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Factors affecting the Faradaic efficiency of Fe(0) electrocoagulation

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ABSTRACT

Electrocoagulation (EC) using Fe(0) electrodes is a low cost water treatment technology that relies on efficient production of Fe(II) from the electrolytic dissolution of Fe(0) electrodes (i.e. a high Faradaic efficiency). However, the (electro)chemical factors that favor Fe(0) oxidation rather than O\textsubscript{2} evolution during Fe(0) EC have not been identified. In this study, we combined electrochemical methods, electron microscopy and Fe measurements to systematically examine the interdependent effects of current density (i), anodic interface potential (E\textsubscript{a}) and solution chemistry on the Faradaic efficiency. We found that Fe(0) oxidation was favored (Faradaic efficiency > 0.85) in chloride and bromide solutions at all i, whereas carbonate, phosphate, citrate, and nitrate solutions lead to Faradaic efficiencies < 0.1. The anodic reaction (i.e. Fe(0) oxidation or O\textsubscript{2} evolution) only depended on i in the sulfate and formate solutions. Experiments in binary-anion solutions revealed that molar ratios of [HCO\textsubscript{3}\textsuperscript{-}] / [Cl\textsuperscript{-}] near 100 and [NO\textsubscript{3}\textsuperscript{-}] / [Cl\textsuperscript{-}] near 20 separated the electrochemical domains of Fe(0) oxidation and O\textsubscript{2} evolution in the EC system. These molar ratios were supported by experiments in synthetic groundwater solutions. We also found that the E\textsubscript{a} vs i curves for solutions with poor Faradaic efficiency overlapped but were situated 2–4 V vs Ag/AgCl higher than those of solutions with high Faradaic efficiency. Therefore, the position of the E\textsubscript{a} vs i curve, rather than the E\textsubscript{a} alone, can be used to determine unambiguously the reaction occurring on the Fe(0) anode during EC treatment.

1. Introduction

Electrocoagulation (EC) using Fe(0) electrodes is a promising, low cost treatment technology for drinking water and wastewater. This technology is particularly applicable in decentralized and rural areas because of its low infrastructure requirements, ease of use and potential for scalability [1,2]. While EC systems are often accompanied by supplementary treatment steps, including settling tanks and filters [3], the technology centers on applying an electric current to Fe(0) electrodes in contact with an electrolyte solution to promote the oxidative release of soluble Fe(II) ions [4,5]. The Fe(II) ions migrate from the Fe(0) anode surface to the bulk solution, where contaminant removal occurs by several possible pathways. For example, Fe(II) ions can immobilize target species through direct reduction reactions (e.g. chromate) [6–9] or they can be oxidized by dissolved oxygen and Fenton-type intermediates (e.g. O\textsubscript{2}^\textsuperscript{-}, H\textsubscript{2}O\textsubscript{2}, and Fe(IV) or ‘OH [10,11]) to produce reactive Fe(III) (oxyhydr)oxides that remove contaminants by sorption (e.g. arsenic [12,13], heavy metals [14], dyes [15,16], perfluoroalkyl acids [17], and pathogens [18]). In addition, the oxidized Fe(II) ions form solid flocs that can be used as a pre-treatment step for additional purification processes, such as membrane filtration or advanced oxidation [19,20]. In recent years, the molecular scale mechanisms of contaminant removal in the bulk solution during Fe(0) EC treatment have been the focus of considerable research [19,21–23]. However, few EC studies have examined the electron transfer reactions occurring at the Fe(0) anode [24,25], which are responsible for the production of Fe(II). Because Fe(0) EC treatment relies on the production of Fe(II) from the Fe(0) anode [26,27], it is essential to understand the factors that promote efficient Fe(II) generation.

In EC, the concentration of Fe(II) (mol/L) generated at the Fe(0) anode is related to the applied current by Faraday’s law:

\[
[\text{Fe}] = (\text{It})/(\pi F V) \tag{1}
\]

where \(I\) represents the current (C/s) applied for time, \(t\) (s), \(n\) represents the number of electrons transferred (\(n = 2\) for Fe(II) production, consistent with previous work [4]), \(F\), the Faraday constant (96,485 C/mol) and \(V\), the solution volume (L). From Eq. (1), we can define the Faradaic efficiency as the Fe concentration measured in the bulk solution,
normalized by the theoretical Fe(II) concentration calculated by Faraday’s law. Therefore, a Faradaic efficiency near 1 indicates the production of Fe(II) and optimum system performance. Conversely, a drop in Faradaic efficiency, and thus treatment efficiency, can occur when the applied current drives the anodic oxidation of H₂O or coexisting solutes, rather than Fe(0). As shown in Fig. 1a, the anodic oxidation of Fe(0) to form Fe(II) occurs at a low standard redox potential (E⁰), whereas the oxidation of H₂O, chloride, or bromide can be favored as the anodic interface potential increases above 1.0 V vs Ag/AgCl. Therefore, knowledge of the interface potential, which relates to the thermodynamic driving force of electrode reactions [28], is critical to deconvolute the factors that affect the Faradaic efficiency of Fe(0) EC.

Based on previous research, we can identify a number of (electro)chemical parameters that can potentially influence the Faradaic efficiency and interface potential [24,25,29]. For example, the Butler-Volmer relationship shows that the applied current is directly proportional to the interface potential:

\[ i = i_0 \left( \exp(a_A E_A) - \exp(a_C E_C) \right) \]

where \( i \) represents the current density (mA/cm²), \( A \), the electrode surface area (cm²), \( i_0 \), the exchange current density (mA/cm²), \( a_A \) and \( a_C \) represent constants (detailed in Supplementary material) and \( E_A \) and \( E_C \) represent the anode and cathode interface potentials (V). Therefore, an increase in \( i \) (or \( I \) for a constant electrode surface area) can increase the anodic interface potential (\( E_A \)) to levels high enough to yield O₂, Cl₂ or Br₂ (Fig. 1a), which would decrease the Faradaic efficiency. This relationship between \( i \) and the Faradaic efficiency in Al(0) and Fe(0) EC studies has been reported for a limited range of electrolyte solutions (e.g. nitrate [30] and sulfate [24,25,29]). However, the absence of \( E_A \) measurements in most Fe(0) EC investigations has prevented reliable explanations of the impact of \( i \) on the Faradaic efficiency.

Another factor likely to influence the Faradaic efficiency is the ionic composition of the solution. For example, Arroyo et al. (2010) measured a Faradaic efficiency of ~1.0 for Fe(0) EC experiments in a chloride solution [31], whereas a reduced Faradaic efficiency has been reported in sulfate and nitrate solution at circumneutral pH [25,30]. However, these previous Fe(0) EC studies investigated a limited range of \( i \) and did not examine the impact of a wide variety of electrolyte solutions. A number of other inorganic and organic ions that are ubiquitous in natural systems, including carbonate, phosphate, formate and citrate, have been shown to interact with Fe(0) surfaces to inhibit the oxidation of Fe(0) in passive corrosion systems [32,33]. The impact of these environmentally relevant ions on the Faradaic efficiency of Fe(0) EC is poorly constrained, leading to a critical knowledge gap. Because Fe(0) EC treatment for a wide range of influent compositions hinges on maintaining a high Faradaic efficiency and minimizing electrode passivation, it is essential to investigate systematically the relationship between \( i \), \( E_A \) and Faradaic efficiency in a diverse set of solution compositions.

In this work, our objective was to determine the (electro)chemical factors that govern the Faradaic efficiency of Fe(0) EC. To this end, we investigated the interdependent effects of \( i \), \( E_A \) and solution composition by combining time dependent measurements of \( E_A \) and total Fe released from the Fe(0) anode as a function of \( i \) and solution chemistry. To encompass the range of current densities likely applied in Fe(0) EC field treatment [3], the \( i \) of EC experiments was adjusted over two orders of magnitude (0.25–50 mA/cm²). The impact on the Faradaic efficiency of a variety of dissolved species common in natural waters was studied by systematically varying the ionic composition and pH of the solution, building in complexity from single and binary anion systems to synthetic groundwater matrices. Micrometer scale modifications on the surface of Fe(0) anodes used in the EC experiments were probed by scanning electron microscopy (SEM). Our approach allowed us to elucidate the impact of solution chemistry and EC operating parameters on the redox reactions occurring at the Fe(0) anode, which has not been investigated previously and is critical for predicting and optimizing the performance of Fe(0) EC treatment under a wide range of source water chemistry. The ultimate aim of this study was to identify simple and low cost strategies to improve Fe(0) EC treatment in waters having compositions that result in poor Faradaic efficiency.

2. Materials and methods

2.1. Preparation of the electrolyte solutions

All chemicals used to prepare the stock solutions were reagent grade and all glassware was acid washed and rinsed three times with 18 MΩ·cm Milli-Q deionized (DI) water prior to use. Stock solutions of sodium chloride (NaCl, 0.5 M), potassium bromide (KBr, 0.2 M), sodium sulfate (Na₂SO₄, 0.2 M), sodium bicarbonate (NaHCO₃, 0.5 M),
sodium phosphate (Na2HPO4, 0.2 M), sodium nitrate (NaNO3, 0.2 M), sodium formate (NaHCOO, 0.2 M) and sodium citrate (Na3C3H5O
tions. In these experiments, solution pH drifted to neutral during pre-
cumneutral pH, we also examined solutions with a range of pH from

during the reaction. In addition to experiments performed at cir-

The amount of acid/base added changed the anion concentration by

Autolab potentiostat (Utrecht, the Netherlands) operated in a three

2.2. Electrochemical experiments

After adding all components to the single or binary anion solutions,

As our measurements of total Fe do not discriminate between Fe

AgCl reference electrode and Fe(0) working and counter electrodes
(spaced 2 cm apart). The trace metal composition of the Fe(0) elec-
trodes used in our study was reported previously [45]. Before each ex-
periment, the Fe(0) electrodes were cleaned and polished with fine
grained sandpaper. Because a positive electrochemical potential was
applied to the working electrode in each of our experiments, we refer to
the Fe(0) working electrode as the anode and the Fe(0) counter elec-
trode as the cathode in this work. Experiments were initiated by ap-
plying a galvanostatic current to the electrochemical cell with the

electrolyte solution mixed by a magnetic stir bar. Experiments were

Table 1
Composition, conductivity and $E_a$ ($i = 1 \text{ mA/cm}^2$) for single anion electrolyte solutions.

| Electrolyte | Anion | Ionic Strength (M) | Conductivity $^a$ (dS/m) | $E_a$ at $i = 1 \text{ mA/cm}^2$ (V vs Ag/AgCl) |
|-------------|-------|-------------------|--------------------------|------------------------------------------|
| Chloride    | 10    | 0.010             | 0.79                     | −0.1                                     |
| Bromide     | 10    | 0.010             | 0.79                     | 0.0                                     |
| Sulfate     | 5     | 0.015             | 1.18                     | 0.1                                     |
| Carbonate   | 10    | 0.010             | 0.79                     | 1.6                                     |
| Phosphate   | 5     | 0.015             | 1.18                     | 1.3                                     |
| Nitrate     | 10    | 0.010             | 0.79                     | 1.4                                     |
| Formate     | 5     | 0.010             | 0.79                     | 1.5                                     |
| Citrate     | 5     | 0.030             | 2.36                     | 1.5                                     |

$^a$ Conductivity was determined using the Griffin-Jurinak relation [62].

Table 2
Compositions and conductivity of the synthetic groundwaters.

| Electrolyte | Ionic Strength (dS/m) | $E_a$ at $i = 1 \text{ mA/cm}^2$ (V vs Ag/AgCl) |
|-------------|-----------------------|------------------------------------------|
| Chloride    | 0.646                 | 0.068                                    |
| Bromide     | 0.153                 | 0.055                                    |
| Sulfate     | 0.341                 | 0.055                                    |

$^a$ SBGW also contained 1.1 mM H2SiO4. Sodium is not included in this table but is used to balance electroneutrality. The total bicarbonate is represented by [HCO3$^-$/]. The open circuit potential was $−0.2 \text{ V vs Ag/AgCl}$ for all solutions except NO3-GW 2, which had an open circuit potential of $−0.5 \text{ V vs Ag/AgCl}$. $^b$ Conductivity was determined using the Griffin-Jurinak relation [62].

2.2. Electrochemical experiments

Electrochemical experiments were performed with a Metrohm-
Autolab potentiostat (Utrecht, the Netherlands) operated in a three

electrode system (Fig. 1b). The electrochemical cell consisted of an Ag/
consistent with the electrochemical production of only Fe(II). Our results are plotted as the average and standard deviation, where applicable, of replicated experiments, but we note that the variation for many data points (experiments with Faradaic efficiency < 0.1 and most interface potential measurements) was smaller than the size of the plotted data point.

2.3. Scanning electron microscopy with energy dispersive X-ray spectroscopy

Scanning electron microscopy (SEM) and energy dispersive X-ray spectroscopy (EDXS) were used to track surface modifications of the Fe (0) anodes caused by electrolysis. Iron anode samples used in four different electrolyte solutions were prepared by performing 10 of the previously described EC experiments over 3 d in each of the chloride, bromide, nitrate, and citrate solutions. Anodes were rinsed briefly with DI water after each EC experiment and were air dried overnight, which is similar to the operation and drying cycles of electrodes during long term Fe(0) EC field treatment [3]. The surface morphology (SEM) and chemical composition (EDXS) of the Fe(0) anodes were examined before electrolysis (0 runs, cleaned and polished) and after 2 runs (1 d), 6 runs (2 d) and 10 runs (3 d). SEM images were collected with a FEI Quanta 3D FEG SEM equipped with a voltage contrast detector. EDXS spectra were obtained using an Oxford X-Max 20 mm² EDXS spectrometer. An accelerating voltage of 20 kV and a current of 8 nA were used during data collection. A conductive coating was not used on the Fe(0) anodes.

3. Results

3.1. Time dependent Fe concentration and anodic interface potential

In Fig. 2, we present an overview of representative data obtained in each electrochemical experiment (i = 1 mA/cm²) for three electrolyte solutions: chloride, sulfate and nitrate. The Fe generated (Feexp) in both the chloride and sulfate solutions follows closely the theoretical value (Fe0) for the production of Fe(II), which is highlighted by one of the shaded regions in Fig. 2a. A subtle trend in the production of Fe with time is also observed for these two electrolytes, with slightly decreased Feexp values relative to Fe0, assuming the constant production of Fe(II) (Eq. (1)). Nevertheless, the Feexp in the chloride and sulfate electrolyte solutions generally increased linearly with time and a Faradaic efficiency > 0.85 was obtained at all time steps. By contrast, the Feexp in the nitrate solution was negligible, yielding a Faradaic efficiency < 0.1 throughout electrolysis.

The time dependent anodic interface potential (Ea) measurements for the three electrolyte solutions are shown in Fig. 2b. The Ea in the nitrate solution was substantially higher (1.6 V vs Ag/AgCl) than that measured in the chloride or sulfate solutions (< 0.5 V vs Ag/AgCl), which is consistent with the differences in time dependent Feexp. For the chloride and sulfate solutions, the Ea at i = 1 mA/cm² never increased above the E for O2 or Cl2 production (E ≈ 1.02 V vs Ag/AgCl for O2 at pH 7; E ≈ 1.68 V vs Ag/AgCl for Cl2 at 10 mM chloride). This result indicates that significant anodic production of O2 and Cl2 was unlikely. Although the Ea in the sulfate solution decreased slightly over time, it was relatively stable in the chloride and sulfate solutions after the onset of electrolysis. There were no rapid fluctuations to indicate a change in the dominant anodic redox reaction.

Analogous time dependent measurements of Feexp and Ea were obtained for every electrochemical experiment in our study. However, to simplify the presentation of our results, we note that each data point presented in subsequent figures corresponds to the Faradaic efficiency at the end of electrolysis (i.e. the final time step in Fig. 2a) and Ea measured at the midpoint of electrolysis.

Fig. 2. Measurements of (A) Fe in the bulk electrolyte solution and (B) the anodic interface potential as a function of time for selected solution compositions: 10 mM chloride (crosses), 5 mM sulfate (circles), and 10 mM nitrate (squares). The i was 1 mA/cm² and the initial pH was 7.5. The shaded regions in (A) show the hypothetical Faradaic Fe concentrations, assuming the production of Fe(II) and Fe(III).

3.2. Influence of solution composition

3.2.1. Chloride, bromide and sulfate

The Faradaic efficiencies (top panels) and Ea (bottom panels) are plotted as a function of a wide range of i (0.25–50 mA/cm²) for 8 different inorganic and organic single anion solutions in Fig. 3. The Faradaic efficiencies for the chloride and bromide electrolytes were largely independent of i, with values > 0.8 observed at all i (Fig. 3a). Coinciding with these high Faradaic efficiencies, the Fe(0) anode surfaces after electrolysis in both chloride and bromide solutions developed macroscopic surface modifications that were low in contrast on SEM images, consistent with the formation of less electron dense Fe (oxyhydr)oxide corrosion layers (Fig. 4 and Figs. S1 and S2 in the Supplementary material). By contrast, the Faradaic efficiency in the sulfate solution was strongly dependent on i. At i < 3 mA/cm², the Faradaic efficiency decreased to < 0.1, which was similar to that for the bromide and chloride solutions. However, at i ≥ 3 mA/cm², the Faradaic efficiency decreased to < 0.1, which indicates that Fe(II) generation was not the dominant anodic reaction.

The shape and position of the Ea vs i curves for the chloride and bromide solutions matched closely (Fig. 3b), which is consistent with their similar Faradaic efficiencies. On a log scale, the Ea increased exponentially from ~0.1 to > 5.0 V vs Ag/AgCl with increases in i for both the chloride and bromide solutions. However, there was a linear relationship between Ea and i when i was plotted on a linear scale (Fig. S3). Despite the high Faradaic efficiencies in the chloride and bromide solutions, the Ea at i > 20 mA/cm² exceeded the standard state E° for the O2/H2O, Cl2/Cl−, and Br2/Br− redox couples. Although experiments at high i yielded Ea well above the potential required for evolution of Cl2 (E ≈ 1.68 V vs Ag/AgCl), Br2 (E ≈ 1.41 V vs Ag/AgCl) and O2 (E ≈ 1.02 V vs Ag/AgCl), a high Faradaic efficiency was measured and no bubble production was observed. These results indicate that Ea
levels above 5.0 V vs Ag/AgCl do not necessarily entail significant O2 evolution or chloride/bromide oxidation in these solutions.

Consistent with the distinct i-dependence of the Faradaic efficiency observed in the sulfate solution, the $E_A$ vs i curve for sulfate differs relative to that of chloride and bromide. At $i < 3$ mA/cm$^2$, the $E_A$ vs i curve for sulfate follows that of chloride and bromide. However, a nonlinear jump in $E_A$ occurs at $i > 3$ mA/cm$^2$, leading to a ~2 V vs Ag/AgCl increase in $E_A$ relative to that of chloride and bromide electrolyte solutions for $i > 3$ mA/cm$^2$. Notably, this abrupt increase in $E_A$ coincided with the dramatic decrease in the Faradaic efficiency observed in the same i-range. This indicates the existence of a threshold $i$ in the sulfate solution that delineates the electrochemical domain of Fe (0) oxidation.

3.2.2. Carbonate, phosphate and nitrate

In contrast to the chloride and bromide solutions, poor Faradaic efficiencies were observed in the carbonate, phosphate and nitrate solutions, regardless of $i$ (Fig. 3c). Even for experiments performed at the lowest $i$, which should lead to the lowest $E_A$ based on Eq. (2), the Faradaic efficiency never exceeded 0.1 for these single anion solutions. In addition, the Fe(0) anode showed no evidence for any Fe (oxyhydr) oxide corrosion layers after 10 runs in the nitrate electrolyte (Fig. 4, Figs. S1 and S2). This observation contrasts with the behavior of the chloride and bromide solutions and is consistent with the passivation of the anode surface [32,33].

Because the ionic strength of the nitrate and halide ion electrolytes was identical (Table 1), these results imply that ionic composition, rather than conductivity, plays a central role in determining the Faradaic efficiency of Fe(0) EC. The $E_A$ vs i curves for the carbonate, phosphate and nitrate electrolytes had similar shape to that of chloride and bromide but were uniformly offset by $+2$ to $+4$ V vs Ag/AgCl (Fig. 3d). The $E_A$ vs i curves for the carbonate, phosphate and nitrate electrolytes overlap almost perfectly, which is consistent with the similarly low Faradaic efficiencies for these electrolytes. Interestingly, at $i > 3$ mA/cm$^2$, the $E_A$ measurements for these electrolytes are identical to those of the sulfate electrolyte, which also had poor Faradaic efficiency in this i-range. The high $E_A$ (> 1.4 V vs Ag/AgCl) observed in the carbonate, phosphate and nitrate solutions at all $i$ was accompanied by the formation of gas bubbles on the anode (Fig. S4) and poor Faradaic efficiency. In addition, dissolved O2 measurements in the nitrate solution (Fig. S5) revealed a rapid increase in dissolved O2, with values reaching above the air saturated maximum of approximately 9 mg/L, which provides conclusive evidence for the anodic formation of O2 in these three solutions.

3.2.3. Citrate and formate

The properties of the formate solution were most similar to those of the sulfate solution, with the Faradaic efficiency exhibiting a strong i-dependence (Fig. 3c). However, even at low $i$, the Faradaic efficiency in the formate solution was never as high as for sulfate. The Faradaic efficiency in the formate solution also decreased gradually with $i$, i.e. a transitional Faradaic efficiency near 0.5 was observed at $i = 1$ mA/cm$^2$. For the citrate solution, the Faradaic efficiency resembled that of the nitrate solution, with negligible electrochemical Fe(0) oxidation throughout the entire i-range. Furthermore, consistent with the nitrate solution, the surface of the Fe(0) anode after 10 electrochemical experiments in the citrate solution resembled that of a freshly polished Fe (0) anode (Fig. 4, Figs. S1 and S2), indicating surface passivation.

The $E_A$ at the lowest $i$ (highest Faradaic efficiency) in the formate solution was similar to that of chloride/bromide, but at $i > 1$ mA/cm$^2$, the $E_A$ vs i curve followed that of nitrate. Similar to the results in the sulfate solution, the crossover $i$ between high and low Faradaic efficiency in the formate solution coincided with an abrupt increase in $E_A$. For citrate, the position and shape of the $E_A$ vs i curve resembled that of the nitrate solution, which is consistent with its poor Faradaic efficiency.

3.3. Impact of pH on the Faradaic efficiency and $E_A$

The influence of pH was examined in the sulfate and phosphate solutions because these ions affected the Faradaic efficiency differently in single anion systems (Fig. 3) and also have distinct pH dependent speciation, i.e. the dominant sulfate species remains as SO4$^{2-}$ above pH 3 ($pK_{a1} = 3$, $pK_{a2} = 1.9$), whereas phosphate deprotonates ($pK_{a1} = 2.1$, $pK_{a2} = 7.2$, $pK_{a3} = 12.7$) [47]. For both sulfate and phosphate solutions, the Faradaic efficiency and $E_A$ ($i = 1$ and 3 mA/cm$^2$) depended strongly on pH. At i-values of 1 and 3 mA/cm$^2$, the Faradaic efficiencies in the sulfate and phosphate solutions were above 1.0 at acidic pH (4.0 ± 0.5) and below 0.1 at basic pH (11.0 ± 0.5). Faradaic efficiencies above 1.0 for both acidic sulfate and phosphate solutions.
solutions can be explained by chemical induced release of Fe, likely a result of proton promoted Fe(0) oxidation. The major difference between the impact of pH in the sulfate and phosphate solutions is the threshold pH and i at which the Faradaic efficiency changes from high to low. For phosphate, efficient electrochemical Fe(0) oxidation was only observed at acidic pH, whereas the Faradaic efficiency remained high at circumneutral pH in the sulfate solution at i = 1 mA/cm².

The same inverse relationship between Faradaic efficiency and $E_A$ found in the single anion solutions was also observed in the sulfate and phosphate solutions at different pH values. For example, the lowest $E_A$ for both solutions was observed at acidic pH, which showed the highest Faradaic efficiencies, whereas $E_A$ was greater than 1.0 V vs Ag/AgCl for all samples with poor Faradaic efficiency. It is also noteworthy that decreasing solution pH from 7.5 ± 0.5 to 4.0 ± 0.5 decreased the

Fig. 4. Scanning electron micrographs of a polished Fe(0) anode (A and B), and Fe(0) anodes after 10 runs in various electrolyte solutions: bromide (C and D), chloride (E and F), citrate (G and H), nitrate (I and J). The panels on the right are magnified from the region in the dotted rectangles on the left.
3.4. Faradaic efficiency in chloride containing, binary anion solutions

In Section 3.2, we found that the behavior of the chloride solution contrasted that of the nitrate and carbonate solutions with respect to Faradaic efficiency and \( E_A \). Therefore, to evaluate the relative impact of ubiquitous ions that facilitate or prevent electrochemical Fe(0) oxidation, we performed EC experiments \( (i = 1 \text{ mA/cm}^2) \) in binary anion solutions with a wide range of nitrate to chloride or carbonate to chloride molar ratios \( ([\text{NO}_3^-]/[\text{Cl}^-]) \) or \( ([\text{HCO}_3^-]/[\text{Cl}^-]) \). Despite the 15 fold increase in conductivity for the NO3-GW 1 chloride, the NO3-GW 1 solution still displayed poor Faradaic efficiency. This result suggests that the [NO3\(^-\)]/ [Cl\(^-\)] molar ratio above 30 for this solution, this result suggests that the [NO3\(^-\)]/ [Cl\(^-\)] and [HCO3\(^-\)]/ [Cl\(^-\)] ≥ 1.

At [NO3\(^-\)]/[Cl\(^-\)] molar ratios below 20, the nitrate and chloride binary anion solution behaved similarly to the chloride solution, with high Faradaic efficiencies and \( E_A < 0.5 \text{ V vs Ag/AgCl} \). The persistence of efficient electrochemical Fe(0) oxidation and low \( E_A \) at [NO3\(^-\)]/[Cl\(^-\)] ratios near 1.0 indicates that the presence of chloride outweighs the detrimental impacts of nitrate on an equimolar basis. However, at [NO3\(^-\)]/[Cl\(^-\)] ratios above 30, the Faradaic efficiency decreased and the \( E_A \) increased, leading to a solution with properties akin to the nitrate single anion system. These abrupt changes in Faradaic efficiency and \( E_A \) suggest that a narrow range of [NO3\(^-\)]/[Cl\(^-\)] ratios near 20 to 30 separate the electrochemical domains of Fe(0) oxidation and O\(_2\) evolution in the Fe(0) EC system.

The relationship between Faradaic efficiency and \( E_A \) with increasing [HCO3\(^-\)]/[Cl\(^-\)] was similar as that for the nitrate and chloride system, but the transition from low to high Faradaic efficiency occurred at a greater [HCO3\(^-\)]/[Cl\(^-\)] ratio. Whereas the drop in Faradaic efficiency and spike in \( E_A \) occurred above [NO3\(^-\)]/[Cl\(^-\)] ratios of 20, the critical [HCO3\(^-\)]/[Cl\(^-\)] ratio was 100. This indicates that carbonates solutions require a lower chloride content to ensure efficient electrochemical Fe(0) oxidation than nitrate solutions.

3.5. Faradaic efficiency as a function of \( i \) in synthetic groundwaters

In Fig. 7, the Faradaic efficiency and \( E_A \) are plotted as a function of \( i \) for synthetic Bangladesh groundwater (SBGW) and for two synthetic nitrate contaminated groundwaters (NO3-GW 1 and NO3-GW 2). The ionic composition and pH of these synthetic groundwaters, which cover a range of potential source waters for Fe(0) EC treatment, are summarized in Table 2. High Faradaic efficiency in the SBGW solution was observed for all \( i \), which is consistent with its high chloride concentration and low [HCO3\(^-\)]/[Cl\(^-\)] and [NO3\(^-\)]/[Cl\(^-\)] ratios. Coinciding with the high Faradaic efficiency, the \( E_A \) vs \( i \) curve for the SBGW solution also matched the shape and position of that for the chloride single anion solution. By contrast, the poor Faradaic efficiency and high \( E_A \) for the NO3-GW 1 solution resembled those of the nitrate solution, which agrees with the [NO3\(^-\)]/[Cl\(^-\)] molar ratio above 30 for this solution. Despite the 15 fold increase in conductivity for the NO3-GW 1 solution relative to the nitrate solution and the presence of 0.5 mM chloride, the NO3-GW 1 solution still displayed poor Faradaic efficiency. This result points to the key role of the [NO3\(^-\)]/[Cl\(^-\)] molar ratio in determining the Faradaic efficiency, rather than the conductivity or absolute chloride concentration.

For the NO3-GW 2 solution, the Faradaic efficiency was greater than 0.9 for \( i \leq 10 \text{ mA/cm}^2 \) but dropped abruptly to below 0.1 at \( i > 20 \text{ mA/cm}^2 \). Given the moderate [NO3\(^-\)]/[Cl\(^-\)] ratio of ~18 mol/mol for this solution, this result suggests that the [NO3\(^-\)]/[Cl\(^-\)] ratio dividing the domains of Fe(0) oxidation and O2 evolution also depends on \( i \). Furthermore, although acidic pH improved the Faradaic efficiency in sulfate and phosphate solutions (Section 3.3), the results in the NO3-GW 2 matrix indicate that high \( i \) can still promote poor Faradaic efficiency in acidic solutions. The \( E_A \) vs \( i \) curve for the NO3-GW 2 solution increased sharply at the \( i \)-value for which the Faradaic efficiency decreased, which is similar to the \( E_A \) vs \( i \) curve for sulfate. Despite having a region of where Faradaic efficiency was poor, the \( E_A \) vs \( i \) curve for the NO3-GW 2 solution was positioned ~0.5 V vs Ag/AgCl lower than that of the chloride solution. We attribute this difference in the position of the \( E_A \) vs \( i \) curve for the NO3-GW 2 matrix to its acidic pH, leading to a lower open circuit potential (~0.6 V vs Ag/AgCl) relative to the single anion solutions (~0.2 V vs Ag/AgCl). This was also observed in the acidic sulfate and phosphate solutions.

4. Discussion

4.1. Factors affecting the Faradaic efficiency

4.1.1. Ionic composition

For the majority of single anion solutions and synthetic groundwater matrices, the Faradaic efficiency was largely independent of \( i \). Rather, the type of anion in the solution played a more critical role in governing the Faradaic efficiency and \( E_A \), and thus the potential for electrode passivation. For example, regardless of \( i \), Faradaic efficiency was high in the chloride and bromide solutions, whereas it was minimal in nitrate and citrate solutions. Modifications of the Fe(0) anode surface mirrored these trends in Faradaic efficiency, with the chloride and bromide solutions producing macroscopic surface layers of oxidized Fe and the nitrate and citrate solutions yielding passivated Fe(0) anode surfaces that resembled cleaned electrodes (Figs. 4, Figs. S1 and S2). Although previous EC studies have identified conductivity as a key operational parameter [7,31,48], our results demonstrate substantial differences among solutions with identical conductivity, which reveals the critical importance of the ion composition.

Comparing the Faradaic efficiency and \( E_A \) measurements as a function of \( i \) for all solutions in Fig. 3 (Section 3.2), it is possible to rank the anions in the following order of negative influence on Fe(0) EC performance: chloride < bromide < sulfate < formate < carbonate < phosphate ≤ citrate ≤ nitrate. With the exception of nitrate, this sequence of anions generally follows the same order of increasing stability constants with Fe(II) or Fe(III) (Table S1). For example, chloride and bromide do not form strong complexes with Fe(II/III), whereas phosphate and citrate have a high affinity for Fe(II/III) [49,50]. A similar anion sequence has been observed in previous studies investigating the reactivity of Fe(0) metal toward contaminant reduction, which reported that chloride enhanced electron transfer, whereas the sorption of phosphate and citrate diminished the redox activity of Fe(0) metal by blocking reactive sites [32,33]. Therefore, based on this connection between detriment to Faradaic efficiency and the Fe(II/III) stability constants, the effect of single anions can be explained by the bonding of high affinity anions (e.g. phosphate, citrate) to oxidized Fe at the electrode surface, resulting in a thin, but impermeable passivating film that prevents the release of Fe(II/III) to solution. Although a molecular-scale investigation of the surface layers formed on the electrodes having poor Faradaic efficiency is beyond the scope of our work, the formation of a nanoscale passivating film is consistent with the absence of corrosion spots on the anodes used in the citrate experiments and is supported by previous studies reporting the formation of a phosphate-bearing layer that inhibits the corrosion of Fe and stainless steel in phosphate solutions [51,52].

The exception to the trends in anion sequence is nitrate, which inhibited Fe(0) oxidation but does not readily complex with Fe(II/III). However, the mechanism by which nitrate decreases the Faradaic efficiency is expected to be different than for anions with a high affinity for Fe(II/III). Rather than complexing with Fe(II/III) at the Fe(0) surface, nitrate is known to oxidize Fe(0) metal, producing a thin, Fe (oxyhydr)oxide passive film that inhibits ion diffusion [32]. Despite the different inhibition mechanisms expected for nitrate and high affinity anions, both mechanisms are expected to lead to nanoscale passivating films on the Fe(0) anode surface, which are imperceptible with SEM [53,54].
4.1.2. Range of \( i \)

For the single anion sulfate and formate solutions and the synthetic NO\(_3\)-GW 2 matrix, the \( i \)-range influenced the domains of Fe(0) oxidation and O\(_2\) evolution. Importantly, the three solutions also displayed intermediate properties with respect to ionic composition and pH. For example, the stability constants of sulfate and formate for Fe(II/III) lie between those of chloride/bromide and phosphate/citrate [50]. In addition, the [NO\(_3\)^−]/[Cl\(^−\)] ratio for the NO\(_3\)-GW 2 solution (18 mol/mol) occurs near the threshold ratio (20–30 mol/mol) that separates Fe(0) oxidation and O\(_2\) evolution (Fig. 5). Although \( i \) was less important for end member solutions of the anion sequence described in Section 4.1.1, this parameter played an important secondary role for solutions with intermediate compositions.

We attribute the impact of \( i \) on the anodic reaction and potential for electrode passivation to the Butler-Volmer relationship, given in Eq. (2), which shows that an increase in \( i \) leads to an increase in \( E_A \). Because an increase in \( E_A \) can promote anodic reactions with high \( E^\circ \) (Fig. 1a), an increase in \( i \) can lead to the oxidation of H\(_2\)O or other solutes instead of Fe(0), which would decrease the Faradaic efficiency. Consistent with this relationship, we observed a nonlinear jump in \( E_A \) when the dominant anodic reaction switched from Fe(0) oxidation to O\(_2\) evolution (Fig. 3) for solutions in which the Faradaic efficiency depended on \( i \) (e.g. sulfate). For the sulfate solution, the magnitude of this jump in \( E_A \) (∼2 V vs Ag/AgCl) was the exact value separating the \( E_A \) vs \( i \) curves for Fe(0) oxidation (chloride/bromide) and O\(_2\) evolution (citrate/nitrate). Therefore, we conclude that the measurements of the position of the \( E_A \) vs \( i \) curve can be a useful and low cost indicator of the anodic reaction occurring during Fe(0) EC.

4.1.3. Solution pH

For both the sulfate and phosphate solutions, pH influenced the Faradaic efficiency considerably, with efficient Fe(0) oxidation occurring only at low pH. Considering that phosphate is a detrimental end member species in the anion sequence (Section 4.1.1), the high Faradaic efficiency observed in the phosphate solution at low pH suggests that pH effects can overcome the negative impact caused by high affinity anions, particularly at low \( i \). The pH dependence of the Faradaic efficiency observed in our study can be explained by a number of contemporaneous processes. First, as pH increases, \( E^\circ \) for the H\(_2\)O/O\(_2\) redox couple decreases [50], leading to a narrower field of H\(_2\)O stability and increased likelihood of O\(_2\) evolution for a given \( E_A \). Second, increasing pH leads to orders of magnitude faster rates for Fe(II) oxidation by O\(_2\) [55] as well as decreased Fe(III) solubility (see Supplementary material for speciation calculations and Fig. S5 for Fe(III) saturation indices). Both of these processes, which are applicable in all solutions regardless of ion composition, are expected to promote the formation of Fe(III) (oxyhydr)oxide passivating films that inhibit electron transfer across the anode and Fe(II/III) release. Finally, for the phosphate electrolyte, the solubility of Fe(II) in the presence of phosphate depends strongly on pH. For example, as pH increases from 4 to 7, the saturation index of vivianite (Fe\(_3\)(PO\(_4\))\(_2\)·8H\(_2\)O) increases ∼10 orders of magnitude (−2.9 to 7.4, Fig. S6), leading to conditions that favor the formation of anodic surface layers that inhibit Fe(II) release. These pH dependent changes in solid phase solubility in the phosphate solution, which do not occur to the same extent in the sulfate solution (Fig. S6), explain the stronger impact of pH in the presence of phosphate relative to sulfate (Fig. 6).

4.2. Implications for EC treatment in the field

4.2.1. Field treatment recommendations

Effective treatment of contaminated waters by Fe(0) EC relies on the production and transport of Fe(II) from the Fe(0) anode surface to the bulk solution. For example, Cr(VI) is removed in EC systems by reaction with EC generated Fe(II), to form insoluble Cr(III) [6,7], whereas As(III) is removed by oxidation to As(V) by Fenton type intermediates and sorption to EC generated Fe(III) precipitates [35,56]. Therefore, regardless of the contaminant removal mechanism, a critical loss of treatment efficiency occurs when the applied current drives O\(_2\) evolution at the expense of Fe(0) oxidation. In addition, as shown in this work, a drop in Faradaic efficiency is accompanied by a spike in \( E_A \), which is consistent with electrode passivation. This increases the total EC cell voltage and can result in higher power consumption during EC treatment. Therefore, EC system effectiveness in the field relies on...
avoiding poor Faradaic efficiency.

Based on our results, the i-range can impact the Faradaic efficiency for specific solution compositions, with efficient Fe(0) oxidation favored at $i < 5 \text{ mA/cm}^2$, which is similar to the i-range used in previous Fe(0) EC field experiments [3]. However, although $i$ should not be ignored in field treatment, our results indicate that the solution composition is the dominant parameter governing the Faradaic efficiency. Source waters with low chloride levels, alkaline pH and high concentrations of nitrate, carbonate and other high affinity oxyanions can be a barrier to high Faradaic efficiency. Treatment influents with such solution chemistry can be encountered in nitrate contaminated groundwater systems, including those impacted by agricultural runoff [57], sewage [58], and radionuclide enrichment activities [37,42,44]. Furthermore, chloride concentrations in groundwaters contaminated by geogenic arsenic can be low and $[\text{HCO}_3^-]/[\text{Cl}^-]$ ratios above the critical ratio of 100 mol/mol (Fig. 6) are not uncommon in regions targeted for Fe(0) EC treatment in South Asia (e.g. Table 7.1 in the British Geologic Survey analysis of Bangladeshi groundwater [39]).

Although decreasing pH to near 4 can negate some of the impacts of ion composition, adjusting pH at field treatment scale is impractical for several reasons, e.g. hazardous and costly chemicals, pH readjustment of treatment effluent, skilled labor requirement. Instead, our results indicate that an influent composition favoring $\text{O}_2$ evolution can be overcome easily and at low cost by simple additions of small quantities of NaCl (table salt) prior to electrolysis. Although the importance of halide ions to Fe(0) EC performance has been suggested previously [25,31,48], our results have uncovered distinct $[\text{NO}_3^-]/[\text{Cl}^-]$ and $[\text{HCO}_3^-]/[\text{Cl}^-]$ molar ratios that separate the Fe(0) oxidation and $\text{O}_2$ evolution domains (Fig. 6). Based on this ratio, the addition of 1 mM NaCl can overcome nitrate and carbonate concentrations up to 20 and

![Fig. 6. Influence of the (A) $[\text{NO}_3^-]/[\text{Cl}^-]$ and (B) $[\text{HCO}_3^-]/[\text{Cl}^-]$ molar ratios on the Faradaic efficiency and $E_a$ for Fe(0) EC experiments performed at $i = 1 \text{ mA/cm}^2$. Filled symbols represent Faradaic efficiency; empty symbols indicate $E_a$.](image)

![Fig. 7. Faradaic efficiency (A) and anodic interface potential (B) as a function of $i$ for three synthetic groundwaters (composition presented in Table 2).](image)
100 mM, respectively. This result suggests that a single purchase of 10 kg NaCl can provide enough chloride to treat over 150 m³ of groundwater containing moderate to high nitrate or carbonate levels. Previous analyses of fully operational Fe(0) EC systems (10,000 L/d capacity) have estimated total costs to be < $1.50/m³ of treated water [5]. Chloride containing, 

[II,III] [50], promote Fe(0) oxidation. In chloride containing, 

capacity) have estimated total costs to be < $1.50/m³ of treated water groundwater containing moderate to high nitrate or carbonate levels. Therefore, assuming a conservative price of table salt ($10/10 kg), the proposed addition of small quantities of NaCl to ensure high Faradaic efficiency would increase these estimated costs < 5%. Although this option adds a pretreatment step, it is very simple and does not require hazardous chemicals or major supply chain infrastructure and would render the electrolyte solution suitable for a wide range of i. In addition, our results indicate that measurement of $E_A$, which does not require expensive equipment, and comparison of this value with i can inform system operators when NaCl treatment is required.

4.2.2. Insights into field treatment performance over time

Several previous Fe(0) EC field studies have reported a drop in system performance over extended operating times (weeks to months) and cycles of electrode polarization, i.e. periods of operation (wet) and storage (dry) [45,59]. One previously proposed explanation for this decreased field performance is a drop in Faradaic efficiency resulting from O₂ evolution after many cycles of operation. However, extended EC field trials typically contain few systematic measurements of key parameters over time, i.e. no regular measurements of $E_A$ or Fe in suspension [3]. As a consequence, explanations of the efficiency drop observed during field treatment remain speculative.

We observed a subtle decrease in the Faradaic efficiency on single run time scales (Fig. 2), which was reproduced in both the sulfate and chloride solutions. Over the 10 runs required to produce anode samples for SEM imaging, a decrease in the Faradaic efficiency was also observed (Fig. S7). These drops in Faradaic efficiency over a single run (Fig. 2) or multiple runs (Fig. S7) did not coincide with an offset in $E_A$ indicative of O₂ evolution (or Fe(III) production). Although Fe(0) EC field systems differ in size and treatment time relative to our experimental setup [1,3,59], the absence of an increase in $E_A$ over several days (10 runs) in this work suggests that O₂ evolution after extended treatment cycles in the field is not the primary cause of decreased system performance. This conclusion is consistent with the relatively low i (~ 1 mA/cm²) used in previous Fe(0) EC field trials [3]. Rather, in light of the build up of anodic surface layers in the chloride/bromide solution (Fig. 4, Figs. S1 and S2), our results support the hypothesis that field treatment performance decreases over time because EC generated Fe(II) is trapped at the anode by macroscopic Fe (oxyhydr)oxide surface layers (e.g. magnetite, goethite) formed on the electrode after extended operation [45].

5. Conclusions

The ionic composition and pH of the electrolyte solution largely determined the Faradaic efficiency of Fe(0) EC, with changes in i influencing the domains of Fe(II) production and O₂ evolution only in certain electrolyte solutions (e.g. sulfate and formate). By ranking single anion solutions according to their detriment to Faradaic efficiency (chloride ≫ bromide ≫ sulfate ≫ carbonate ≪ phosphate ≪ citrate ≪ nitrate), we elucidated a useful relationship between the propensity of ions to form strong complexes with Fe(II,III) and low Faradaic efficiency. The exception to this trend is nitrate, which is expected to interact with the Fe(0) anode differently [32]. Based on this relationship, we expect that solutions containing ions with strong affinity for Fe(II,III), such as silicate, would favor O₂ evolution, whereas solutions containing ions such as iodide and fluoride, which have a weaker affinity for Fe(II,III) [50], promote Fe(0) oxidation. In chloride containing, 

binary anion solutions, we found the domains of Fe(0) oxidation and O₂ evolution (i = 1 mA/cm²) were separated by discreet [NO₃⁻]/[Cl⁻] and [HCO₃⁻]/[Cl⁻] molar ratios, which were confirmed in synthetic groundwater solutions. This result indicates that a simple amendment of NaCl to source waters with compositions that favor O₂ evolution (e.g. sites of nitrate pollution or arsenic contaminated groundwater with low chloride content) can ensure high Faradaic efficiency, which is critical to optimize the performance of Fe(0) EC systems in the field.

The $E_A$ vs i curves for the chloride and bromide solutions overlapped and reached $E_A$ far greater than the $E^0$ of H₂O/O₂ Cl⁻/Cl⁻ and Br⁻/Br₂ couples, but high Faradaic efficiency was still observed in these solutions. This conclusion demonstrates that significant anodic Cl₂ and Br₂ formation is unlikely to occur during Fe(0) EC even at extreme $E_A$ (and i), although their formation has been reported for different electrode materials (e.g. boron-doped diamond or Al(0) electrodes) [60,61]. Finally, the $E_A$ vs i curves for the citrate and nitrate solutions, which exhibited poor Faradaic efficiency, also overlapped and were situated 2–4 V vs Ag/AgCl above that of the chloride/bromide solutions. Comparison between the $E_A$ vs i curves of both high and low Faradaic efficiency solutions showed that the $E_A$ vs i curve, rather than the $E_A$ alone, is necessary to identify the primary reaction occurring on the Fe (0) anode. This information can be used to accurately determine in real time if Fe(0) EC systems are operating at high Faradaic efficiency using only simple, low cost, and chemical-free field measurements of $E_A$ with knowledge of i.

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Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at http://dx.doi.org/10.1016/j.jece.2017.09.008.

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