Title
Electrochemical polarization and impedance of reinforced concrete and hybrid fiber-reinforced concrete under cracked matrix conditions

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1. Introduction

The electrochemical phenomenon of corrosion is a global source of deterioration of reinforced concrete structures. In environments lacking chloride ions (chlorides), low-carbon steel reinforcing bars (rebar) embedded within concrete form a passive film due to the high pH of the concrete pore solution and are generally not expected to exhibit any significant corrosion activity during the designed service life of a structure. However, external chlorides, which originate from sources such as ocean saltwater or deicing salts placed on roadways, can permeate porous concrete cover layers and depassivate the steel rebar after a critical threshold of local chloride concentration has been reached at the rebar surface [1,2]. After depassivation, corrosion activity of the steel rebar readily increases. As the volume of corrosion products is greater than the sum of the reactant volumes, internal expansion of these products causes concrete cracking [3], subsequently increasing the chloride solution permeability of the concrete [4,5] and reducing the mechanical performance of the composite [6,7]. Premature replacement of damaged structures and associated mass consumption of additional construction materials are undesirable due in part to the high energy demands required for concrete production [8].

Because reinforced concrete degradation is dependent on the cracked state of the composite matrix, research of novel crack-resistant construction materials has gained popularity in recent decades [9,10]. In particular, hybrid fiber-reinforced concrete (HyFRC) is a candidate material to reduce corrosion-induced cracking damage due to the mechanical toughening provided by the inclusion of different types of short (e.g., 8-mm to 60-mm long), discontinuous fibers dispersed throughout its cementitious matrix [11,12]. A crack-resistant concrete such as HyFRC is further advantageous considering that a primary functional purpose of reinforced concrete is to resist mechanical loads, requiring any newly implemented construction material to be damage-tolerant against not only corrosion-induced cracking but also structural loading [13,14]. An increase in the time to corrosion initiation was observed for reinforced HyFRC (i.e., HyFRC composite with embedded steel rebar) compared to reinforced concrete after subjecting samples to flexural stress [15,16]. While a lower corrosion current density $i_{corr}$ of reinforced HyFRC was also measured compared to reinforced concrete, indicating favorable durability performance after active corrosion had begun, electrochemical characteristics of samples
were limited to corrosion potential, polarization resistance, and corrosion current density based on assumed Stern-Geary coefficients. Several corrosion-related studies with different types of fiber-reinforced concrete have utilized similar techniques [17–20], making detailed information from certain other electrochemical tests, such as Tafel polarization and electrochemical impedance spectroscopy (EIS), rarely available for fiber-reinforced concrete composites.

As mentioned, \( i_{corr} \) is an experimental parameter that is often of interest when evaluating the corrosion behavior of reinforced concrete, and may be calculated using the Stern-Geary equation if the Stern-Geary coefficient \( B \) and the polarization resistance \( R_p \) are known [21]. For corrosion concrete studies, a number of researchers have relied on the assumed values of \( B = 52 \text{ mV} \) and \( 26 \text{ mV} \) for passive and active steel reinforcement, respectively, referencing the paper of Andrade and González [22] published in 1978. The analysis of this paper considered a comparison between gravimetric steel mass loss and electrochemical mass loss, which was based on time-integration of \( i_{corr} \) measurements, for validation of the proposed \( B \) values. While the authors did not report values for the cathodic (\( \beta_c \)) or anodic (\( \beta_a \)) Tafel coefficients, which are typically required for determination of \( B \) (Eq. (1)), Chang et al. [23] noted that \( B = 52 \text{ mV} \) may be obtained by setting \( \beta_c = 120 \text{ mV/decade} \) and allowing \( \beta_a \) to reach infinity. Similarly, \( B = 26 \text{ mV} \) may be obtained by letting both Tafel coefficients be equivalent to 120 mV/decade. Though not mentioned, the assignment of \( \beta_c = 120 \text{ mV/decade} \) may be attained by evaluating the definition of the anodic Tafel slope (Eq. (2)) under standard temperature \( T \) and setting the product \( \alpha_c n = 0.5 \), where \( \alpha_c \) is the anodic charge transfer coefficient and \( n \) is the number of transferred electrons involved in a rate-determining step. Furthermore, \( R \) is the universal gas constant and \( F \) is the Faraday constant. A similar determination of \( \beta_c = 120 \text{ mV/decade} \) may be performed by allowing \( \alpha_c n = 0.5 \), where \( \alpha_c \) is the cathodic charge transfer coefficient. When assuming \( n = 1 \), an electrochemical cell with \( \alpha_c = 0.5 \) and \( \alpha_c = 0.5 \) is possible, though the charge transfer coefficients should not be arbitrarily assigned to such values [24].

\[
B = \frac{\beta_c \beta_a}{2.303 (\beta_c + \beta_a)} \quad (1)
\]

\[
\beta_a = \frac{2.303RT}{\alpha_a n F} \quad (2)
\]

Elsewhere in the literature, reported values of \( \beta_c, \beta_a, \) and \( B \) have generally been inconsistent, as surveyed in Table 1. Using a variety of specimen types and exposure durations, several authors have found the cathodic Tafel coefficient in actively corroding samples to be in the approximate range of 100–250 mV/decade [25–28], as well as in the higher range of 300–450 mV/decade [27,29,30]. Anodic Tafel coefficients of corroding steel have shown more scatter, ranging from 200 mV/decade to values approaching infinity [25–27,29,30]. While \( B \) values for active corrosion have been determined in the range of 7–15 mV [31], greater values of 43–86 mV have also been reported [25,27].

Cracking damage of samples, if any, was not reported in several polarization studies surveyed in Table 1 despite experimental durations reaching or exceeding 2 years [27–30]. Babaee and Castel [25] noted no corrosion-induced cracks were observed around the embedded steel rebar at the conclusion of experiments, while Grubb et al. [31] observed only microcracking in plain mortars but no cracking in fiber-reinforced mortars. Subramaniam and Bi [26] performed flexural cracking in reinforced concrete beam samples prior to chloride exposure to determine the effects of localized transverse cracking on microcell and macraccole corrosion rates, though they did not consider the effects of corrosion-induced cracking on polarization behavior.

EIS has become a common monitoring technique for reinforced concrete corrosion research in recent decades [32,33], though few researchers have considered the influence of matrix cracking on the electrochemical impedance of corroding steel reinforcement. Feng et al. [34] subjected reinforced concrete beams to varied magnitudes of flexural stress prior to chloride exposure. The authors found that the concrete cover resistance, when modeled as an

<table>
<thead>
<tr>
<th>Reference</th>
<th>Binder type*</th>
<th>Total exposure duration</th>
<th>Concrete cracking in samples</th>
<th>Passive</th>
<th>Active</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>( \beta_c ) (mV/dec.)</td>
<td>( \beta_a ) (mV/dec.)</td>
</tr>
<tr>
<td>Babae and Castel</td>
<td>FA, ultra-fine FA, GGBFS, NaOH solution</td>
<td>0.9 years(^b)</td>
<td>No cracks found</td>
<td>30–45</td>
<td>Infinity</td>
</tr>
<tr>
<td>Chang et al. [27]</td>
<td>C, FA, GGBFS</td>
<td>4.7 years</td>
<td>Not reported</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Alonso et al. [29]</td>
<td>OPC</td>
<td>6 years</td>
<td>Not reported</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Al-Tayyib and Khan</td>
<td>OPC</td>
<td>2 years</td>
<td>Not reported</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Subramaniam and Bi</td>
<td>OPC</td>
<td>0.5 years</td>
<td>Crack induced by mechanical loading prior to chloride exposure</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Glass and Chadwick</td>
<td>OPC</td>
<td>2.5 years(^b)</td>
<td>Not reported</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Andrade and González</td>
<td>OPC</td>
<td>Slag cement Pozzolanic cement</td>
<td>Not reported</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Grubb et al. [31]</td>
<td>OPC</td>
<td>0.4 years(^b)</td>
<td>Microcracks found in mortar; No microcracks found in fiber-reinforced mortar</td>
<td>–</td>
<td>–</td>
</tr>
</tbody>
</table>

\(^a\) Binder type abbreviations: OPC - ordinary portland cement; FA - fly ash; GGBFS - ground granulated blast-furnace slag; C - unspecified cement.

\(^b\) Estimated duration in years.
element within an equivalent electrical circuit, was not highly sensitive to the magnitude of applied flexural stress when surface concrete cracks could not be visually detected after flexural loading. However, greater applied stresses tended to decrease the circuit element value representing polarization resistance due to increased rebar-matrix interfacial damage. Similar effects of applied load-induced interfacial damage on polarization resistance [35] and charge transfer resistance [36] have been reported when using an equivalent circuit to model the electrochemical impedance spectra of reinforced concrete. While EIS can provide useful information on the effect of matrix cracking on the corrosion behavior of reinforced concrete, application of this technique on crack-resistant fiber-reinforced concrete composites remains scarcely reported in the literature. Grubb et al. [31] performed EIS on reinforced fiber-reinforced mortars subjected to chloride solution intrusion and determined that the presence of steel fibers influenced the measured impedance spectra due to the electrical conductivity of the fibers, though the authors did not provide results on equivalent circuit modeling.

Due to the aforementioned influence of cracking on the durability of reinforced concrete, a study detailing the electrochemical behavior of reinforced concrete with cracking damage was warranted to better elucidate the processes that occur while embedded steel rebar are corroding. The research presented herein investigates the long-term influence of concrete cracking on the electrochemical behavior of reinforced concrete and reinforced HyFRC in chloride environments. Prismatic samples each containing a single strand of steel reinforcing bar were subjected to external NaCl solution intrusion conditions for approximately 2.5 years, during which corrosion potential measurements, linear polarization resistance tests, and Tafel polarization tests were periodically performed to monitor changes in corrosion activity. EIS was performed at the conclusion of the environmental conditioning period to evaluate changes in the impedance responses due to accumulated matrix cracking damage. To simulate the matrix cracking that may occur while reinforced concrete structures are in service, samples were subjected to an applied tensile load prior to, and during, exposure to the corrosion-inducing conditions.

2. Materials and methods

2.1. Materials and specimen

A hybrid fiber-reinforced concrete (HyFRC) developed by Blunt and Ostertag [12] was selected for study, and its batch proportions are presented in Table 2. The total fiber volume fraction, based on composite volume, was 1.5%. Fiber hybridization consisted of a blend of 0.04-mm diameter, 8-mm long polyvinyl alcohol (PVA) fibers; 0.55-mm diameter, 30-mm long hooked-end steel fibers; and 0.75-mm diameter, 60-mm long hooked-end steel fibers. The chemical compositions of the steel fibers, as well as of the ASTM type I/II [37] portland cement used in the mixtures, are summarized in Table 3. A combination of inductively coupled plasma optical emission spectroscopy (ICP-OES), combustion, colorimetric tests, and gravimetric tests was performed on the steel fibers, while X-ray fluorescence (XRF) was performed on the cement. For the XRF testing, SO₃ could not be quantified due to its volatility during high temperature fusion. The HyFRC was evaluated against a plain concrete mixture that had the same cement content and water-cement mass ratio as HyFRC but lacked fiber reinforcement. After casting of the fresh mixtures, concrete and HyFRC specimens were cured within covered molds for 7 days, demolded, and cured an additional 21 days in ambient laboratory conditions (approximately 23.5 °C, 45% relative humidity) prior to tensile loading, if applicable. The curing regime was selected to better represent reinforced concrete construction in professional practice, where formwork is often removed a few days after casting. When formwork is stripped, the concrete is exposed to ambient conditions. Compression testing of cylinders with a 100-mm diameter and 200-mm height revealed that the concrete and HyFRC had similar 28-day compressive strengths of 42.6 MPa and 44.2 MPa, respectively.

Geometric detailing of the specimen design is presented in Fig. 1a. The rectangular prisms had a length of 610 mm and a square cross section with side lengths of 127 mm. A single ASTM A706 [38] steel reinforcing bar with a nominal yield stress of 410 MPa and a nominal diameter of 16 mm was placed centrally within the specimen’s cross section. The alloy composition of the steel is summarized in Table 3 and was reported by a mill certificate provided upon receipt of the rebar from its commercial supplier. Prior to sample casting, the mill scale of the rebar was removed by sandblasting. The middle 406-mm length of rebar was defined to be the region of interest where corrosion was allowed to occur, and was in direct contact with the cementitious matrix during casting. Elsewhere, the reinforcing bar was coated with an electrically-insulating lacquer and then tightly wrapped with vinyl insulating tape to prevent electrical and physical contact with concrete. The extruded portions of the steel bar were threaded to allow for later tensile loading. A loading frame consisting primarily of light-gauge perforated tube steel was constructed such that specimens would remain in a stressed state while the frame would resist reactionary forces (Fig. 1c).

Ponding of 3.5% w/w NaCl solution through the concrete cover

### Table 2

Batch proportions per cubic meter of composite.

<table>
<thead>
<tr>
<th>Composite</th>
<th>Portland cement (kg)</th>
<th>Aggregates (kg)</th>
<th>Water (kg)</th>
<th>Fibers [% vol. composite]</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Coarse</td>
<td>Fine</td>
<td></td>
</tr>
<tr>
<td>Concrete</td>
<td>423</td>
<td>742</td>
<td>872</td>
<td>228</td>
</tr>
<tr>
<td>HyFRC</td>
<td>423</td>
<td>775</td>
<td>853</td>
<td>228</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.2</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.5</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.8</td>
</tr>
</tbody>
</table>

### Table 3

Chemical composition of select reinforced concrete materials, expressed in mass percentage of material.

<table>
<thead>
<tr>
<th>Type I/II portland cement</th>
<th>CaO</th>
<th>SiO₂</th>
<th>MgO</th>
<th>Al₂O₃</th>
<th>Fe₂O₃</th>
<th>K₂O</th>
<th>Na₂O</th>
<th>TiO₂</th>
<th>P₂O₅</th>
<th>MnO</th>
</tr>
</thead>
<tbody>
<tr>
<td>A706 rebar steel</td>
<td>0.29</td>
<td>0.17</td>
<td>0.010</td>
<td>0.041</td>
<td>0.030</td>
<td>0.06</td>
<td>0.86</td>
<td>0.08</td>
<td>0.28</td>
<td>0.004</td>
</tr>
<tr>
<td>Fiber (30-mm length) steel</td>
<td>0.062</td>
<td>&lt;0.005</td>
<td>0.04</td>
<td>0.011</td>
<td>0.006</td>
<td>&lt;0.005</td>
<td>0.005</td>
<td>0.30</td>
<td>0.05</td>
<td>0.08</td>
</tr>
<tr>
<td>Fiber (60-mm length) steel</td>
<td>0.074</td>
<td>&lt;0.005</td>
<td>0.15</td>
<td>0.015</td>
<td>0.007</td>
<td>&lt;0.005</td>
<td>0.005</td>
<td>0.32</td>
<td>0.05</td>
<td>0.09</td>
</tr>
</tbody>
</table>
was achieved by fixing a polyvinyl chloride ponding dam to a horizontal face of each of the prismatic specimens using a rubber cement adhesive, as in Fig. 1b. All vertical concrete surfaces, along with the portion of the ponded specimen face not enclosed by the adhered dam, were coated with a moisture-resistant sealer to prevent leakage of solution during experimentation. As summarized in Table 4, a total of 4 specimen types were studied, varying in composite type (concrete (C) or HyFRC) and loading state during environmental conditioning (nonloaded (0) or loaded (L)).

2.2. Tensile loading

Select specimens of each composite type were prescribed for tensile loading during chloride exposure to account for the loading of reinforced concrete that occurs in practice, which includes permanent gravity loading from the weight of supported structures as well as temporary loading such as vehicle traffic or seismic events. These loads may cause tensile concrete cracking, which is allowed under United States codes when the element is properly designed [39]. In this study, samples were loaded such that concrete cracking occurred while the steel rebar remained elastic. Based on the tensile testing results reported by Moreno et al. [40], who tested prismatic reinforced concrete and reinforced HyFRC samples with the same cross-sectional geometry and rebar size as the samples described in Fig. 1, a load magnitude of 53 kN was expected to achieve the desired deformation characteristics.

The setup for loading is shown in Fig. 2a while a schematic of the loading procedure is presented in Fig. 2b. To load the specimens, a hydraulic pump was used to longitudinally displace one end of the steel rebar, resulting in a tensile stress and strain response along the specimen. The other, opposite end of the extruded rebar was anchored and fixed against a steel plate. After a load of 53 kN was reached, a nut located near the bearing plate where the hydraulic pump was positioned was tightened against the plate, transferring the point of force application on the rebar from the hydraulic pump to the nut. To prevent any effects of galvanic coupling between the extruded steel reinforcing bar and the steel loading frame, the frame’s bearing plates were coated with an electrically-insulating spray solution and covered with vinyl insulating tape where they were to be in direct contact with the extruded rebar or bearing nut.

2.3. Environmental conditioning

To saturate the concrete after curing in ambient laboratory conditions, specimens were externally ponded with 3.5% w/w NaCl solution for 13 consecutive weeks. Afterwards, the ponding dam was completely relieved of solution for 2 weeks and then refilled with fresh solution for 2 subsequent weeks. These steps were repeated in a cyclic manner to simulate field conditions where sodium chloride solution ingress into reinforced concrete is not a continuously occurring event. After 29 total weeks of environmental conditioning, the frequency of the cycles was increased such that specimens were under drying conditions for 1 week, followed by wetting for 1 week. The total experimental corrosion monitoring time of the samples was 132 weeks, or approximately 2.5 years.

2.4. Electrochemical measurements

Corrosion activity measurements were taken periodically with a potentiostat under ambient laboratory temperature throughout the environmental conditioning period, with all measurements occurring while the specimen was in a ponded state. The potentiostat was equipped with a positive feedback iR compensation feature that corrected for the concrete cover resistance. All tests utilized a three-electrode setup, as shown in Fig. 3, with the working electrode being the steel reinforcing bar of the specimen, the counter electrode being stainless steel, and the reference electrode being a saturated calomel electrode (SCE). Stainless steel was selected as the counter electrode material based on its reported use for reinforced concrete research [41,42]. The counter electrode and the reference electrode were immersed in a 3.5% w/w NaCl solution contained by the ponding dam. During the initial 13 weeks of environmental conditioning, measurements were taken at intervals no more than 2 weeks apart. Afterwards, wet-dry cycling of

Table 4

Specimen type designation.

<table>
<thead>
<tr>
<th>Reinforced composite</th>
<th>Nonloaded</th>
<th>Loaded</th>
</tr>
</thead>
<tbody>
<tr>
<td>Reinforced concrete</td>
<td>C-0</td>
<td>C-L</td>
</tr>
<tr>
<td>Reinforced HyFRC</td>
<td>HyFRC-0</td>
<td>HyFRC-L</td>
</tr>
</tbody>
</table>

Fig. 1. Details of specimen design: (a) Geometric drawing; (b) Plan view of specimen with ponding dam fixed to concrete surface; (c) Specimen in loading frame.
samples occurred and the measurement frequency was relaxed to approximately 4-week intervals. Once 100 total weeks of environmental conditioning had elapsed, measurement sessions were further relaxed to every 6 to 10 weeks.

In a typical sample-testing session, measurement of the corrosion potential $E_{corr}$ was first performed with a duration of at least 10 minutes, during which $E_{corr}$ was not to deviate by more than 1 mV to ensure quasi-steady state conditions. Afterwards, linear polarization resistance testing, considering a potential domain of $-10 \text{ mV}$ to $10 \text{ mV}$ versus $E_{corr}$, was performed to obtain the polarization resistance $R_p$. The corrosion potential was again measured to ensure quasi-steady state conditions prior to conducting Tafel polarization tests, which polarized samples from a potential of $75 \text{ mV}$ below the corrosion potential to a maximum value of $75 \text{ mV}$ above the corrosion potential at a potential sweep rate of 0.1 mV/s. In the literature, others have utilized a magnitude of overpotential that was greater than or equivalent to $120 \text{ mV}$ [25–27,29,30]. An overpotential value of $75 \text{ mV}$ was selected for this study to minimize possible damage introduced by large applications of overpotential while being sufficiently great to allow for extraction of cathodic $\beta_c$ and anodic $\beta_a$ Tafel coefficients. These coefficients were determined by curve fitting of quasi-linear regions of the Tafel polarization response. After calculating the Stern-Geary coefficient $B$ through Eq. (1), the corrosion current $I_{corr}$ can be obtained through Eq. (3):

$$I_{corr} = \frac{B}{R_p}$$

Electrochemical impedance spectroscopy (EIS) was conducted at the conclusion of the environmental conditioning period. The frequency domain for testing was defined to be from $10 \text{ mHz}$ to $100 \text{ kHz}$ and the amplitude of the perturbation potential was $10 \text{ mV}$.

### 2.5. Destructive testing

After conclusion of the environmental conditioning phase, specimens were subjected to destructive tests to evaluate chemical, electrochemical, and physical changes caused by chloride intrusion and corrosion activity. After removing the ponding dams, cylindrical cores with a diameter of 29 mm were extracted from select samples using a coring drill press. For loaded samples, one core was taken directly over the applied load-induced macrocrack and a secondary core was taken 76 mm from the crack, as shown in Fig. 4. Otherwise, a core was taken near the midlength of nonloaded samples. Coring initiated at the ponding surface and continued into the depth of the specimens, with the core drill positioned such that it would not make contact with the embedded reinforcing bar. The extracted concrete cores were cut into segments, each with a height of 25 mm, and pulverized such that the final concrete material could pass a sieve with a maximum opening of 0.84 mm. The experimental testing standard of ASTM C1152 [43], utilizing potentiometric titration, was performed on the pulverized samples to determine their acid-soluble chloride content, which is expected to be equivalent to the total (i.e., bound and free) amount of chlorides in the cementitious matrix. Because the test standard is specific to cement-based materials, steel fibers from HyFRC cores were manually removed after pulverizing to avoid possible artifacts introduced by oxidation of the steel fibers in acid solution. After an equivalence point of titration was determined, the chloride content was calculated as a percentage of the pulverized concrete mass used in the titration test. For HyFRC samples, this pulverized concrete mass did not include the mass of steel fibers.

Remaining specimens not subjected to coring were prepared for optical cross-sectional imaging by sawing the samples to produce a prismatic element with the steel rebar located at the approximate center of the prism, as schematically shown in Fig. 5. The prismatic element was then sawed into plates approximately 13 mm thick. In the figure, hatched (i.e., shaded) portions of the extracted prism and plate indicate where the imaging surface was located. Prior to sawing, all appreciable surface cracks were impregnated with a blue-colored, low-viscosity epoxy to maintain existing crack widths and minimize any damage caused by the cutting process.
3. Results

3.1. Observed damage

Typical surface cracking of reinforced concrete and reinforced HyFRC, after application of an applied tensile load and prior to chloride exposure, is shown in Fig. 6. Tensile macrocracks occurred near specimens’ midlength with an approximate maximum crack width of 0.42 mm and 0.22 mm for concrete and HyFRC, respectively. For the same magnitude of applied tensile load, a smaller maximum crack width in HyFRC is expected as a result of the crack control provided by hybrid fiber reinforcement [40].

Final accumulated damage of reinforced concrete and reinforced HyFRC at the conclusion of the 132-week environmental conditioning duration is shown in Figs. 7 and 8, respectively. The images were obtained from an extracted prism taken from a sample of each specimen type, as previously detailed in Fig. 5. Binary images (Fig. 7a and f, Fig. 8a and f) show a vertical surface of the extracted prism. Observed macrocracks, shown in white, are contrasted with the matrix, shown in black. The middepth of the images may be slightly offset from the actual centerline of the longitudinal steel rebar embedded within the specimens. Colored lines superimposed on the binary images indicate the location of presented transverse section cuts. For instance, the transverse section cuts of sample C-0 found in Fig. 7b and c are located where the green lines occur in Fig. 7a. In all transverse section images, surface voids and cracks are colored blue due to the impregnation of the samples with an epoxy containing a blue pigment. Finally, the transverse section images are positioned such that the centroid of the rebar cross section nearly coincides with the center of the presented image.

Differences in the cracking severity between concrete and HyFRC are noticeable. Matrix damage of reinforced concrete samples, regardless of applied loading state, was primarily characterized by splitting crack propagation over a considerable length of the specimens (Fig. 7a and f). Splitting cracks are identified in a longitudinal view as cracks that form along the length of a sample, and in a transverse view as cracks that emanate radially from the rebar. The presence of splitting cracks at a rebar surface generally coincided with a corroded surface (Fig. 7b–e), though it is unclear which event (i.e., matrix splitting crack propagation or rebar corrosion) preceded the other for a particular transverse section. Regardless, the widespread propagation of these cracks allowed a free path for external deleterious substances, such as NaCl solution and gaseous O₂ and CO₂, to reach the rebar surface far from the site of initial rebar corrosion and to promote depassivation or further corrosion activity.

In a nonloaded HyFRC-0 sample, transverse or splitting cracks could not be visually detected and rebar corrosion damage appeared negligible (Fig. 8a–c). While under applied load, splitting cracks formed within the HyFRC-L matrix, though the crack opening and propagation length (Fig. 8d–f) were significantly less severe compared to reinforced concrete samples. Splitting cracks could not propagate to the exterior of the sample due to the crack resistance of the HyFRC cover, as no surface splitting cracks were observed. Where the splitting crack widths were fine, as in Fig. 8d, steel mass loss was marginal, indicating that rebar corrosion activity was generally limited to the site of the induced tensile crack near the section shown in Fig. 8e.
Steel fibers embedded within chloride-contaminated HyFRC were observed to have insignificant corrosion damage, except those fibers located very near a matrix crack or located at the ponding surface itself. Similar observations of the greater chloride threshold required to corrode steel fibers as opposed to conventional steel rebar have been summarized elsewhere in the literature [10,44].

3.2. Tafel polarization

Typical Tafel polarization curve fitting results for a passive and an active specimen are presented in Fig. 9, showing the fitted cathodic and anodic polarization curves and the net current produced by the two polarization curves. The potential where net current is naught ($E_{\text{cell}}$) and the corrosion current ($I_{\text{corr,T.}}$) can be extracted from the intersection of the fitted cathodic and anodic polarization curves. The subscript in $I_{\text{corr,T.}}$ indicates the result is based directly on Tafel (T.) polarization curves as shown. Generally, the most negative potentials of the raw data were ignored for fitting due to the uncharacteristically high current that occurred at these overpotentials, an experimental observation that was also noted by Chang et al. [23]. Compared to active behavior, passive behavior is distinguished by greater values of potential, lower values of current, and an anodic portion of the curve exhibiting a low-angle slope as plotted. The anodic polarization curve slope is a consequence of the near-potential independence of current in the passive regime.

Cathodic and anodic Tafel coefficients $\beta_\text{c}$ and $\beta_\text{an}$, respectively, are plotted in Fig. 10 with $E_{\text{cell}}$ labeled as the abscissa, while Table 5 summarizes the mean and standard deviation of $\beta_\text{c}$, $\beta_\text{an}$, and the Stern-Geary coefficient $B$. Two distinct clusters of data may be observed for each subplot, generally separated by corrosion potentials greater than $-100$ mV vs. SCE and less than $-300$ mV vs. SCE. The former set of measurements, totaling 71 data points, is associated with passivated samples while the latter set, totaling 156 data points, is associated with active samples. Samples not satisfying either criterion were not considered for reported statistics.

The cathodic Tafel coefficient $\beta_\text{c}$ differed between the passive regime ($\beta_\text{c} = 19.8$ mV/dec) and the active regime ($\beta_\text{c} = 102$ mV/dec) for steel rebar. Babaei and Castel [25] reported a similar discrepancy in $\beta_\text{c}$ values between passive and active steel rebar embedded within geopolymer concrete. Changes in $\beta_\text{c}$ for the same metal could be attributed to different electrochemical reduction reactions that occur at different corrosion potentials. The reduction of dissolved oxygen within basic concrete pore solution, as shown in reaction (4), is often assumed to be the reduction reaction involved in steel rebar corrosion. At lower corrosion potentials, the reduction of hydrogen ions (i.e., $H_2O$, as indicated in reaction (5)), could also be considered. Using the Nernst equation (Eq. (6)) at room temperature ($25^\circ$C), the equilibrium reduction potential $E$ may be calculated for a given reduction reaction.

$$\begin{align*}
\frac{1}{2}O_2 + H_2O + 2e^- &\rightarrow 2(OH)^- \\
2H_2O + 2e^- &\rightarrow H_2 + 2(OH)^- \\
E &= E^0 - \frac{0.0592}{z} \log(Q)
\end{align*}$$

In Eq. (6), $E^0$ is the standard reduction potential, $z$ is the number of electrons transferred in the cell reaction, and $Q$ is the reaction
differences in measured cathodic Tafel coefficients at varied corrosion potentials.

For active samples, a mean $B = 28.1$ mV was found and is similar to the well-cited value of $B = 26$ mV proposed by Andrade and González [22] for reinforced concrete materials. Fig. 11 plots the correlation between the corrosion current $I_{corr, S.-G.}$ based on the Stern-Geary equation, which incorporates $B$ directly, and the corrosion current $I_{corr,T}$. Found by extrapolation of fitted Tafel polarization responses. The dashed line in the plots is where equivalent measurements of $I_{corr, S.-G.}$ and $I_{corr,T}$ occur. For active samples, the corrosion currents from both methods of derivation are similar in magnitude and the mean values show reasonably good correlation (13.7% error). For passive samples exhibiting lower values of corrosion current, being approximately less than 10 μA based on either determination method, use of the Stern-Geary equation resulted in an underestimation of corrosion current compared to $I_{corr,T}$. When minimizing the residual between the mean values obtained from each method, an empirical Stern-Geary coefficient $B = 24.0$ mV was found to be more appropriate for passivated samples.

3.3. Corrosion rate monitoring

The averaged corrosion currents $I_{corr}$ of specimens are plotted as a function of elapsed environmental conditioning time $t$ for the initial 12 weeks of experimentation and for the entire duration in Fig. 12a and b, respectively. The corrosion current measurements are based on Eq. (3) and directly incorporate the Stern-Geary coefficient $B$. For presumably passive samples, the empirical value $B = 24.0$ mV was used for calculation. Otherwise, $B$ was determined directly from a sample’s experimental Tafel polarization response at a given time. In reinforced concrete research, the corrosion current density $I_{corr}$ is often reported in the literature and is typically determined by normalizing $I_{corr}$ by the nominal surface area of the rebar in contact with cementitious matrix, as the actual corroding surface area of an embedded rebar is generally not known without the use of advanced materials characterization techniques. Considering the nonuniform corrosion activity of samples in this study, as visualized in Figs. 7 and 8, communication of the corrosion current density normalized by a constant active surface area may result in inaccurate interpretations of the actual corrosion current density of samples.

Initial active corrosion of a sample was detected by an increase in $I_{corr}$ on the order of one magnitude between consecutive measurements. Among all specimen types, loaded reinforced concrete samples (C-L) exhibited the shortest time to corrosion initiation, requiring approximately 1.0 week to become active. After 5.0 weeks, corrosion initiation was detected in loaded HyFRC-L, which was delayed compared to C-L due to the smaller crack widths produced in HyFRC upon loading. Nonloaded reinforced concrete samples (C-0) first exhibited active corrosion between 43.1 and 63.3 weeks. Compared to C-0 samples, the time to corrosion initiation for any nonloaded HyFRC sample (HyFRC-0) was at least 41.7 weeks longer and at least a factor of 1.7 longer.

The time to corrosion initiation was influenced by the diffusion rate of chlorides through the cementitious matrix. Fig. 13a and b plot the chloride content $C_{Cl^-}$ of reinforced concrete and reinforced HyFRC, respectively, as a function of depth $d$ from the ponding surface of the specimens. The chloride content is the mass of chlorides normalized by concrete mass in a tested sample. For HyFRC samples, steel fibers were excluded from the testing mass, as previously mentioned in Section 2.5. At the depth to the centroid of the steel reinforcing bar ($d = 63.5$ mm), the chloride content of C-L was 0.38% wt. concrete at the cracked location and 0.32% at a noncracked location, highlighting the faster rate of chloride ingress...
where a crack was present and correlating with the shorter time to corrosion initiation of cracked samples as compared to noncracked samples. For HyFRC specimens not containing a macrocrack, the chloride contents were found to be in the range of 0.07–0.11% wt. HyFRC excluding steel fibers at the depth where the reinforcing bar was located, reduced from the chloride content found in non-cracked conventional concrete at the same depth by a factor of at least 2.9. Because the concrete and HyFRC were designed with the same cement weight content and same water-cement mass ratio, differences in chloride binding from cement were likely negligible between composite types. The reduction in chloride penetration could then be attributed to the presence of fiber reinforcement in sound matrices. The inclusion of PVA or steel fibers has been experimentally observed to retard the rate of chloride ingress in cementitious composites [45–47]. In the absence of macrocracks, hybrid fiber reinforcement is an effective corrosion initiation retardant, though the precise influence of fibers on chloride diffusion requires further investigation.

3.4. Electrochemical impedance spectroscopy

Selection of an appropriate equivalent electrical circuit that models the potentiostatic EIS results of a metal considers three parallel paths: (1) the impedance associated with the oxidation reaction; (2) the impedance associated with the reduction reaction; and (3) the impedance associated with the capacitance of the electrode-electrolyte solution interface. The oxidation reactions result in pitting corrosion and passive film growth. While reduction reactions can take place within a pit, only the reduction of oxygen on the steel surface outside a pit is considered herein. The impedance related to pitting corrosion was reported by Wenger et al. [48] to short-circuit the impedance associated with passive film growth and reduction of oxygen.

A circuit containing two time constants, as shown in Fig. 14a, was suitable for modeling the behavior of conventional reinforced concrete and is similar to the equivalent circuit described by others of steel reinforcing bars undergoing corrosion within reinforced
Constant-phase elements (CPE) were used in the equivalent circuit to account for the nonideal behavior of capacitor elements. The impedance $Z_{CPE}$ of a CPE is described by Eq. (7):

$$Z_{CPE} = \frac{1}{Y_0(j\omega)^\alpha} \quad (7)$$

$Y_0$ is the admittance modulus of an ideal capacitor, $j$ is the imaginary unit, $\omega$ is the angular frequency, and $\alpha$ is an exponent ranging from 0 to 1. A value of 1 for $\alpha$ describes an ideal capacitor, a value of 0.5 describes an infinite Warburg diffusion-controlled element, and a value of 0 describes a resistor. At the base of corrosion pits, the element designated as $CPE_{DL}$ represents the double-layer capacitance while the element $R_{CT}$ accounts for the charge transfer resistance of the oxidation reaction. These two parallel elements are connected in series to the resistor $R_{so}$, which accounts for the resistance related to ion transport through the electrolyte solution in corrosion pits. The remaining constant phase element, $CPE_{pass}$, relates to the capacitance of the double-layer at the interface between the passive film and concrete pore solution and the capacitance of the passive film. Capacitances related to $CPE_{pass}$ occur in series, with the double-layer capacitance likely greater than that of the passive film, resulting in the total capacitance $CPE_{pass}$ approximated as the double-layer capacitance. Finally, the ohmic resistance of the concrete solution is represented by the element $R_{soln}$ and is attributed to solution-filled pores and cracks in the matrix.

After unsatisfactory fitting of data from HyFRC samples to the described equivalent circuit, a constant phase element ($CPE_{fiber}$) was placed parallel to $R_{soln}$, as shown in Fig. 14b, to account for steel fiber-matrix interfaces that occur for an arrangement of continuously connected fibers. Within the cementitious matrix, fibers are in contact with each other due to their nonuniform spatial distribution resulting from concrete mixing and casting [52]. Any resistance from the concrete pore solution between the imperfect contact of steel fibers was assumed to be negligible relative to $R_{soln}$ which considers a relatively large concrete cover (56 mm) compared to the small distances between nearly contacting steel fibers in the matrix.

Bode plots comparing the fitting results of the impedance

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**Fig. 8.** Sectional images of reinforced HyFRC specimens at the conclusion of environmental conditioning: (a)–(c) HyFRC-0; (d)–(f) HyFRC-L. Colored vertical lines on binary images indicate locations of indicated transverse section cuts. Binary images are used to highlight crack locations on a vertical surface of the extracted prism (as in Fig. 5). A blue-colored epoxy was used to impregnate cracks and voids in transverse section views. (For interpretation of the references to color in this figure caption, the reader is referred to the Web version of this article.)
modulus $|Z|$ and negative phase shift $-\phi$ to acquired EIS data are presented in Fig. 15a and b, respectively. EIS measurements were taken at the conclusion of the environmental conditioning period, and thus consider the influence of matrix cracking, if any. Good correlation is observed between the experimental data and the model fit. At intermediate to high testing frequencies $f$ (i.e., $f$ greater than approximately 1 Hz–10 Hz), the impedance modulus of reinforced HyFRC samples decreases more rapidly with increasing frequency compared to that of reinforced concrete samples. Grubb et al. [31] observed a similar disparity between the impedance modulus spectra of steel rebar embedded within steel fiber-reinforced mortar and plain mortar. Nyquist plots relating the negative imaginary part $-Z_{im}$ of the complex impedance to the real part $Z_{re}$ of the complex impedance are presented in Fig. 15c and d at different plot scales. In these plots, a lower value of $Z_{re}$ is associated with a greater testing frequency. Compared to the responses of C-0 and C-L, as shown in Fig. 15c, the HyFRC-L response contains a characteristic minimum located at low values of $Z_{re}$, or conversely, high tested frequency $f$. This minimum at high tested frequency is indicative of steel fiber inclusion in the HyFRC matrix, and is also present in the response of HyFRC-0. Similar effects of steel fibers on Nyquist plot responses of concrete materials have been reported by others [53–55], though these authors did not consider the impedance spectra with a steel rebar embedded within concrete. Passivity of the rebar within the HyFRC-0 sample dominated its impedance response, resulting in a considerably different response compared to HyFRC-L at lower tested frequencies, as exhibited in Fig. 15d.

Values of the parameters used in the equivalent circuit modeling, in addition to corrosion current, are summarized in Table 6 for considered specimens. The corrosion currents were determined in the same manner as described in Section 3.3. Similar to the reporting of corrosion current rather than corrosion current density, the EIS parameter values are not scaled by the area affected by pitting corrosion due to this area not being accurately known during electrochemical testing. The ohmic concrete resistance $R_{soln}$, determined to be 3.07–6.93 $\Omega$ for reinforced concrete, was greater for HyFRC-L (28.8 $\Omega$) and was one order of magnitude greater for HyFRC-0. Due to the presence of longitudinal splitting cracks that reached the specimen surface in both reinforced concrete specimens, a direct path with greater void volume saturated by solution was available from the steel rebar to the surface of the concrete. Cracks within the HyFRC-L matrix were finer compared to those in reinforced concrete, resulting in a greater electrolyte solution resistance for the HyFRC sample. Lower charge transfer resistances $R_{CT}$ were noted for samples with greater $I_{corr}$. While $R_{CT}$ of loaded samples were similar in magnitude to the nonloaded reinforced concrete sample C-0, the charge transfer resistance of HyFRC-0 was four orders of magnitude greater than the other considered specimens due to the oxidation of this specific specimen attributed almost entirely to passive film growth and to marginal amounts of pitting corrosion. As previously visualized in Figs. 7 and 8, noticeable pitting corrosion was found for the HyFRC-L, C-0, and C-L sample types, with greater corrosion damage observed for greater matrix cracking severity. Similarly, the $R_{pit}$ value for HyFRC-0 was greater than that of other considered samples.
Circuit element values for the constant phase elements $CPE_{DL}$ and $CPE_{pass}$ differed between sample types. For reinforced concrete and reinforced HyFRC samples, the exponent term $\alpha$ for $CPE_{DL}$ was determined to be 0.297 to 0.336 and 0.632 to 0.807, respectively. The values of $\alpha$ were low (i.e., on the order of 0.3) for the reinforced concrete samples. Similarly low values were reported by Dhouibi-Hachani et al. [56] for reinforced concrete samples immersed in solutions containing chloride, sulfate, or both. The authors found that the value of $\alpha$ correlated with the nonuniform arrangement of corrosion pits as well as the heterogeneity of corrosion products on the steel bar surface and in the concrete pores adjacent to the steel bar surface, with lower $\alpha$ associated with greater levels of inhomogeneity. Similar remarks concerning the relationship between $\alpha$ and surface heterogeneity have been reported in the literature [57,58]. Lower $\alpha$ values were noted for reinforced concrete samples compared to those of reinforced HyFRC due to more extensive pitting corrosion on the steel rebar surface.

The $\alpha$ values for $CPE_{pass}$ in reinforced HyFRC were determined to be 0.502 to 0.512. As $\alpha = 0.5$ describes an infinite Warburg element, it is suggested that $CPE_{pass}$ is a diffusion-controlled element for the HyFRC samples. The passive film of steel in simulated concrete pore solution was modeled by Sánchez et al. [59] as a Warburg element representing the diffusion of oxygen vacancies through the passive film. Compared to noncracked conventional concrete sections, noncracked HyFRC sections were found to exhibit a relatively low chloride content in the matrix pore solution at the depth of the reinforcing bar, as detailed in Fig. 13. It is likely that the HyFRC generally maintained an insignificantly damaged passive film at cathodic surfaces, while the higher chloride content in the pore solution of conventional reinforced concrete caused damage to the film, resulting in lower $\alpha$ values (0.306–0.325).

Fitting of $CPE_{fibre}$, which accounted for the impedance contribution of embedded steel fibers in the HyFRC matrix, resulted in an exponent term $\alpha$ in the range of 0.493–0.508. The steel fibers in the matrix are also best described as a Warburg diffusion element due to their porous nature.

### Table 5

<table>
<thead>
<tr>
<th></th>
<th>Passive</th>
<th></th>
<th></th>
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<th></th>
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<tr>
<td></td>
<td>$\beta_\alpha$ (mV/dec)</td>
<td>$\beta_\beta$ (mV/dec)</td>
<td>$\beta_\gamma$ (mV)</td>
<td>$\beta_\delta$ (mV/dec)</td>
<td>$\beta_\epsilon$ (mV/dec)</td>
<td>$\beta_\zeta$ (mV)</td>
<td>$\beta_\eta$ (mV/dec)</td>
<td>$\beta_\theta$ (mV)</td>
</tr>
<tr>
<td>Mean value (standard deviation)</td>
<td>19.8 (4.26)</td>
<td>562 (201)</td>
<td>8.31 (1.81)</td>
<td>102 (9.74)</td>
<td>180 (40.4)</td>
<td>28.1 (2.83)</td>
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</table>

* Empirical mean value based on fitting of corrosion current values from Stern-Geary equation to extrapolated corrosion current values from Tafel polarization.
to $\alpha$ being nearly equivalent to the value of 0.5, suggesting that the bulk of fibers should be in a passivated state. As previously mentioned in Section 3.1, steel fibers embedded within sound HyFRC did not appear to exhibit corrosion damage, indicating the fitting results coincided with visual observations.

4. Discussion

4.1. Early-age corrosion propagation

Changes in the Tafel polarization responses of select samples during the initial 40 weeks of active corrosion are presented in Fig. 16. The indicated time in the plots is the elapsed active corrosion time, which is considered to be 0 weeks upon first detection of corrosion initiation. HyFRC-0 did not produce sufficient active corrosion measurements during the experiment to consider for appropriate comparison.

The early-age corrosion propagation characteristics were influenced by composite type and presence of matrix cracks. For the nonloaded sample C-0, insignificant changes are observed for the polarization curves at 6 and 20 weeks. Later, a shift in the polarization response at 40 weeks is characterized by a large current increase in the anodic reaction curve, as highlighted by the vertical arrow in Fig. 16a. By comparison, the cathodic reaction curve is only slightly shifted to lesser values of potential. An increase in anodic current may be caused by an increase in chloride concentration of the local solution at the rebar surface \cite{60-63}, an increase in the actively corroding surface area, or both. Cracking of the concrete matrix would allow both events to occur, as splitting cracks with sufficient width increase the chloride solution permeability of the concrete cover \cite{4,5} and subsequently cause passivated surfaces within the splitting crack wake to become active. The large shift of the anodic curve results in an increase in corrosion current from 20 to 40 weeks, as highlighted in Fig. 16d. While loaded specimen C-L exhibits a greater corrosion current at 6 weeks compared to C-0, the polarization response of C-L does not significantly change from 6 to 40 weeks. Because corrosion products preferentially form within defects (i.e., cracks) located near the steel rebar surface \cite{64,65}, the corrosion rate of C-L is initially higher due to products more readily forming within empty crack openings than in a denser rebar-matrix interface. Corrosion-induced matrix cracking occurs when the accumulation of expansive corrosion products fills a void volume and places sufficient stresses on the matrix, an event which likely did not occur until after 40 weeks of active corrosion. Due to a lack of additional matrix cracking, the corrosion rate does not increase as rapidly compared to that of C-0 despite being initially greater.

Although HyFRC-L was also exposed to a chloride environment while in a cracked and loaded condition, its corrosion current was lower compared to C-L due to the fineness of the cracks in the HyFRC
matrix, resulting in a smaller initial anodic surface where the crack-rebar interface was located. A decrease in HyFRC-L corrosion potential occurs at 40 weeks and the corrosion rate (107 μA) is slightly reduced from the measurement obtained at 20 weeks (113 μA) despite an increase in the anodic reaction curve to greater current. Lowering of the corrosion rate is caused in part by shifting of the equilibrium potential $E$ of the cathodic reaction to a more negative value. Based on the Nernst equation and the oxygen reduction reaction previously presented in \(4\), a change in the equilibrium potential of the cathodic reaction may be anticipated based on changes in solution chemistry as described by Eq. (8):

$$E = E^0 - \frac{0.0592}{2} \log \left( \frac{[\text{OH}^-]^2}{p_{\text{O}_2}^{1/2}} \right)$$  \(8\)

<p>| Element values for equivalent electrical circuit modeling. |
|----------------|----------------|----------------|----------------|----------------|----------------|</p>
<table>
<thead>
<tr>
<th></th>
<th>$I_{corr}$</th>
<th>$R_{so} \Omega$</th>
<th>$R_{ct} \Omega$</th>
<th>CPE$_{Pt}$-Y$_0$ mS $s^\alpha$</th>
<th>CPE$_{Pt}$-α</th>
<th>CPE$_{Pt}$-Y$_0$ mS $s^\alpha$</th>
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<th>CPE$_{F}$-Y$_0$ mS $s^\alpha$</th>
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<td>33.6</td>
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<td>41.0</td>
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<td>0.306</td>
<td>–</td>
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<td>63.3</td>
<td>81.3</td>
<td>0.297</td>
<td>56.7</td>
<td>0.056</td>
<td>0.325</td>
<td>–</td>
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<td>HyFRC-0</td>
<td>526</td>
<td>74.1</td>
<td>3.07 $\times$ 10$^4$</td>
<td>12.2</td>
<td>0.807</td>
<td>431</td>
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<td>0.512</td>
<td>5.86 $\times$ 10$^{-3}$</td>
</tr>
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<td>HyFRC-L</td>
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<td>28.8</td>
<td>77.7</td>
<td>73.3</td>
<td>0.632</td>
<td>61.5</td>
<td>2.58</td>
<td>0.502</td>
<td>5.67 $\times$ 10$^{-3}$</td>
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</tbody>
</table>

Fig. 15. Plots showing EIS results and fitted results: (a) Bode plot with impedance modulus $|Z|$ versus frequency $f$; (b) Bode plot with negative phase shift $-\phi$ versus frequency $f$; (c) Nyquist plots with the negative imaginary part $-Z_{im}$ of the complex impedance versus the real part $Z_{re}$ of the complex impedance. EIS measurements were performed at the conclusion of the environmental conditioning period.
$E^0$ is the standard reduction potential, $z$ is the number of electrons transferred in the cell reaction, $[\text{OH}^-]$ is the concentration of OH$^-$ in solution, and $P_{O_2}$ is the partial pressure of dissolved O$_2$ in solution. The equilibrium potential lowers for increasing concentration of hydroxide or decreasing partial pressure of dissolved oxygen. The former change in chemistry is unlikely due to the inherently high pH of concrete pore solution (approximately 13 to 13.5) and a tendency for pH to decrease due to carbonation or calcium hydroxide leaching of the cementitious matrix. A lowering of oxygen partial pressure is more likely and may be caused by several phenomena, including the formation of corrosion product scales on the steel rebar surface, which impede the diffusion of oxygen through the scale layers [66,67], and a greater consumption of oxygen in electrode reactions than replenishment [68,69], which leaves the oxygen partial pressure lowered at the time of measurement.

Fig. 16. Tafel polarization responses of select specimens over elapsed active corrosion time, up to 40 weeks: (a) C-0, additionally showing influence of factors on polarization curvature; (b) C-L; (c) HyFRC-L; (d) Superposition of $\log I_{corr} - E$ relations from (a)–(c). Numbers adjacent to polarization curves indicate time, in weeks, of elapsed active corrosion.

4.2. Late-age corrosion propagation

Tafel polarization diagrams showing responses for up to 90 weeks of corrosion activity for C-0 and up to 130 weeks of corrosion activity for C-L and HyFRC-L are presented in Fig. 17. C-0 is characterized by the greatest increases in corrosion current as corrosion propagation time increases, followed by C-L and then HyFRC-L. Greater corrosion current was correlated with greater splitting crack damage, which was previously mentioned to increase the permeability of the cementitious matrix and allow for the ingress of chlorides and oxygen to the steel rebar surface through cracks. For C-0 and C-L, the maximum surface splitting crack widths were measured to be 0.32 mm and 0.14 mm, respectively. While HyFRC-L exhibited local splitting crack damage near actively corroding rebar surfaces, as previously shown in Fig. 8, no surface splitting cracks could be visually detected on the exterior surface of any HyFRC sample. Correspondingly, the corrosion rate of HyFRC-L shows small increases from 40 to 70 weeks though does not significantly
Evaluation of the fitted cathodic and anodic reaction curves from polarization responses taken at 90 weeks of active corrosion, as shown in Fig. 18, highlight the dominating effect of the anodic response on $I_{\text{corr}, T}$ compared to the cathodic response. Due in majority to the increased anodic current, the corrosion currents of C-0 and C-L are greater than the corrosion current of HyFRC-L, designated as $I_{\text{corr}, T\text{(HyFRC-L)}}$, by factors of 5.78 and 2.21, respectively. Hybrid fiber reinforcement reduces the total corrosion current of reinforced concrete materials over long-term conditions due to its effective crack control, which restricts pitting corrosion of large surfaces of steel rebar far from the initial site of corrosion.

5. Summary and conclusions

The corrosion behavior of conventional reinforced concrete and reinforced hybrid fiber-reinforced concrete (HyFRC) was found to be highly dependent on the cracked state of the composite matrix. A corrosive environment was prepared by allowing 3.5% w/w NaCl solution to permeate the porous cementitious matrix of samples containing a single steel reinforcing bar (rebar) in a wet-dry cyclic manner over a 2.5-year experimental duration. To account for cracks that are present in civil engineering structures in service, subsets of specimens were subjected to the same applied tensile load during environmental exposure. Corrosion potential measurements, linear polarization resistance tests, and Tafel polarization tests were periodically conducted to monitor the electrochemical response of samples during the experiment. In addition, electrochemical impedance spectroscopy (EIS) was performed to evaluate the differences in the impedance responses between samples as a result of accumulated matrix cracking. The following conclusions are made:

1. HyFRC composites require a longer time to initiate steel rebar corrosion compared to conventional reinforced concrete. While placed under tensile stress and subsequently cracked, the delay is attributed to the crack control of fiber reinforcement restricting the maximum opening of induced cracks, which slows the rate of external NaCl solution penetration. For non-loaded samples, the inclusion of polyvinyl alcohol (PVA) and...


steel fibers reduces the effective chloride ion diffusion rate through the matrix and lengthens the time to achieve active corrosion by a factor of at least 1.7, though the detailed mechanism of retardation requires additional study.

2. The cathodic and anodic Tafel coefficients for actively corroding samples, regardless of composite or loading condition, were found to have mean values of $\beta_c = 102 \text{ mV/decade}$ and $\beta_a = 180 \text{ mV/decade}$, respectively. A mean Stern-Geary coefficient of $B = 28.1 \text{ mV}$ was also calculated, similar to the well-cited value of $B = 26 \text{ mV}$ for active corrosion of reinforced concrete proposed by Andrade and González [22]. For passive samples, use of $B$ based on fitted Tafel coefficients resulted in an underestimation of the corrosion current compared to the corrosion current extrapolated directly from Tafel polarization curves. In this case, an adjusted Stern-Geary coefficient of $B = 24.0 \text{ mV}$ was used for passive samples based on empirical fitting.

3. At the conclusion of testing, matrix cracking damage was most severe for nonloaded reinforced concrete (C-0). In order of descending observed damage, loaded reinforced concrete (C-L), loaded reinforced HyFRC (HyFRC-L), and finally nonloaded reinforced HyFRC (HyFRC-0) followed. Upon initial active corrosion, stress accumulation and subsequent cracking from expansive corrosion product formation occur more rapidly for C-0 due to the formation of such products in a dense rebar-matrix interface rather than within a crack void. HyFRC was significantly more effective than conventional concrete at restricting the opening and propagation of corrosion-induced cracks.

4. Correlating with the most severe cracking damage among sample types, C-0 was associated with the lowest charge transfer resistance and lowest ohmic concrete pore solution resistance, as well as the highest corrosion current. Correspondingly, the charge transfer and pore solution resistances were greater and the corrosion current was lower for HyFRC-L compared to reinforced concrete samples due to less severe matrix cracking.

5. The exponent term $\alpha$ for constant phase elements of equivalent circuits for EIS modeling had lower magnitude for greater pitting corrosion severity. Considering the constant phase element representing the interfacial capacitance and passive film of steel rebar (CPFpass), $\alpha$ was determined to be approximately 0.5 for reinforced HyFRC samples, indicating that the element acted as a diffusion-controlled infinite Warburg element that was not significantly damaged. The result correlates well with low measured chloride ion contents in the matrix of noncracked HyFRC sections.

6. While the analysis of Tafel polarization curves indicates that the partial pressure of reducible oxygen in the concrete pore solution decreased over time, which would otherwise lower the corrosion rate if no other changes occurred, increasing corrosion currents occurred for reinforced concrete samples during the testing duration while active reinforced HyFRC did not significantly change in corrosion rate for the final 40 weeks of experimentation. Increases in corrosion current were primarily attributed to the formation of additional active steel rebar surfaces beyond the initial site of corrosion, which was made possible by the formation of matrix splitting cracks along the length of the rebar and was more widespread for reinforced concrete. Control of the anodic current response is regarded as being critical for preventing increasing corrosion rates in reinforced concrete composites undergoing wet-dry cycling.

Acknowledgments

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