight.

Sensitivity readings

DO concentration of stirred deionized (DI) water was measured using the DO sensor with clean, biofouled, phage-treated and bleached membranes installed. Two membranes were used per condition and the measurements were repeated three times with each membrane. Each experiment was performed in triplicate. The response time for each membrane was recorded as the time the sensor readings showed fluctuations $<\pm 0.01$ mg L⁻¹ for 30 seconds.

Fluorescence microscopy

Biofouled and phage-treated membranes were fixed with 4% formaldehyde in PBS for 30 min and rinsed with DI water. Fixed samples were stained with 400 μ L of 25 μ g mL⁻¹ Hoechst 33342 in DI water for 1 h and rinsed lightly with DI water afterwards to remove unbound fluorescence dye. Images were recorded using Nikon Eclipse Ti2-E motorized inverted fluorescence microscope, and the results were quantified using ImageJ image processing software. For quantification, two membranes were used, and seven frames of each membrane were quantified.

Scanning electron microscopy (SEM)

Membranes were cut out of their cassette and rinsed with PBS three times. Following fixation with 30% glutaraldehyde for 30 min at 4 °C, the membranes were dehydrated in a gradient of ethanol solutions (10%, 30%, 50%, 70%, 90%, 100%) for 10 min each, then dried using a Leica EM CPD300 critical point dryer (Leica Mikrosysteme GmbH, Austria). Those were then coated with 20 nm of gold using a Polaron Model E5100 sputter coater (Polaron Equipment Ltd., UK). SEM images were obtained using a Tecan VEGA II LSU SEM (Tecan USA, PA, USA) at 10 kV and 4700 \times magnification.

Statistical analysis

Data are presented as means \pm standard deviation. Statistical significance was assessed using analysis of variance and *post-hoc* analysis with Tukey's test, where significant difference was considered as $P < 0.05$. All experiments were performed in triplicates unless otherwise specified.

Results and discussion

In this study, PA biofilms were grown on DO sensor membranes following baseline measurements (Fig. 1A). Biofouled membranes were then subjected to overnight incubation in a phage suspension consisting of a 1 : 1 mixture of phages vB_Pae-Kakheti25 and vB_Pae-Tbilisi32 in LB media. The choice of phage was based on previous experience in our lab with a library of PA phages and their performance against PA biofilms in a stationary microtiter plate model. We chose two phages for treating the biofilm instead of one phage, because it has been shown that using more than one phage decreases the frequency of evolution of phage-resistant bacteria,^{43–45} thus increasing the efficiency of phage treatment. To justify the choice of a phage mixture, we demonstrated that the two phages



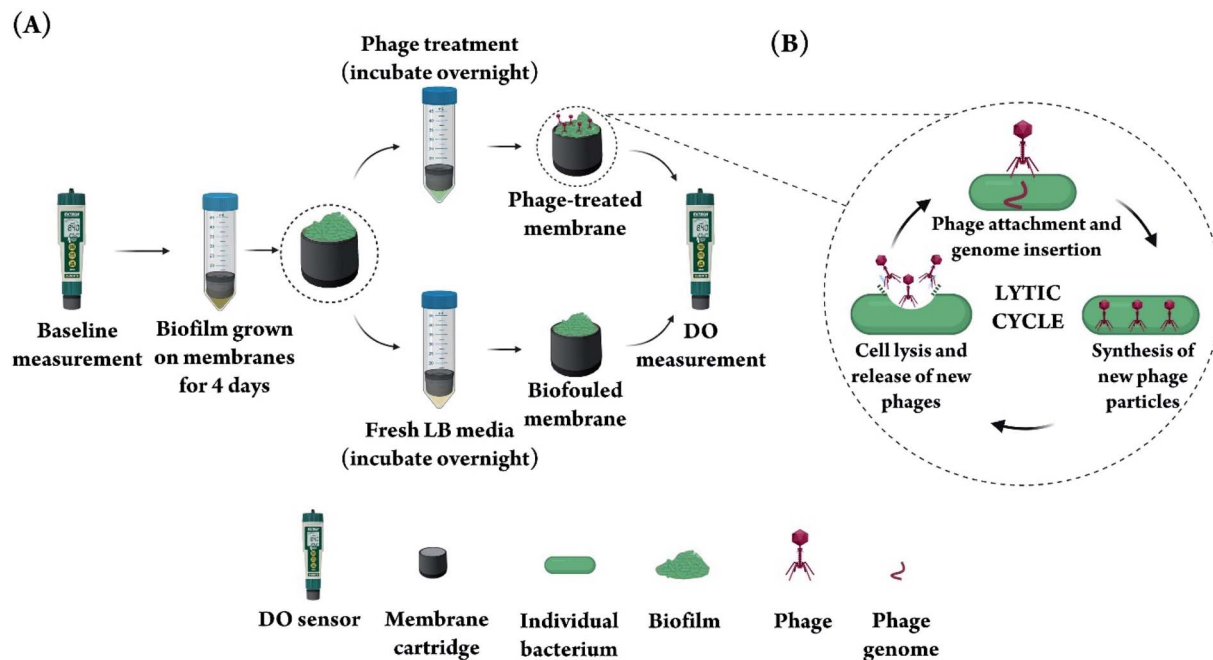


Fig. 1 (A) Schematic diagram showing the workflow for artificially fouling the membranes of DO sensors and subsequent phage treatment. After baseline measurement with the DO sensor, the membranes were incubated with a PA culture for 4 days to form a thick bacterial biofilm, then treated with phage overnight before subsequent DO measurements. (B) A simplified diagram of the phage lytic cycle that leads to lysis and destruction of the bacterial cells comprising the biofilm on the sensor membrane. First, phage attaches to specific receptors on the bacteria cell surface and injects its genome into the bacterium, thus taking over the cell reproduction machinery. Hundreds of phage particles are synthesized and assembled in each bacterium and the bacterium is subsequently lysed to release the new phages into the environment to infect more bacterial cells.

used lead to a significantly higher biofilm reduction (in a microtiter biofilm model) when used in combination, compared to the case when each phage is used alone (Fig. S1†). Fig. 1B shows a simplified schematic of the phage lytic cycle that leads to the destruction of bacterial cells in a biofilm as a result of phage action. Control biofilms were incubated overnight in fresh LB media without any phage.

Biofilm removal with phage

To ensure a thick biofilm forms on the sensor membranes, we first assessed the appropriate length of time needed to allow for biofilm formation on the membrane by quantifying biofilm growth in a 96 well plate for the duration of four days. The biofilm level was quantified using crystal violet, a commonly used dye that binds and stains both bacteria and the surrounding matrix.⁴² As shown in Fig. 2A, the level of total biomass steadily increased over the four day period. The SEM images of PA biofilm grown on DO sensor membranes in Fig. 2B confirm the trend shown in Fig. 2A and correlate with the well-documented stages of a PA biofilm lifecycle.^{46–48} As depicted in Fig. 2C, the biofilm developmental process begins with (i) attachment of PA cells to a surface. This is followed by (ii) the cells spreading across the surface instead of upward, as can be seen in our 1 day and 2 day biofilm SEM images by visibility of the membrane underneath the cells, and the beginning of EPS production. A consistent increase in cell clustering and matrix production can be seen between day 1–3 images as the biofilm

matures. The next stage of growth (iii) is the development of water channel structures and an increase in biofilm height, exhibited in the 3 day and 4 day SEM images where the membrane surface is not visible anymore (Fig. 2B). The 4 day biofilm SEM image shows full maturation (this is stage iv in Fig. 2C) as evidenced by encasement of the cells in EPS, which appears as the glue-like substance between the bacterial cells (Fig. 2B). Mature biofilms are very difficult to eradicate, even with harsh chemical and/or mechanical treatments.^{49,50} Additionally, clusters from a mature biofilm commonly detach,⁵¹ allowing the bacterial cells to spread and establish biofilms elsewhere (this is stage v in Fig. 2C). Based on these results, we chose 4 days of incubation as the time point that would result in a strong, well-developed biofilm and a thick matrix for all subsequent experiments, in an effort to properly demonstrate the biofilm-fighting ability of our phage treatment. We have used a microtiter plate model to quantify PA biofilm formation over time using a well-accepted and simple biomass quantification method, namely the crystal violet method. This method stains all biomass and will thus provide an overall value for the amount of bacteria plus the extracellular matrix at any given time. The crystal violet method, however, could not be applied directly to the DO sensor membranes because of non-specific membrane staining. Therefore, we quantified the biofilm using a microtiter plate model to demonstrate the increase in biomass over time, and used electron microscopy of biofilms grown on the DO sensor membranes to qualitatively confirm the quantitative data obtained using crystal violet method.



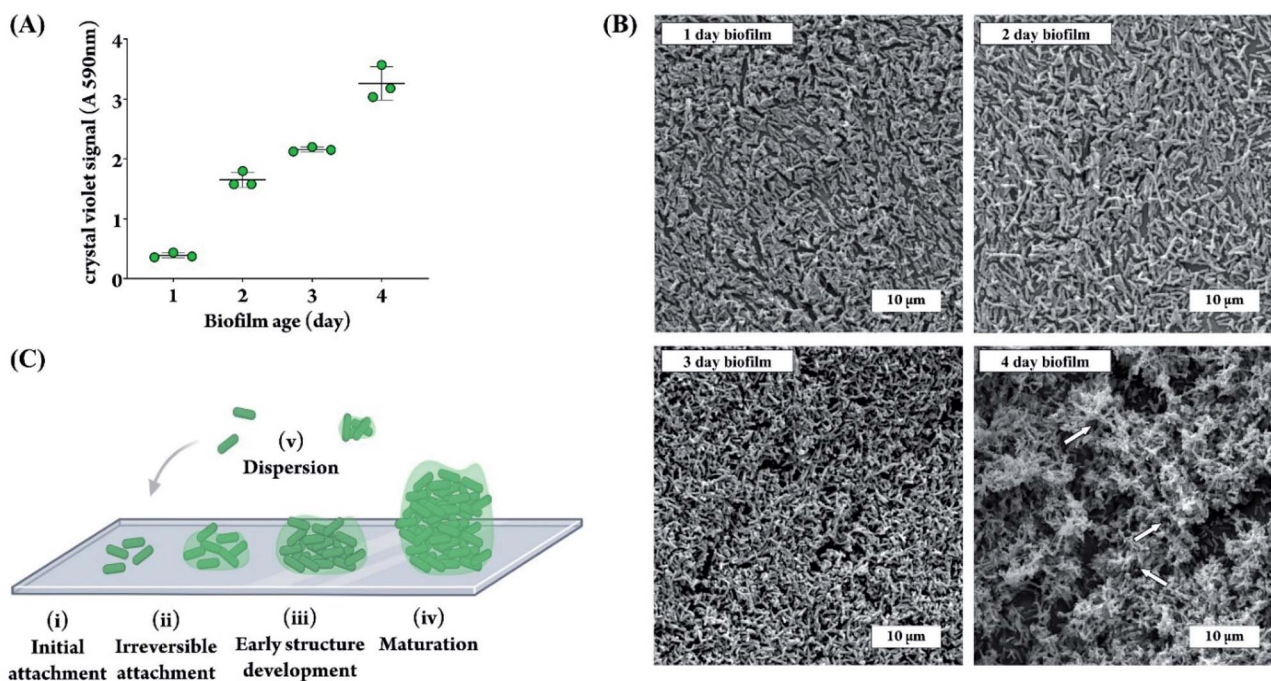


Fig. 2 (A) Amount of PA biofilm grown in a microtiter plate for the duration of one to four days, quantified as absorbance of re-solubilized crystal violet stain at 590 nm. Each data point represents the average of 20 technical replicates ($n = 3$). (B) SEM images of PA biofilm grown on DO sensor membranes for one, two, three, and four days. White arrows indicate some of the water channels visible in the matrix structure. (C) Schematic of a PA biofilm lifecycle: (i) initial attachment of cells to the surface; (ii) lateral spreading of cells and production of EPS, leading to irreversible attachment; (iii) early development of biofilm structure; (iv) mature biofilm architecture; (v) dispersion of cells from the biofilm and continuation of the cycle.

Next, we determined the required length of time for phage treatment to reduce biofilm levels. We chose 6 h and overnight (16–18 h) as these reflect timepoints commonly found to be effective in our lab for various phage-host pairs. The amount of biofilm remaining on the sensor membranes was quantified before and after phage treatment by staining the biofilm with Hoechst, a deoxyribonucleic acid (DNA)-binding fluorescent dye that can penetrate the bacterial cell wall and stain the bacterial genome.⁵² As shown in Fig. 3A, a 6 h phage treatment was capable of diminishing the fluorescence signal from stained bacteria, but an overnight treatment was the most effective at reducing the amount of biofilm on the DO membrane.

The fluorescence images from treated and non-treated membranes reflect the quantitative measurements, with an overnight phage treatment resulting in the least amount of staining on the membrane (Fig. 3B). It should be noted that the biofilm EPS is composed of exopolysaccharides, proteins, and extracellular DNA (eDNA),⁴ and therefore the DNA-binding fluorescent dye can bind to both the eDNA and the DNA inside the bacterial cells. To confirm the trends observed with fluorescence imaging of the membranes, SEM was used to visualize a biofouled membrane following phage treatment. The SEM images show that the phages were able to clear the dense matrix structure and no intact bacterial cells were visible (Fig. 3C). It is noteworthy that we chose a phage concentration of 10^9 PFU mL⁻¹ mainly to speed up the biofilm removal process. With these particular phages, we have observed in our lab that lower phage concentrations are also effective (10^6 – 10^8

PFU mL⁻¹), but it takes a longer time to achieve similar levels of biofilm dispersal with lower concentration. Should the reader choose to adapt this method to *in situ* treatment, we recommend that the phage treatment be applied by submerging the sensor in a closed container with phage, regardless of the phage concentration used because the dispersal and thus reduction of phage concentration will prolong treatment. As shown in Fig. 1B, lytic phages self-propagate by lysing bacterial cells and propagating themselves in the process, on site, which results in biofilm reduction over time. There are, however, traces of what appears to be cellular debris on the phage-treated membrane, which is expected to be a result of phage-mediated bacterial lysis. The full removal of the biofilm matrix is an interesting observation because it suggests the involvement of matrix-degrading enzymes. Phages coevolved with bacteria over millions of years and have developed specific strategies to combat biofilms. Some phages carry or express enzymes to degrade extracellular proteins and polysaccharides, key components of a biofilm matrix, and reach the bacterial cells embedded inside.⁵³ Alternatively, some phages can induce such enzymes from within the bacterial genome.⁵³ We would like to highlight the fact that genomes for the two selected phages have been sequenced and reported in the literature (NCBI: txid1141525 and NCBI: txid1141526), but only a fraction of those genes have a known function, with many classified as hypothetical proteins of unknown function. This is a well-known gap in knowledge in the phage field. A more comprehensive bioinformatic analysis would have to be performed on



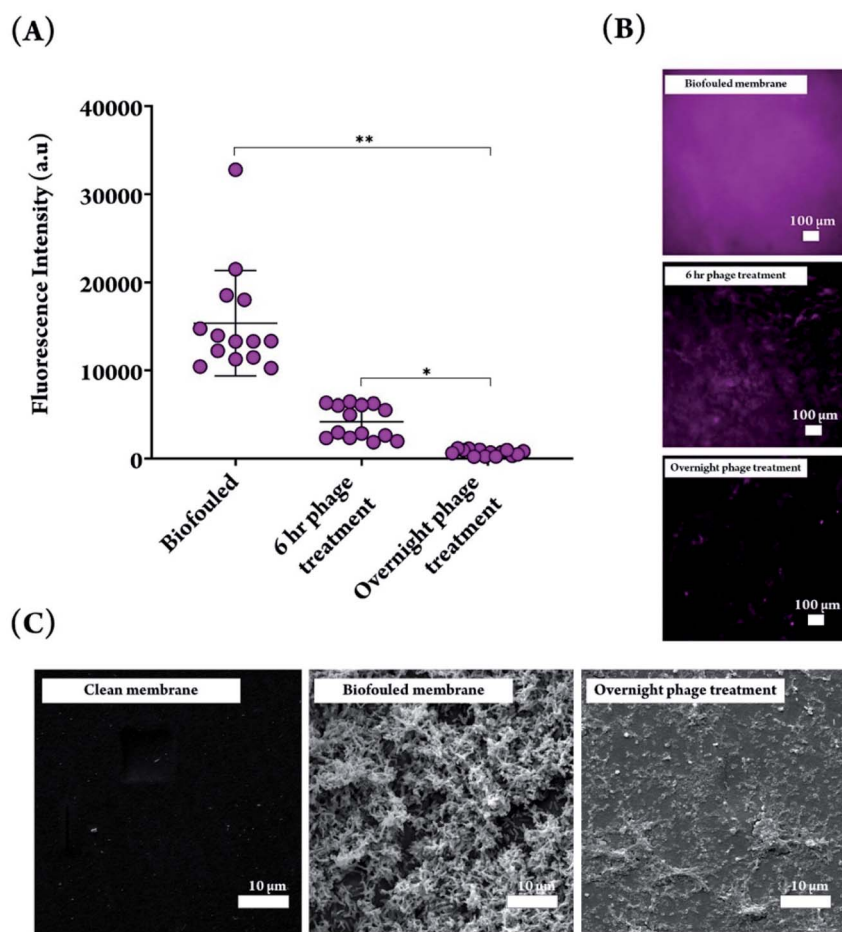


Fig. 3 (A) Fluorescence intensity of biofouled and phage treated membranes for 6 h and overnight phage treatments, stained with Hoechst dye and quantified with ImageJ software. Significant difference in fluorescence intensity was observed between biofouled and phage treated membranes. Overnight phage treatment resulted in a significantly lower fluorescence signal, compared to 6 h treated phage. Significant difference when comparing biofouled membranes with phage treated membranes, and overnight versus 6 h treatment (* $P < 0.05$, ** $P < 0.001$). (B) Fluorescence micrographs of biofouled, 6 h- and overnight phage treated membranes. The fluorescence signal does not differentiate between intracellular and extracellular DNA. (C) SEM images depicting a clean DO sensor membrane, a biofouled membrane with a 4 day PA biofilm, and a biofouled membrane subjected to an overnight phage treatment.

the phage genomes to uncover putative enzymes involved with biofilm degradation. We cannot, therefore, comment on the presence of genes encoding for matrix-degrading enzymes. However, determining appropriate phages for treating environmental biofilms does not necessarily require identification of anti-biofilm genes, but can be done with simple high-throughput assays which assess biofilm reduction in response to phage addition, which we have performed in our lab using our in-house phage library to choose the appropriate phage for this work.

Sensor signal recovery

As is evident from the fluorescence microscopy images of phage treated samples, overnight phage treatment led to a significant decrease in the amount of biomass, and successfully removed biofilms from the surface of DO sensor membranes. As a result, these membranes were used for further analysis to measure DO concentrations in DI water, and the resulting sensor signals

were compared with clean and biofouled membranes. The sensor used in this study is an electrochemical sensor that contains an electrode, electrolyte and a selectively permeable membrane. This polytetrafluoroethylene (PTFE)-based membrane allows dissolved oxygen in water to pass through and hinders the diffusion of water and other large components, thus playing a central role in the generation of an accurate electrochemical DO signal.⁵⁴ Once passed through the membrane, diffused oxygen undergoes an electrochemical reaction within the electrolyte solution, producing an electrical signal that is shown in the sensor display.

The DO sensor signal decreased from $7.96 \pm 0.27 \text{ mg L}^{-1}$ for clean membranes to $5.39 \pm 1.63 \text{ mg L}^{-1}$ after growing biofilms for four days on those membranes (32% decrease in signal after biofouling). Biofilm formation prevents proper oxygen diffusion through the small pores on the surface of permeable membranes, therefore significantly decreasing the signal and increasing the noise (note the increase in standard deviation). After incubating the biofouled membranes in the phage



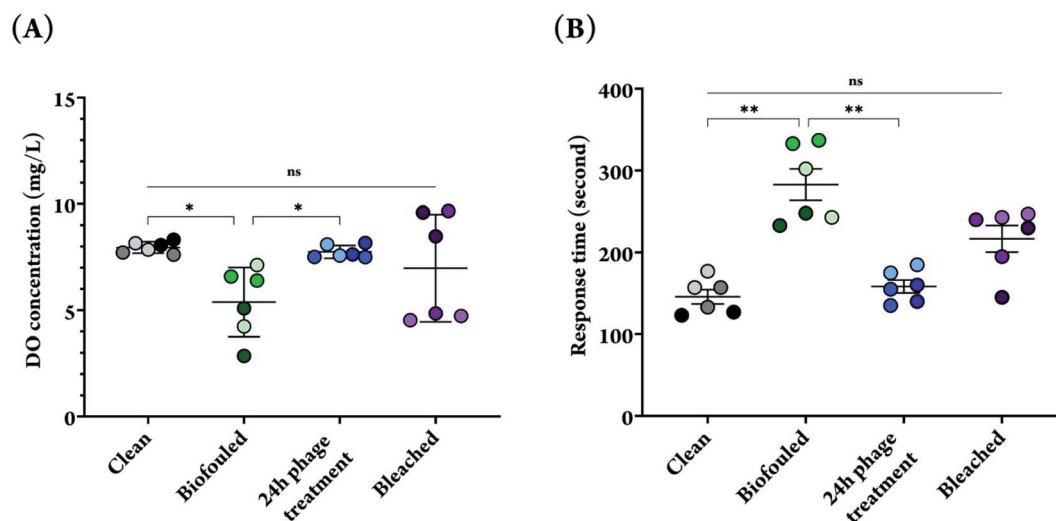


Fig. 4 (A) DO measurement for clean, biofouled, phage treated, and bleached sensor membranes in DI water. (B) Response time of DO sensor measured with clean, biofouled, phage treated and bleached membranes. The data points show the average of three measurements with each membrane. Experiments were performed in triplicates, and for each replicate, 2 membranes were used, bringing the total number of membranes used for each condition to six. Different color shades have been used to differentiate the data points for each biological replicate. Significant difference when comparing phage treated membranes with biofouled and clean membranes (* $P < 0.05$, ** $P < 0.01$).

suspension overnight, which allowed for biofilm removal from the surface of the membranes, the signal to noise ratio increased significantly (8 fold increase) and allowed for recovery of the absolute DO sensor measurement ($7.75 \pm 0.30 \text{ mg L}^{-1}$, Fig. 4A). Although some cellular and matrix debris still remained on the surface, results from DO measurements indicate that this did not affect the measurements, and the drastic decrease in amount of biomass following phage treatment worked well to enhance oxygen diffusion through the permeable membrane. In addition, the sensor response time that had increased following the biofilm formation on the membrane was also recovered after phage treatment, decreasing from $282.7 \pm 47.12 \text{ s}$ for a biofouled membrane to $158.3 \pm 19.41 \text{ s}$ for a phage treated membrane, which was comparable to the response time of a clean sensor $145.7 \pm 21.27 \text{ s}$ (Fig. 4B).

On the other hand, when a 10% bleach solution, a common bactericidal chemical, was used to remove biofilms from the surface of DO sensor membranes, a high variability in DO measurement and response time were observed. Although the results do not show significant difference from phage treated membranes, it is not a hidden fact that using chemicals such as bleach have adverse consequences including but not limited to corrosion of metals and painted surfaces, reaction with other chemicals and release of toxic gases and products, which can be extremely harmful to both humans and the environment.⁵⁵

Dissolved oxygen is one of the pivotal parameters defining the health of water bodies and marine environments. Aquatic creatures rely on dissolved oxygen and they require at least 5 mg L^{-1} to survive.⁵⁴ Electrochemical DO sensors, such as the one used for our investigation, are known for accuracy and fast response time, both of which have been significantly affected by biofouling of the sensor membrane, as indicated by our data, clearly demonstrating how biofouling could pose a major

challenge for the use of DO sensors in marine environment monitoring. A single phage treatment, however, regenerated the sensor surface and completely recovered the signal to its original value, as well as lowering the response time to that of a clean sensor. This demonstrates the utility of phage for non-invasive cleaning of DO sensors and suggests potential for cleaning of other sensor surfaces that have been subjected to biofouling.

Conclusion

We present an effective method for cleaning biofouled sensors and completely restoring the sensor signal to its original value with no harsh chemicals or mechanical treatments required, which is beneficial for use on delicate membranes and sensitive sensing surfaces. We achieved these results with only a single phage treatment, and without the need for repeated treatments, owing to the self-propagating nature of bacteriophages.⁵⁶ Phages can replicate on the sensor membrane, as long as they have access to viable host bacteria, allowing for a continuous cycle of infection until the biofilm is cleared and resulting in recovery of the DO sensor function. Aside from being polymicrobial, which would require the use of multiple phages, environmental biofilms typically develop a strong matrix, which would require the use of phage with matrix degrading enzymes. This means that biofilm control using bacteriophages requires a thorough understanding of the candidate bacteriophages. Not all bacteriophages perform equally for biofilm removal and this is an area of active research in our lab and others.

The specificity of phages in being able to only infect a few closely related strains of bacteria means they are non-destructive towards beneficial microbes present in the environment in which the sensor is installed,⁵⁷ unlike harsh



cleaning agents. The lack of environmental toxicity means phage treatments could, in principle, be readily applied on site, without the need to remove the sensor from the point at which it was installed. In that case, the phage treatment can be applied by submerging the sensor in a closed container with phage.

It is worth noting that phage-host specificity would require the real-life application of our proposed method to involve a pre-screening for specific lytic phages that can infect the bacteria comprising the biofilm. In this study, we grew mono-species biofilms using a lab strain of bacteria known to be a strong biofilm former as a demonstration of the effectiveness of phage treatment against sensor biofouling. Environmental biofilms are typically a mixture of multiple species,⁵⁸ which would require the use of more than one type of phage. Finding phages that infect the multispecies environmental biofilm is usually as easy as screening the same aquatic environment for phage, because phages reside in the same niche as their host. In fact, even for therapeutic use in humans, phages are routinely isolated from environmental reservoirs such as ponds, lakes, streams, or even sewage.¹⁹ This means that with seasonal changes in the composition of environmental biofilms formed in aquatic environments, we have access to an endless supply of phages that change and evolve with changes in the host population. With time, this would lead to a library of phages that can be used for a certain niche to tackle environmental biofilms on sensors and can be easily screened against new target bacteria. Generating phage preparations for environmental applications is relatively low cost because, unlike therapeutic applications, extensive purification may not be required.

In summary, with the correct selection of phage, bacteriophage treatment could find broad utility in regenerating not only DO or membrane-based sensors, but any sensor surface that is subject to extensive biofouling.

Abbreviations

DO	Dissolved oxygen
EPS	Extracellular polymeric substance
PA	<i>Pseudomonas aeruginosa</i> PAO1
DNA	Deoxyribonucleic acid
eDNA	Extracellular DNA
PBS	Phosphate buffered saline
DI water	Deionized water
PTFE	Polytetrafluoroethylene
SEM	Scanning electron microscopy
OD	Optical density
PFU mL ⁻¹	Number of plaque forming units per mL

Author contributions

The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript. DM and FB contributed equally to the manuscript.

Funding sources

ZH and TFD each acknowledge support from NSERC Discovery Grant and McMaster start up funds.

Conflicts of interest

The authors declare that there are no conflicts of interest.

Acknowledgements

Electron microscopy was performed at the Canadian Centre for Electron Microscopy (also supported by NSERC and other government agencies). All illustrations were created with Bio-render (<https://BioRender.com>).

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