Short-term Inactivation Rates of Selected Gram-Positive and Gram-Negative Bacteria Attached to Metal Oxide Mineral Surfaces: Role of Solution and Surface Chemistry

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Abstract

Metal oxides such as ferric or aluminum oxides can play an important role in the retention of bacteria in granular aquatic environments; however, their role in bacterial inactivation is not well understood. Herein, we examined the role of water chemistry and surface chemistry on the short-term inactivation rates of three bacteria when adhered to surfaces. To evaluate the role of water chemistry on the inactivation of attached bacteria, the loss in membrane integrity of bacteria attached to an iron oxide (Fe₂O₃) surface was measured over a range of water ionic strengths of either monovalent or divalent salts in the absence of a growth substrate. The influence of surface chemistry on the inactivation of attached bacteria was examined by measuring the loss in membrane integrity of cells attached to three surfaces (SiO₂, Fe₂O₃ and Al₂O₃) at a specific water chemistry (10 mM KCl). Bacteria were allowed to attach onto the SiO₂ or metal oxide coated slides mounted in a parallel-plate flow cell, and their inactivation rate (loss in membrane integrity) was measured directly without removing the cells from the surface and without disturbing the system. X-ray photoelectron spectroscopy analysis revealed a high correlation between the amounts of C-metal or O-metal bonds and the corresponding bacterial inactivation rates for each surface. Finally, for all three surfaces, a consistent increase in inactivation rate was observed with the type of bacterium in the order: *Enterococcus faecalis*, *Escherichia coli* O157:H7, and *Escherichia coli* D21f2.
Introduction

The occurrence of microbial pathogens in drinking water sources is recognized as an important threat to public health. A better understanding of the key processes governing the fate of microbial pathogens in groundwater aquifers or deep-bed granular filters used in water treatment operations can help mitigate the risk of drinking water contamination. The attachment of pathogens to aquifer or filter grain surfaces and the inactivation of attached and suspended pathogens are key processes that attenuate the concentration of viable pathogens in potable water supplies.\(^1\), \(^2\) Although substantial research effort has been aimed at elucidating the role of various physical, chemical and biological factors on the attachment of microorganisms to aquifer or filter grain surfaces,\(^3\), \(^4\) less is known about the factors influencing microbe inactivation in these environments. The impact of various environmental factors such as water chemistry, temperature, and organic matter content on the inactivation rate of viruses, bacteria, and/or protozoa suspended in the aqueous phase has been examined.\(^2\), \(^5\), \(^6\) However, our understanding of microbe inactivation when attached to grain surfaces is limited.

The surfaces of aquifer sediments or filter grains are commonly coated with patches of minerals such as ferric and aluminum oxides that may be positively charged at certain pH ranges (e.g., the pH\(_{\text{pzc}}\) for Al\(_2\)O\(_3\) is between 8 to 10).\(^7\)–\(^9\) A number of researchers have demonstrated that such metal oxide coatings can provide a favorable surface for the deposition of negatively charged colloids or microorganisms and thus play an important role in their retention in granular porous matrices.\(^10\)–\(^14\) It has also been shown that polyvalent cations increase attachment of bacteria\(^15\) and that the adsorption of dissolved organic matter onto iron oxides reduces adhesion by masking positive
charge. Although it has been established that microbes are preferentially retained on the surface of positively charged metal oxide patches under environmentally relevant conditions, the impact of these coatings and their surface chemistry on the inactivation of retained microorganisms has not been well examined. There is limited literature on techniques to measure the loss of viability of microbes attached to natural or model surfaces in an aqueous environment. Most of the methods rely on detachment of microbes from the surface, which may damage or kill cells and thereby introduce experimental error into the measurement. To overcome this and other experimental challenges in evaluating the role of aquifer or filter grain surface chemistry on microbial inactivation, we recently developed an experimental technique for in situ real-time evaluation of the inactivation kinetics of bacteria adhered onto a surface in model environmental conditions.

Poortinga et al. measured electron transfer during bacterial adhesion to indium tin oxide and titanium oxynitride coatings suggesting that chemical reactions may occur during adhesion. In another study, Kerisit et al. investigated the rate of direct electron transfer across the cytochrome-iron oxide interface. Thus, reactions between bacterial surfaces and mineral coatings on grain surfaces could also influence the attachment and subsequent inactivation of those attached microbes.

The objective of this study is to evaluate the extent of bacterial attachment and subsequent inactivation on model grain surfaces, representative of both bare silica and metal oxide portions of aquifer or filter grains. The bare sand surface is modeled using a clean glass (SiO₂) slide whereas the metal oxide coated patches found on the surface of aquifer or filter grains are modeled using glass slides coated with thin films of iron oxide (Fe₂O₃) or alumina (Al₂O₃). The bare or coated glass slides are mounted in a parallel plate flow chamber (PPFC), and the extent of bacterial attachment and
subsequent inactivation (loss in membrane integrity) of three different bacterial strains (*E. coli* D21f2, *E. coli* O157:H7, and *E. faecalis*) is examined in real-time without disturbing the experimental system. To evaluate the role of water chemistry on bacterial inactivation, the loss in membrane integrity of bacteria attached to an iron oxide (Fe$_2$O$_3$) surface was measured over a range of solution ionic strength (IS) (1 to 100 mM) of either monovalent or divalent salts (KCl and CaCl$_2$). The influence of the substratum surface chemistry on bacterial inactivation kinetics was examined by measuring the loss in membrane integrity of cells attached to three different surfaces (SiO$_2$, Fe$_2$O$_3$ and Al$_2$O$_3$) at a specific water chemistry (10 mM KCl). Finally, using high resolution X-ray photoelectron spectroscopy (XPS) analysis, the abundance of specific bonds between the bacterial cells and the three mineral surfaces was determined in an effort to assess the role of chemical bonding in the bacterial inactivation mechanism.

**Materials and Methods**

**Selection of Bacteria.**

Two Gram negative bacteria, non-pathogenic *E. coli* D21f2 and pathogenic *E. coli* O157:H7 (ATCC 700927), and a Gram positive organism, pathogenic *E. faecalis* (ATCC 29212) were used as the test bacteria. *E. coli* O157:H7 was selected as a representative bacterial pathogen that may be found in contaminated drinking water supplies. *E. coli* D21f2 was selected as it is a well-characterized non-pathogenic laboratory strain of *E. coli* used in several studies of environmental fate. It is also of interest to evaluate whether the non-pathogenic *E. coli* D21f2 could be a good surrogate for the inactivation behavior of *E. coli* O157:H7. To evaluate how the bacterial cell wall
structure may influence inactivation of cells attached to the metal oxide surfaces, *E. faecalis* was selected as a representative pathogenic Gram positive bacterium.

**Physico-chemical Characterization of Bacterial Cells.**

The electrophoretic mobility (EPM), hydrodynamic diameter and relative hydrophobicity of the selected bacteria were characterized (details are provided in the Supporting Information).

**Characterization of Silica and Metal Oxide Surfaces.**

A glass microscope slide (clear and corrosion-resistant silica, Fisher Scientific) was used as the bare silica surface. The glass samples coated with metal oxide films were prepared by the MicroFabrication Facility at McGill University. The 7 and 200 nm-thick films of Fe₂O₃ and Al₂O₃ were produced by low pressure chemical vapor deposition (Tylan/Tytan furnace). The thickness of the metal oxide films on the surface was determined using ellipsometry. The sputtering targets with dimensions of 3 in (diameter) × 0.125 in (thickness) were 99.99% pure Fe₂O₃ and Al₂O₃ (Kurt J. Lesker Company).

The relative hydrophobicity of the three types of surfaces was evaluated by water contact angle measurements at room temperature using an OCA-30 goniometer (Future Digital Scientific Corp., NJ, USA) equipped with a microliter syringe. Measurements were made on sessile drops (5 μL droplets) by measuring the tangent to the drop at its intersection with the surface. Values reported are the mean of the tangents taken at both sides of five droplets. The contact angle was measured from a snapshot taken 3 sec after the initial contact. All measurements were repeated on at least three different samples.
The streaming potential of bare and coated glass slides was measured using an electrokinetic analyzer fitted with an asymmetric clamping cell (Anton Paar, Graz, Austria). The measurements were performed in triplicate at room temperature in 1, 10, and 100 mM KCl or CaCl₂, pH 5.8. Streaming potential measurements were converted to zeta-potentials using the modified Smoluchowski-Helmholtz formulation.

The nature of the surface coating was also examined using XPS (VG ESCALAB 3 Mark II). The excitation source was non-monochromatic magnesium X-rays with a wavelength of 0.989 nm which corresponds to photon energy of 1253 for silica and alumina surfaces and non-monochromatic aluminum X-rays with a wavelength of 0.83386 nm which corresponds to photon energy of 1486.7 eV for the Fe₂O₃ surface. The binding energies for elements of interest were referenced to the binding energy of C 1s at 285.0 eV.

**Bacterial Adhesion Experiments.**

Bacterial adhesion to all three surfaces was examined using a PPFC with dimensions of 47.5 mm×12.7 mm×1.6 mm (Model FC81, Biosurface Technologies). As described previously, the PPFC was first rinsed with ethanol and DI water using a syringe pump (Model 200, KD Scientific) at 2 mL/min for duration of 10 min and then equilibrated with sterile electrolyte, either KCl or CaCl₂ (1, 10 or 100 mM IS), for 10 min, taking care to remove all air bubbles from the system. Subsequently, bacterial suspensions (3×10⁶ cells/mL) were injected into the flow cell (0.5 mL/min, 10 min), followed by an injection of only KCl or CaCl₂ of equivalent concentration (0.5 mL/min, 10 min) to remove any unattached cells from the chamber. Adhered bacteria were then stained using Live/Dead BacLight viability kit, imaged by fluorescence microscopy (IX-71, Olympus), and
enumerated with image analysis software (ImagePro). Images were acquired from the bottom slide
and mid-point length of the PPFC; on the surface of the bare or coated glass slide on an hourly basis
and for up to 6 hours. The top plate of the chamber was a clean microscope cover slide. The PPFC
was always maintained at room temperature (25°C). The images were then analyzed to calculate the
number of viable and total adhered bacteria as a function of time.

All experimental variables except the bacterial strain, electrolyte composition or surface type
were held constant, including cell concentration, exposure time, chamber size and solution volume.
An extent of attachment (EA) was quantified by normalizing the number of adhered bacteria onto
the desired surface immediately following a 10-min bacteria injection (N_{PPFC}) by the corresponding
number determined under favorable conditions for deposition (N_{PPFC,fav}) which represents the
maximum possible number of adhered bacteria:

\[ EA = \frac{N_{PPFC}}{N_{PPFC,fav}} \]  \hspace{1cm} (1)

Favourable, nonrepulsive conditions were achieved in the PPFC system by using 3-
aminopropyltriethoxysilane (APTES)-coated glass slides having a net positive zeta potential (+16±2
mV).\textsuperscript{18} Briefly, PPFC experiments were conducted using an APTES-coated glass slide and the
number of attached bacteria (N_{PPFC,fav}) was enumerated following a 10-min injection of cells followed
by a 10-min flushing of electrolyte (using the three bacteria described above, separately).

**Determiniation of Attached Bacterial Inactivation Rate.**

The viability of cells adhered to bare and coated glass surfaces was evaluated using a
commercially available kit (Live/Dead BacLight viability kit, Molecular Probes) which includes two
DNA-binding stains: the green fluorescent stain, SYTO-9, and the red fluorescent stain, propidium
iodide (PI). With this kit, determination of bacterial viability is based on differences in membrane integrity. SYTO-9 labels all bacteria, whereas PI penetrates bacteria with compromised membranes. Thus, bacteria with intact membranes fluoresce green and are considered to be viable while bacteria with compromised membranes fluoresce red.\textsuperscript{24} Attached bacteria were stained by filling the chamber of the PPFC with a solution of 100 \( \mu \)L BacLight stain in 5 mL electrolyte for 15 min. To prevent photobleaching, the entire experimental setup (PPFC + tubing) was protected from light and the image acquisition time was minimized. Stained bacteria were imaged using fluorescence microscopy. Direct counts were made for at least 20 images from the center of the PPFC at each time point. As a control, the inactivation of suspended (planktonic) bacteria was also measured over the same time period in the same solution conditions, and using the same BacLight solution in a 1:1 volumetric ratio. At each time point, 5 \( \mu \)L samples of stained planktonic cells (stored in glass vials at room temperature) were imaged by fluorescence microscopy for quantification of viable and nonviable suspended bacteria. Here, viability is defined in terms of preservation of membrane integrity as the percentage of bacterial cells with intact membranes (stained with SYTO-9) to the total number of cells. By evaluating the change in the number of viable (intact) cells as a function of time, the bacterial inactivation rate can be obtained. Specifically, the bacterial inactivation rate \((r)\) is calculated from the initial (first 3 h) slope of the viability curve as the \% loss of viability per hour.

To investigate the role of solution IS and cation valence on bacterial attachment and inactivation, the experiment was conducted with Fe\textsubscript{2}O\textsubscript{3}-coated surfaces at 1, 10, and 100 mM IS of KCl or CaCl\textsubscript{2}, for the three bacterial strains. The solution chemistries tested were selected to be representative of the ions and range of IS found in natural waters.\textsuperscript{25} To further study the role of
surface chemistry on bacterial adhesion and inactivation, the same experiment was repeated using two other substrates; namely, bare glass (SiO₂) and Al₂O₃-coated glass at a single IS (10 mM KCl). Each experiment was performed in triplicate.

**XPS Analysis for Surface Characterization and Identification of Chemical Bonding between Cells and Surfaces.**

The nature of the surface film before and after bacterial attachment was examined using XPS (VG ESCALAB 3 Mark II). The excitation source was non-monochromatic magnesium X-rays with a wavelength of 0.98 nm which corresponds to photon energy of 1253 eV for silica and alumina surfaces and non-monochromatic aluminum X-rays with a wavelength of 0.83 nm which corresponds to photon energy of 1486.7 eV for the Fe₂O₃ surface. Sample preparation involved placing 200 µL of a fresh bacterial suspension (in 10 mM KCl) on a 1 cm² sample of bare or coated-glass surface for 3 h which is the same length of time as that used to evaluate the initial inactivation rate. Next, the excess cell suspension was removed using a pipette. The surfaces were rinsed with 10 mM KCl to remove the unattached cells. As a control, the same surfaces with no exposure to bacteria were also prepared. The constant pass energy of the analyzer for high resolution scan was 20 eV and it was calibrated with reference to the binding energy of C1s at 285.0 eV. The spectra were recorded following the sequence C1s, O1s, N1s, Si2p, Fe2p, and Al2p.
Results and Discussion

Bacteria Properties.

Cell surface charge has been shown to play an important role in bacterial attachment to surfaces; hence, the cell surface (zeta) potential was evaluated from EPM measurements conducted over the range of experimental conditions investigated. The zeta potentials of the *E. coli* D21f2, *E. coli* O157:H7 and *E. faecalis* cells indicate that all three bacterial strains used in this study were negatively charged over the range of IS and at the pH (5.6-5.8) examined. The zeta potential results are presented in Figure S1. For *E. coli* D21f2 and *E. faecalis*, the absolute magnitude of the cell zeta potential decreased with an increase in IS of either KCl or CaCl$_2$, as expected based on compression of the electrical double layer. In contrast, for *E. coli* O157:H7, the cell zeta potential remained close to neutral over a range of IS in both electrolytes. The other bacteria exhibited a much more negative zeta potential when suspended in KCl versus CaCl$_2$ solutions. This is likely due to Ca$^{2+}$ electrical charge screening and complexation with bacteria surface proteins resulting in charge neutralization. The bacterium with the least negative zeta potential was *E. coli* O157:H7 in either KCl or CaCl$_2$ solution (~2 mV), while the most negatively charged bacterium was *E. coli* D21f2 in KCl solution (~25 mV in 1 mM KCl). These zeta potential values were used to calculate the DLVO interaction energy profiles between the bacteria and the iron oxide surface, which are discussed later in the paper.

The hydrodynamic diameter of all three bacteria measured by DLS (Table S1) and confirmed by scanning electron microscopy was approximately 1.3±0.17 μm. Also, it was observed that the size of the cells was not greatly affected by the presence of either K$^+$ or Ca$^{2+}$ (Table S1).
These data suggest that the variation in bacterial adhesion observed for the different bacteria in the presence of the monovalent versus divalent cations cannot be attributed to changes in cell size.

**Properties of Silica and Metal Oxide Surfaces.**

The relative hydrophobicity of the three surfaces was determined by water contact angle measurements. The bare glass had the most hydrophilic surface (contact angle: 8°, Table S2). A contact angle of 65° was measured for the Fe₂O₃ surface, and 60° for the alumina surface (Table S2). The bare glass and metal oxide surfaces were negatively charged in all conditions, except for alumina in 1 to 100 mM KCl and in 100 mM CaCl₂. The zeta potential values ranged from -40 to -6 mV for silica, -15 to -1.7 for Fe₂O₃, and -11 to +8, for alumina (Table S2). The thin film of Fe₂O₃ (7 nm) acquired negative charge in the range of tested IS which is in accordance with a previous study. In general, the surface charge should be more positive for a thicker metal oxide layer. The absolute magnitude of the surface zeta potential decreased with an increase in IS of either KCl or CaCl₂ for all three surfaces. These zeta potential values (Table S2) were used in the DLVO interaction energy calculations between the bacteria and the surfaces (Table 1).
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<th>IS (mM)</th>
<th>K⁺</th>
<th>Ca²⁺</th>
<th>K⁺</th>
<th>Ca²⁺</th>
<th>K⁺</th>
<th>Ca²⁺</th>
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<td>NB 0.5</td>
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<td>1.3 ± 0.6</td>
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<tr>
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<td>0.8</td>
<td>0.1 ± 0.1</td>
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</table>

*Hamaker constant of 3.22 × 10⁻²⁰ J was used to calculate interaction energies. \( k_B \): Boltzmann constant, 1.3805 × 10⁻²³ J/K. T: absolute temperature, \( k_B \): energy barrier. Standard deviations for \( r_i \) for planktonic cells and for \( E_A \) are not shown as they were found to be <0.05.

Table 1. Energy barriers \( (\Phi_{max}) \), extent of attachment \( (EA) \), and inactivation rates \( (r_i) \) of all three bacterial strains adhering onto Fe₂O₃ surface at various ionic strengths of KCl and CaCl₂. \( r_i \) values are also provided for planktonic (suspended) cells.

The presence of iron and aluminum elements on the surface of the coated slides was confirmed using XPS analysis. Table S3 shows the elemental analysis for silica, Fe₂O₃, and Al₂O₃ surfaces. The silica surface spectra contained two major peaks for O1s and Si2p at 532.82 eV and 103.23 eV, with relative quantities of 62.36% and 21.59%, respectively. For the Fe₂O₃ surface, the survey showed peak binding energies at 712.07 and 531.45 eV for Fe2p3 (corresponding to Fe(III) in Fe₂O₃) and O1s, respectively contributing to 13.4% and 60.4% of the surface. On the Al₂O₃ coated slide, the relative quantities of O1s and Al2p (in Al₂O₃) were 54.2% and 21.3% at 531.12 eV.
and 74.03 eV, respectively. Thus, the XPS analysis confirmed the presence of Fe₂O₃ and Al₂O₃ deposited on the silica surface.

Effect of Solution Chemistry on Bacterial Adhesion and Inactivation Rate.

To investigate the role of water chemistry on the extent of attachment and subsequent inactivation of attached bacteria, experiments were conducted over a range of IS (1 to 100 mM) of both monovalent and divalent salt solutions for a single surface (Fe₂O₃) (Table 1). As this study focuses on the inactivation of attached bacteria, it was of interest to first evaluate the extent of bacterial adhesion (EA). For all three bacteria, there was an increase in the EA with increasing IS of either KCl or CaCl₂. The ratio EA appears to reach maximum values of 0.5, 0.7, and 0.6 in KCl and approach 1, 0.9, and 0.8 in CaCl₂ for *E. coli* D21f2, *E. coli* O157:H7, and *E. faecalis* adhering onto the Fe₂O₃ surface, respectively. Among the tested bacteria, *E. coli* O157:H7 exhibits the highest adhesion in either KCl or CaCl₂ when depositing onto Fe₂O₃. In general, bacteria suspended in KCl adhered less as compared with cells suspended in CaCl₂ solution where the cells were less negatively charged (Figure S1) and more hydrophobic (Table S1). These variations in adhesion behavior may be attributed to cell electrokinetic properties, surface charge, cell hydrophobicity, and cell surface structures.²⁹,³⁰

To better understand the differences in EA measured in KCl and CaCl₂ solutions for all three bacteria, the classical DLVO theory³¹, ³² is considered and the bacteria-surface interaction energy calculations are presented as the height of the repulsive energy barrier (Φ_max) in Table 1 (details regarding the DLVO calculations are provided in Supporting Information). The DLVO
calculations (Table 1) reveal the presence of a repulsive energy barrier to deposition onto Fe₂O₃ under two experimental conditions only. For each bacterium, the predicted presence of an interaction energy barrier corresponded to the lowest values of $E_A$ observed.

Although previous studies aimed at understanding the fate of bacteria in aquatic environments have previously examined the role of water chemistry on adhesion,²⁸, ²⁹, ³³, ³⁴ those studies have not attempted to quantify the loss in bacterial viability with time; namely, the bacterial inactivation rate for cells remaining attached on the substratum. Here, viability is defined in terms of preservation of membrane integrity as the percentage of bacterial cells with intact membranes to the total number of attached cells. Bacteria were mostly viable (viability>90%) at the beginning of each experiment (when they first adhered onto the Fe₂O₃ surface) and there was a decrease in the observed viability over time (Figure S2). The initial rate of bacterial inactivation for each bacterial strain can be evaluated from the initial slopes of the curves presented in Figure S2 (i.e., from the linear portion of the curves, between $t=0$ and 3 h). ¹⁸ The bacterial inactivation rates are plotted in Figure 1 as the loss in viability with time ($n$) (% loss/h) of cells attached onto Fe₂O₃ for different solution IS of either monovalent or divalent cations. The data is also summarized in Table 1 for direct comparison with the values of $\Phi_{\text{max}}$ and $E_A$.

Figure 1 suggests that the effect of solution chemistry was more pronounced for $E. \text{coli}$ D21f2 and for $E. \text{coli}$ O157:H7, whereas for $E. \text{faecalis}$, there was no significant change in inactivation rate of the adhered cells. The pathogenic $E. \text{coli}$ O157:H7 displays a comparable ability to survive on Fe₂O₃ as the non-pathogenic $E. \text{coli}$ D21f2 in CaCl₂ solution (Figure 1b), where the inactivation rates were 2.1%/h and 2.6%/h in 1 mM CaCl₂ and 1.3%/h and 1.8%/h in 10 mM
CaCl$_2$ for *E. coli* D21f2 and *E. coli* O157:H7, respectively. However, the attached non-pathogenic *E. coli* D21f2 was more vulnerable in KCl solution (Figure 1a). Moreover, *E. faecalis* 29212, a representative Gram-positive bacterium, exhibited a lower inactivation rate in comparison to the two *E. coli* strains and was hence more resistant in the attached mode at the experimental conditions examined here. For comparison, the inactivation rates of suspended (planktonic) cells are also included in Table 1. These data show that bacterial viability was generally higher when the cells were suspended versus attached on Fe$_2$O$_3$ (for all solution chemistries).

**Figure 1.** Inactivation rates based on Live/Dead BacLight viability measurements for different solution ionic strengths (1, 10, and 100 mM) of (a) KCl and (b) CaCl$_2$ for *E. coli* D21f2, *E. coli* O157:H7, and *E. faecalis* adhered onto Fe$_2$O$_3$. The error bars indicate standard deviations.

In general, under all conditions, the three bacteria attached onto Fe$_2$O$_3$ exhibited the highest inactivation rate in the lowest IS solution. When decreasing the solution IS, the cell inactivation rate increased more significantly in KCl versus CaCl$_2$. High osmotic pressure and electrolyte imbalance across the cell membrane are proposed to be the causes of low cell viability at low IS.$^{35}$ However, high viability of planktonic cells over a period of 6 h under all conditions (Table 1) suggests that
other mechanisms are involved in bacterial inactivation when the cells are attached to the surface at low IS.

**Influence of Surface Chemistry on Bacterial Adhesion and Inactivation Rate.**

To better understand the role of substratum surface chemistry on the inactivation of attached cells, we examined the inactivation of bacteria attached on three different surfaces; namely, silica, Fe₂O₃, and Al₂O₃ at a specific solution chemistry (10 mM KCl, pH 5.8).

Figure 2a shows the EA for all three bacteria on the three surfaces at 10 mM KCl. For each organism, the lowest EA was observed on silica and the highest EA was measured on alumina which was positively charged (Figure 2a). On the silica surface, for all three strains, the initial inactivation rate was less than 2%/h (Figure 2b) and there was no significant change in viability over a 6 hour period (data not shown). For the pathogenic *E. coli*, the inactivation rates were comparable on the Fe₂O₃ (-2%/h) and Al₂O₃ surfaces (-3%/h) (Figure 2b). The non-pathogenic *E. coli* D21f2 exhibits the highest inactivation rate (-4%/h) on the alumina surface. For all three surfaces examined, the viability of attached bacteria was greater than 90% at the beginning of each experiment (when they first adhered) and there was a decrease in the observed viability over time, similar to the patterns shown in Figure S2. Of the two metal oxide surfaces, alumina exhibits a greater antimicrobial effect on adhered *E. coli* cells due to its net positive surface charge\(^{17, 36-40}\) compared to the negatively charged Fe₂O₃. Metal ions such as silver ions and metal oxides such as zinc oxide, copper oxide, and aluminum oxide have been shown to exhibit antimicrobial properties.\(^{41-43}\) For instance, reductions of more than 5 log₉ CFU/mL of both *Staphylococcus aureus* and *E. coli* were confirmed after 90 min of
treatment with a silver ion solution. In another study, 8.2% reduction in culturability was reported for *Pseudomonas chlororaphis* O6 after 60 min exposure to ZnO and CuO. In addition, a mild growth inhibitory effect of aluminum oxide nanoparticles were reported for *E. coli*. A 60% growth reduction was observed over a 24-hour period for *P. aeruginosa* in the presence of 250 μg/mL Al₂O₃ nanoparticles. In the same study, ZnO reduced the growth of *S. aureus* and *P. aeruginosa* by 62% and 49%, respectively. In these studies, the bacteria and the metal oxide particles were suspended in a liquid phase, whereas in our study, inactivation occurred after bacterial attachment to the substratum.

Terada et al. investigated the effect of substratum surface charge on *E. coli* adhesion and survival, and found that electrostatic attraction forces induced more bacterial retention and lower viability. In the present study, comparing the results of EA and n (Figure 2) shows that there was no correlation between the extent of attachment and inactivation rate for the tested bacterial strains when considering all three surfaces. For instance, *E. coli* O157:H7 exhibited higher attachment than *E. coli* D21F2 on Fe₂O₃ in 10 mM KCl but its inactivation rate was lower. In another example, *E. faecalis* exhibited comparable attachment to the other tested strains on Fe₂O₃, but it had the lowest inactivation rate.
Figure 2. (a) Extent of bacterial attachment (EA) and (b) inactivation rates (r) for E. coli D21f2, E. coli O157:H7, and E. faecalis adhered to different surfaces (silica, Fe₂O₃, and Al₂O₃) in 10 mM KCl. The error bars indicate standard deviation. An asterisk indicates statistically significant difference in inactivation rates when compared to the silica treatment (determined using one-way ANOVA, p < 0.005).

A potential mechanism that could play a role in bacterial inactivation is loss of membrane structural integrity as a result of interactions between bacteria and minerals. It has been shown that different constituents in cell walls of Gram-positive and Gram-negative bacteria are capable of binding metal ions because of exposed anionic charges associated with them. In this study, for each bacterium, different inactivation rates were observed on surfaces with different chemistry. The selected representative Gram-positive bacterium exhibited the highest resistance on all three surfaces. To determine how the nature of the chemical bonding between the cells and the surfaces may contribute to bacterial inactivation, XPS analysis was used to characterize the cell-surface bonding chemistry. Among different chemical bonds identified using high resolution XPS analysis, the chemical bonds of particular interest to this study were the ones between the oxygen or carbon elements of cell surface functional groups and the metals (Si, Fe, and Al) on the adhering surface (Figures 3 and S3). XPS is a surface chemical analysis technique which does not penetrate the entire cell depth. Because the X-ray cannot penetrate the entire cell depth, the angled X-ray beam is
actually probing the bare surface where there are no cells as well as the edges where bacteria meet the surface (i.e., the bacteria periphery). The analyzed area for each surface was 2 mm×3 mm. In Figure 3, the relative atomic abundance of O-metal or C-metal bonds (evaluated by XPS) was normalized by the number of adhered cells for each surface (this data is represented by the bar graphs). This was achieved by dividing the relative atomic abundance (%) of O-metal or C-metal bonds by the relative atomic abundance (%) of nitrogen (N1s) for each bacterial strain on each surface. As N is only present in cells and not on the adhering surfaces (according to XPS survey of bare surfaces, Table S3), this element provides a fair estimate of the amount of adhered cells. The XPS analysis reveals the presence of C-metal bonds on each type of surface (SiO$_2$, Fe$_2$O$_3$, and Al$_2$O$_3$). This type of bond is not present on the bare surfaces and, thus, the detected C-metal bonds result from bacterial attachment to the surfaces. The interpretation of the abundance of O-metal bonds is more challenging because XPS will also detect the O atoms bound to Al, Fe, or Si in the bare surfaces. Nonetheless, the data in Figure 3 do confirm that XPS detects the contribution of O-metal bonds from the bacterium-surface interface as the technique is sensitive enough to distinguish the different relative amounts of this bond when the three different bacteria are attached to a given surface (e.g., Figure 3d).

The normalized atomic abundance of C-metal and O-metal bonds and corresponding bacterial inactivation rates are plotted for all three bacterial strains adhered onto different surface chemistries (Figure 3). As clearly illustrated in Figure 3, the abundance of C-metal or O-metal covalent bonds was highly correlated with the corresponding inactivation rate of the adhering bacteria on a given surface. For instance, $R$-squared values were 0.997, 0.999, and 0.894 for linear
regressions between the normalized amounts of C-metal bonds and measured bacterial inactivation rates for silica, Fe₃O₅, and Al₂O₃ surfaces, respectively (Figure 3a-c). The same trend was observed for O-metal bonds for all three bacteria adhered onto the three different surfaces (Figure 3d-f).

The inactivation of bacteria following adhesion onto or interaction with different minerals has been the subject of selected recent studies. Cai et al. recently reported that goethite significantly reduces the viability of attached *E. coli* O157:H7. These researchers proposed that the stronger and closer association between positively-charged goethite and bacteria is responsible for the observed lower bacterial viability as compared to cells bound to negatively-charged montmorillonite and kaolinite. Additionally, they hypothesized that the needle-shaped goethite structure can pierce the cell membrane thereby causing cell death. Another study suggests that the antibacterial mechanism of natural clay (illite-smectite or kaolin) is caused by released Fe²⁺ which suppresses outer membrane regulatory proteins by oxidizing to Fe³⁺ and forming toxic hydroxyl radicals. In our study, we observed a direct link between the nature of the bacteria-surface chemical bonding and the loss in cell membrane integrity. These data suggest that bacteria were more likely to become inactivated when attached to a surface with a greater number of covalent bonds; namely, C-metal or O-metal bonds. Although we observe a correlation between the abundance of C- or O-metal bonds and the extent of bacterial inactivation, the XPS analysis does not prove a causative effect. Further studies are needed to better understand the link between the type and abundance of bacteria-surface bonds and the underlying mechanisms of bacterial inactivation.

The Gram-positive bacterium exhibited the lowest extent of inactivation on the three surfaces. It is interesting that this bacterium appeared to form fewer bonds with the metals on the
surface (Figure 3). It has been shown that the metal binding ability of Gram-positive *Bacillus subtilis* is influenced by the proton motive force. In metabolically active cells, there is competition between protons and metal ions for anionic wall sites, resulting in less metal bonds in live cells compared to non-living cells.
Figure 3. Comparison of the normalized atomic abundance (%) of (a-c) C-metal bonds and (d-f) O-metal bonds formed between (a, d) silica, (b, e) Fe₂O₃, and (c, f) Al₂O₃ surfaces and the adhered cells identified by high resolution XPS analysis with the corresponding inactivation rates. The error bars indicate standard deviation for $r_i$. The standard deviation for amount of C-metal and O-metal bonds was less than 0.005.

The experiments presented here were conducted using bare glass (to mimic a sand surface as it is mainly composed of silica) and metal oxide-coated glass (to mimic patches that may be present
on the surface of aquifer or filter grains). The effect of both solution and surface chemistry on the inactivation of adhered cells was investigated. The results show that solution chemistry plays an important role in bacterial adhesion and inactivation. Lower solution IS led to higher inactivation rates of attached bacteria and all the tested strains had higher inactivation rates in KCl compared to CaCl₂. We also observed some variability in the inactivation behavior of the three bacterial strains when adhered on a given surface. Such differences in inactivation rates can presumably be attributed to differences in the structure and composition of the external cell wall of the selected bacteria.¹⁷ Moreover, each bacterial strain exhibited a statistically significant difference in its inactivation rate on alumina (when compared to the silica surface) which can be linked to the net positive charge of the alumina. Other studies have also reported that positively charged surfaces exert an antimicrobial effect on adhering bacteria, while negatively charged surfaces have negligible effects on bacterial viability or metabolic activity.¹⁶,¹⁸,⁴⁶

There are a number of reported laboratory studies of the effects of surfaces on bacterial activity.³⁹,⁵¹,⁵² Attachment of bacteria to surfaces can lead to enhanced or reduced bioactivity depending on the surface properties and charge,⁵³-⁵⁶ and it has been also suggested that some bacteria attach to specific surfaces as a survival strategy,⁵⁷,⁵⁸ but there are also examples of studies where association with a surface had no effect on bacterial activity.³⁹,⁶⁰

In this study, we found that attachment of cells to these model solid surfaces can lead to formation of covalent bonds, specifically O- and C-metal bonds, between the cell and the surface, and this results in reduced bacterial viability (i.e., cell membrane integrity). This knowledge is vital
for understanding how cells interact with solid surfaces as well as how they become inactivated after adhering to surfaces under environmentally relevant conditions.

The results from this study show promise that it may be eventually possible to estimate the rates of inactivation of attached pathogens by accounting for the mineral surface chemistry of sediments, or, barrier materials to inactivate pathogens can be identified based on mineral surface chemistry.
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Supporting Information

Details regarding Live/Dead bacterial viability data and bacterial characterizations such as MATH assay, XPS analysis, as well as figures and tables showing supplementary data. This information is available free of charge via the Internet at http://pubs.acs.org/.
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TOC Graphic