Short Communication

High-throughput corrosion quantification in varied microenvironments

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1. Introduction

As technological advancement enables us to access increasingly harsh environments, ranging from implanted biomedical devices to undersea oil and gas deposits, we continue to test the limits of corrosion susceptibility and require quantitative insights to predict and extend structural reliability. The highly complex nature of corrosion processes, as well as the wide range of relevant environmental variables, complicates rigorous identification of corrosion susceptibility. For example, despite mitigation attempts, microbiologically influenced corrosion (MIC) and associated biofilm formation (biofouling) have been implicated in several rapidly progressing, high-profile failures of buried steel pipeline [1–4]. While sulfate reducing bacteria (SRB) are considered a main culprit in anaerobic environments, the complex consortium of microbial species and the biofilms they produce have obviated mechanistic conclusions. Despite much research of the microbiology and the engineering effects, the mechanisms by which MIC enhances ferrous corrosion and the appropriate strategies to mitigate MIC corrosive loss remain elusive [5,6].

The limitations of prevailing experimental techniques to quantify and predict corrosion in controlled environments have led to the development of several higher-throughput measurement methods (see two recent review articles [7,8]). Current high-throughput approaches use either a single metal bulk substrate with areas controllably exposed to different microenvironments [9,10], deposited metallic thin films in each microenvironment [11–13], or multiple electrode systems (2 or 3) within each microenvironment [14–18]. While existing high-throughput methods hold tremendous promise in certain conditions, their production is not necessarily straightforward and scalable, and their applicability for biotic studies which require complete isolation between microenvironments to prevent cross-contamination and a continuous surface for bacterial colonization is unclear.

Here we describe a high-throughput corrosion testing platform that assays corrosion susceptibility of metals by tracking the changing resistance of thin wires and colorimetric changes of the surrounding solution within a 96-well format. While resistance-based probes have been validated as an effective means to measure corrosion at much larger scales with only a few samples [19–21], multiplexing of these measurements enables more rapid consideration of corrosion susceptibility and rates that requires only sub-ml-scale volumes of fluid. This approach allows for independent control of cues – including chemical composition of surrounding medium, gas concentrations, temperature, type and number of microbial species. Results are obtainable over the course of hours to days. We validate this approach through consideration of low-carbon steel corrosion in both aerobic and anaerobic environments, and in both abiotic and biotic conditions.

2. Experimental

2.1. Multiplexed platform

We designed a novel multiplexed platform to measure in parallel the electrical resistance of metallic samples placed within a
range of controlled fluid microenvironments (see Supplemental methods for a detailed platform description). The multiplexer had 384 addressable channels, used here to conduct 96, 4-wire resistance measurements when coupled with an Agilent 34420A micro-ohm meter. A set a ribbon cables connected the multiplexer to a “test fixture” circuit board designed to sit directly on top of a 96-well plate suspending a u-shaped wire loop within each well. To align the 96-well plate and the test fixture, an aluminum frame was constructed; this allowed the wire array to lower into the center of each well and then hold the two together tightly. The frame also allowed light to pass through wells, as required for optical imaging. Labview 2013 was used to control data acquisition. An Arduino 1.5.4 macro was used to set the current channel on the multiplexer. After setting the channel, the ohmmeter was triggered to begin taking measurements. The voltmeter was set to the 1 Ω range with an integration time of 20 power line cycles and offset compensation. At each time point, five measurements were taken for each channel and averaged after discarding the highest and lowest measurement. All 96 wells could be assessed in this manner in 8 min 15 s (5.15 s/channel).

2.2. Metal sample preparation

Two sources of low carbon steel (1008, McMaster Carr) were used for experiments: a 27 gauge (361 μm diameter) wire used in the as-received condition, and a 12 gauge (2.05 mm diameter) wire drawn into thinner wire segments (see Supplemental methods). The thin black oxide coating on the wires was removed by light sanding with 1500 grit sandpaper and then cleaned by wiping with 70% ethanol. The wire was cut to approximate size and shaped into a “u”-shape by winding tightly between a series of posts of 3.68 mm diameter. This step provided uniform curvature and microstructural damage (due to plastic deformation upon bending) to all wires. Wire diameter was assessed via optical microscopy at a magnification of 20×. Before corrosion susceptibility experiments, the wires were again cleaned and sterilized by soaking in 70% ethanol for 10 min.

2.3. Corrosion assays and analysis

The 96-well plate format allowed us to independently control both the fluid microenvironments and the chemical composition or microstructure of the metal wire within each well. Thus 96 separate experiments were conducted in parallel (a set of experiments, with the potential for replicates of a given condition within that set). Sets were performed either aerobically on the bench top or anaerobically within an anaerobic incubator. Prior to the start of experiments, the wires were again cleaned and sterilized by soaking in 70% ethanol for 10 min.

3. Results and discussion

3.1. Rationale for high-throughput resistance and colorimetric platform

Corrosion of metals involves the loss of structural metal atoms from the bulk to the ionic or complexed forms. A system that can detect very small amounts of material loss could therefore rapidly assess corrosion. Here we employ two techniques to achieve this sensitivity to corrosion susceptibility: the resistance change of thin u-shaped wires loops and the color change of the surrounding fluid medium. Because the resistance of a wire is based on the material resistivity (constant at a given temperature), the length of the wire (constant), and the cross-sectional area (changes with corrosion), monitoring a change in resistance can assess the degree to which the cross-sectional area has decreased due to corrosion (Fig. 1A). Additionally, the darkness and hue of the surrounding medium can give an indication of the course of corrosion and corrosive product. Taken together, they can provide a rapid assessment of the rate, course, and type of corrosion occurring. Fig. 1B shows an example of the change of resistance and a visual representation of a wire corroding in an aerobic salt solution. Correspondingly, the wire exhibited a visible decrease in diameter as the corrosion proceeded.

The 4-wire resistance measurements in Fig. 1B were taken with an Agilent 34420A micro-ohm meter, which nominally has noise of 1–2 μΩ. However, even when using low-resistance Kelvin probes, taking measurements on more than a single wire per experiment required manually attaching, detaching, and replacing the leads; this repeated manual manipulation caused resistance measurements to vary by milliohms among repeated measurements, rendering sensitivity insufficient. Additionally, the time required to move the leads between different experimental wires limited the time resolution of measurements and obfuscated long-term resistance tracking. Therefore, we next designed a high-throughput integrated platform.

3.2. Design of a high-throughput corrosion detection system

An automated system was designed and constructed, consisting of a multiplexer to controllably address a series of channels and a
"test fixture" with integrated 4-wire leads to hold a series of experimental wires (Fig. 2; see Methods). When connected to the micro-ohm meter and computer, this essentially clones the meter for each channel without requiring any manual intervention. This approach allowed us to individually acquire resistance measurements in serial for wires within each well of a 96-well plate at each time point. An upward facing time lapse camera allowed the simultaneous recording of any color changes of the medium.

3.3. Abiotic, aerobic corrosion in salt conditions

As an initial validation of the system, the resistance change of a 361 µm diameter steel wire in the presence of a range of sodium chloride concentrations was measured in aerobic conditions. Each column of the 96-well array represented replicates of a different condition: air, distilled water, and salt concentrations from 1% to 7% (w/v). Thus there were eight replicates per condition. By monitoring the wires for a period of time in air, the baseline resistance (28–34 mΩ) and signal to noise ratio (~1800:1) were determined (see Supplementary results and Fig. S1A–C). Thus, in theory, this approach using this wire diameter could detect corrosion rates of 1, 0.1, and 0.01 mm/year within 30 min, 5 h, and 2 days, respectively.

At time zero, the appropriate liquids were added to the wells. Fig. 3A shows the average and standard error of the mean for each condition, demonstrating that corrosion could be detected within 1 h and distinctions made among different salt concentrations. Fig. 3B shows the resistance change and corrosion rate at a time point 3 h after the addition of the aqueous solutions. The corrosion rate was estimated by calculating the change in wire radius with time, assuming uniform corrosion occurring equally around the circumference of the wire along its full length. Thus, based upon the wire parameters (resistivity $\rho$ and length $L$) and the change from an initial resistance $R_0$ to a resistance $R_t$ at a later time $t$, the corrosion rate is given as:

$$\text{Corrosion Rate} \left( \frac{\text{mm}}{\text{yr}} \right) = \frac{\text{Radius}}{\Delta t} = \frac{\sqrt{\frac{1}{\rho L} \left( \frac{1}{\sqrt{R_0}} - \frac{1}{\sqrt{R_t}} \right)}}{\Delta t}.$$  

Both the salt concentration dependence and the estimated corrosion rate were in agreement with previous studies that required more time and more sample volume [22,23], thus supporting the rapid estimation of corrosion susceptibility and rate in this multiplexed high-throughput format. The process of acquiring the resistance measurements via continuous data acquisition did not noticeably affect the corrosion, as control wires exhibited indistin-
guishable resistance changes when placed within the same micro-environments but with measurements taken at only a few discrete time points over the same total experiment duration (Fig. S2).

Concurrent with these resistance measurements, colorimetric data were also collected. These images were quantified by changes in greyscale intensity (with negative values indicating darker images) (Fig. 4). In the context of this aerobic saline environment, this colorimetric indicator was less sensitive than resistance, in that the well brightness decreased within the first 30 min but no distinction was possible among varying salt concentrations. The colorimetric signal also saturated within one day as the corrosion products settled to the bottom of the wells. However, the mechanistic complementarity of these colorimetric data for even a well-studied model corrosion environment can be appreciated, in that the orange hue of the precipitate is indicative of Fe₂O₃ corrosion products.

Another set of experiments was conducted with iron wires drawn to various diameters (from 65 to 361 μm), all in 3.5% NaCl saline solutions. With decreasing wire diameter, the resistance increased faster and the signal noise became higher (Fig. S3A–C), although the corrosion rate was approximately the same for all but the smallest wires over the first few hours (again, estimated from the change in wire radius) (Fig. S3D). Those spurious corrosion rates for the smallest drawn wires are attributable to imperfections induced in the wire drawing and wire extraction process, which naturally contributed more strongly to measured resistance of smallest-diameter wires. While the manufacturing process could still be improved to produce wires of sub-100 μm diameter, these data demonstrate the ability to multiplex corrosion susceptibility and corrosion rate measurements on a metal not readily available in the desired wire form. This sample preparation method could be considered for other ductile metals and alloys. However, it should be noted that microstructural changes induced by the drawing process may also modulate corrosion susceptibility; in such cases, this wire processing could limit direct application of these results to industrially prevalent material microstructures.

3.4. Anaerobic corrosion in the presence of sulfate reducing bacteria

We next assayed corrosion in biotic environments, specifically anaerobic aqueous environments comprising sulfate reducing bacteria (SRBs). Biofilms produced by such SRBs readily form on the low carbon steel wires. As Fig. 5 demonstrates, wires can be non-destructively imaged while still in fluid, producing an in situ pseudo cross-sectional view from which the morphology and thickness of the biofilm/corrosion product can be quantified. Dry-

![Fig. 3. Wire resistance changes during abiotic, aerobic corrosion in aqueous saline solutions. (A) Average resistance change of wires in purified water containing NaCl in concentrations from 0% to 7% (w/v) as indicated by legend. Eight wires per condition. (B) The resistance change (left axis) and estimated corrosion rate (right axis) after 3 h for each NaCl concentration (same legend).](image)

![Fig. 4. Corresponding colorimetric changes during abiotic, aerobic corrosion in aqueous saline solutions. The average darkening of the medium in NaCl concentrations from 0% to 7% (w/v) as indicated by legend. The salt solutions can be distinguished from the purified water but not from each other. The color signature saturates within one day. Shaded area and error bars ± standard error of the mean. View of the entire plate at time 0, after 3 hr, and after 24 hr.](image)
grown for a month with no medium exchange) to newly inoculated in fresh media. Intermediate to these extremes of bacterial metabolic activity, varying volumes of fresh media were added to bacteria that had just reached their stationary phase after 6 days of growth (as assessed by turbidity). The higher the nutrient availability, the greater the corrosion extent by day 4 (indicated as an estimated radius change). Furthermore, after an initial corrosion rate peak between days 1 and 2 of up to 0.19 mm/year, the corrosion reached a plateau for SRB conditions at various points after the day 1, consistent with nutrient depletion. The corresponding colorimetric data in Fig. 6 complemented these results: black corrosion product indicated iron sulfides rather than iron oxides. The well color rapidly transitioned to black and saturated between days 1 and 2 for conditions with a metabolically active population of SRBs. Freshly inoculated wells turned dark by day 4, and other conditions did not indicate a change in color (and thus a colorimetrically undetectable extent of corrosion over the duration assayed).

Interestingly, all SRB conditions appeared to be passivating at most time points, as compared to the SRB-free medium control. This is likely the case because, despite our attempts to create a truly anaerobic environment, minute concentrations of oxygen below our detection limit (1000 ppm) can facilitate slow oxidative corrosion. In the absence of fluid flow, the biofilms thus appeared to limit the oxygen exposure to the metal surface. The corrosion rate of the control environment was only 0.03 mm/year, or 20- to 30-fold slower than that observed in aerobic aqueous saline conditions. The maximum average measured corrosion rate of the MIC environments was 0.028 mm/year, occurring for conditions that promoted high SRB metabolic activity. In other words, for these specific SRB and environments, MIC included biofilm formation and was slower than abiotic corrosion for low-oxygen environments, and was significantly slower than aerobic corrosion of the same metals in saline environments.

While it is possible that the conductivity of the biofilm itself could mask corrosion-induced resistance increases, our additional control experiments indicated this unlikely to be a compounding factor. SRB biofilms grown on glass capillaries in ferrous ion-containing medium demonstrated that while the biofilm could be conductive, the electrical resistance was still orders of magnitude higher than that of the steel wires used in our experiments (data not shown). Furthermore, growth of ferrous ion-containing biofilms on platinum wires (a metal with extremely low corrosion susceptibility) did not detectably increase measured conductivity (data not shown).

### 3.5. Advantages and limitations

The use of wire in corrosion assays provides several advantages over existing approaches that make it amenable to rapid, high-throughput corrosion analysis. First, wire forms can be adapted easily to almost any conformation, allowing use in confined environments such as 96 well plates and microfluidic channels. Second, the small sample volumes allow rapid assessment of corrosion when compared to traditional studies conducted on larger (cm- to m-scale) samples over the course of months, without the need to scale up metal manufacturing or sourcing of the fluid (such as that from produced water of potential and current oil wells). In addition, wires can be imaged easily, even providing an in situ

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**Fig. 5.** Biofilm formation on wires in anaerobic conditions. Imaging biofilm-coated wires while in liquid provides a pseudo cross-sectional view allowing the biofilm corrosion/product to be readily observed in comparison to a control wire in same conditions without SRBs (initial wire diameters 117 µm). Light and scanning electron microscopy of the dried wire allows further assessment of biofilm and underlying metal morphology. Scale bars 50 µm for optical microscopies and 10 µm (1500 ×) and 5 µm (5500 ×) for the upper and lower electron microscopy images, respectively.

**Fig. 6.** Corrosion in anaerobic aqueous environments comprising sulfate reducing bacteria (SRBs). The estimated radius change for wires in indicated conditions of differing nutrient availability (see Fig. S4 for measurement error) along with colorimetric data at the indicated time points for the same conditions. Data shown as estimated radius change, and not resistance change, to normalize for small differences in initial wire diameter among wells.
pseudo cross-sectional view of the corrosion process. In other words, this approach allows rapid, systematic identification of the least/most susceptible metal composition or microstructure or of the least/most corrosive fluid composition. Such capabilities are well-suited to corrosion susceptibility screening.

The current platform also has certain constraints. Each well contains a static environment with no continuous fluid exchange. While liquid can be exchanged manually through a small hole above each well without interrupting the experiment, corrosion in environments with continuous fluid flow cannot be studied. This also leads to saturation of the colorimetric signal as corrosion products build up and settle to the well floor. Because each well is not sealed completely, evaporation can occur over several days and gas exchange is possible between wells. Future designs incorporating pumps could allow for controlled fluid and gas flow, as well as controlled removal of gaseous byproducts of corrosion or biotic metabolism. Because test materials must be fabricated in wire format, some materials may be precluded from use with this system. Further, resistance measurements alone cannot easily distinguish between uniform and pitting corrosion, although samples can be sacrificed at desired time points to visually assess corrosion modes. Finally, it is important to note that the estimated corrosion rate is sensitive to pitting, as the resistance is modulated by the areas of smallest cross-section.

4. Conclusions

A novel high-throughput approach for the rapid assessment of corrosion susceptibility has been achieved for up to 96 wire samples analyzed in parallel, through the use of resistance and colorimetric measures. The capabilities of this multiplexed approach were demonstrated by assessing the effect of NaCl concentration on aerobic, abiotic corrosion and of nutrient availability on anaerobic, biotic corrosion for a low carbon steel. This platform could be used as a screening mechanism with conditions of interest studied further via other techniques.

These multiplexed electrical/colorimetric analyses are also well suited to studies that leverage directed mutagenesis of bacteria. Such studies could provide insights into the mechanisms by which microbes enhance corrosion or by which antimicrobial reagents work most effectively, but require 100s–1000 s of independent experiments. Thus, to consider corrosive environments of increasing complexity, the rapid and quantitative measurements afforded by this multi-well format can expedite discovery of corrosion susceptibility and also of corrosion mitigation options.

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Appendix A. Supplementary material

Supplementary data associated with this article can be found, in the online version, at http://dx.doi.org/10.1016/j.corsci.2014.07.045.

References