Electrochemical Chloride Extraction: Influence of Concrete Surface on Treatment

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FOREWORD

Many now recognize that once chloride-induced corrosion of the reinforcing steel bars has initiated in a concrete bridge, the only truly effective means of stopping corrosion in the structure is by applying either cathodic protection or the relatively new electrochemical chloride extraction. In addition to providing some beneficial effects to a treated concrete bridge, electrochemical chloride extraction offers some advantages over the use of cathodic protection. Electrochemical chloride extraction extends the functional life of the treated structure, but does not require the long-term commitment generally required of cathodic protection systems. However, both cathodic protection and electrochemical chloride extraction are operated without requiring the excavation of structurally sound concrete that is contaminated with chlorides. This can provide an advantage that does not exist with other restoration techniques.

To facilitate wider application of electrochemical chloride extraction in the rehabilitation of concrete bridges, this investigation was initiated with the objectives of improving the effectiveness of the treatment and contributing to the determination of additional service life that results from treating a structure. This interim report describes the progress made in ascertaining the cause of the abrupt drop in the amount of current that can pass through salt-contaminated concrete. This reduction in current, typically observed during the first several days of treatment, in turn relates to a decrease in chloride removal.

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16. Abstract

One bridge restoration technique available for reducing corrosion-induced concrete deterioration, which removes chloride ions while simultaneously realkalizing the concrete adjacent to the steel, is electrochemical chloride extraction (ECE). Studies have shown that ECE is capable of removing, in a single application, a significant portion of the chloride ions from a reinforced concrete structure. Prior research has also shown that the quantity of chloride ions removed is dependent on numerous factors including quantity and spacing of reinforcing steel, applied voltage, initial chloride concentration, etc. In addition, investigations into chloride binding and competition between other ions as current carriers have helped to clarify the probable mechanisms responsible for decreases in current efficiency with time during chloride removal.

This portion of the investigation has focused on the influence of water-to-cement (w/c) ratio. In addition, an investigation was conducted to identify the cause of decrease in efficiency during chloride removal. A clear relationship between the w/c ratio and the chloride extraction rate was not evident. However, the investigation revealed that the resistance of the concrete surface layer increases considerably during ECE, which effectively restricts the current flow, while the resistance of the underlying layer of concrete either decreases or remains constant. It appears that the increased resistance of the surface layer concrete is accompanied by the formation of a tightly adhering residue on the concrete surface. Preliminary analysis of the surface formation indicates it contains calcium carbonate and calcium chloride.

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INTRODUCTION, LITERATURE REVIEW, AND PURPOSE

Introduction

This report contains typical scientific abbreviations.

It is known that chlorides can lead to corrosion in reinforced concrete structures. A second detrimental factor then transpires because the corrosion product requires a larger volume of space than the original iron. This creates tensile stresses, which makes the concrete more prone to cracking and spalling. If this process continues, premature deterioration of a bridge can result. Therefore, in chloride contaminated bridges it is vital to have a method for alleviating the corrosive attack on the reinforcing steel. Currently, two electrochemical methods are available, cathodic protection (CP) and electrochemical chloride extraction (ECE), each having certain advantages and disadvantages. Although ECE is a proven means of removing chlorides from the concrete while increasing the pH in the region adjacent to the reinforcing steel, extensive use has not developed partially due to a deficient understanding in the following issues:

- The cause of the decrease in current flow and, therefore, the rate of chloride extraction over the duration of a treatment.
- The influence of initial parameters, such as concrete properties and initial chloride concentration, on the required duration of chloride extraction for a concrete structure.
- An estimation of the additional service life following an application.

Interest in the movement of ions through concrete has prompted numerous studies, both theoretical and experimental. Currently, data from field and laboratory experiments indicate certain regions in concrete appear to lead to inefficient chloride extraction. By determining the regions of low efficiency and the controlling mechanisms, questions relating to extraction rates, efficiency, and beneficial life can be addressed. It is anticipated that upon the completion of this project, techniques for altering the procedure and/or materials will provide a means to improve the ECE process.

Literature Review

Corrosion Threshold

Despite differing opinions on corrosion mechanisms and ion ingress into concrete, the concentration and movement of oxygen, chloride, and hydroxyl ions are all considered important factors in the corrosion of reinforcing steel. However, chloride ions are considered the key factor with regard to the corrosion. Li, et al., presented the flow chart shown in figure 1, which is based on the work from several studies that investigated the relationship between chloride concentration and reinforcing steel corrosion. ^[1] This diagram emphasizes the numerous factors that could influence the corrosion threshold value. ^[1]

Early research into the corrosion of reinforcing steel indicated that a change in alkalinity near the reinforcement can significantly influence the steels' susceptibility to corrosive attack. [2] According to Hausmann, the chloride threshold value, which is given in equation 1, is a function of the chloride and hydroxyl ion concentrations ratio and should not exceed 0.61. [2]

$$\frac{\left[Cl^{-}\right]}{\left[OH^{-}\right]} \le 0.61\tag{1}$$

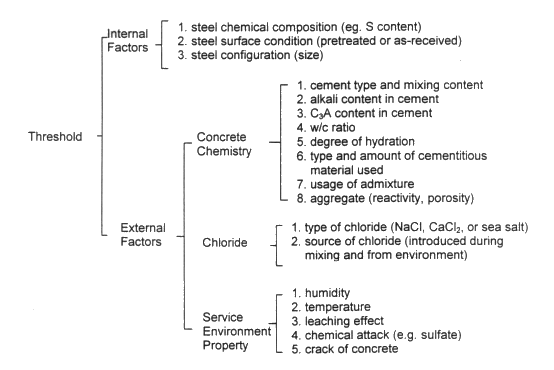


Figure 1. Potential influences on the corrosion threshold for steels exposed to chlorides [1]

Based on the relationship in equation 1, it has been suggested that concrete could have a higher percentage of chlorides if they were introduced during the mixing stage, due to binding of some of the chloride, versus if chlorides diffuse into a cured concrete block. Other researchers have produced estimates for the chloride threshold value that range from 75 - 3640 ppm in concrete. The effects of several other factors on the corrosion threshold value are summarized in Table 1.

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Table I	Hactors int	liiencing c	orrosion	threshold	value ^[3, 5-12]
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Description	Effect on Threshold Value
Dehydration of concrete	Increases
Water saturation resulting in oxygen depletion	Increases
Concrete sealed or pores constricted	Increases
Increase in humidity	Decreases
Changing concrete mixture (i.e. admixtures,	
w/c ratio)	Increases or Decreases

Electrochemical Chloride Extraction

The concept of removing chloride ions from concrete by electrochemical migration was borne in 1973 out of Kansas Department of Transportation (KDOT) experiments on electro-stabilization of clayey

soils. [13] Since then, numerous studies have shown it is possible to remove chlorides from concrete using electrochemical means. [14-17] To facilitate the treatment of vertical surfaces a commercial electrochemical method was developed, which is known as Norcure [TM, [18]] The benefit of removing the chloride ions electrochemically is that contaminated concrete that are still structurally sound would not require excavation and will remain in place after the application of the chloride removal process. This restoration technique has inherited various names; electrochemical chloride removal, desalination, and electrochemical chloride extraction.

Electrochemical Chloride Extraction vs. Cathodic Protection

ECE and CP have some distinct similarities and differences. Although both are DC techniques that cathodically polarize the reinforcing steel and reduce the corrosion rate, CP is usually permanently installed, operates at lower current densities (approximately 10 mA/m^2), and usually requires routine maintenance. Figure 2 is an illustration of a typical CP system installed on a bridge deck with the anode permanently embedded in the concrete overlay. In contrast, ECE is an in-situ restoration technique that is designed to remove chlorides and increase the alkalinity adjacent to the reinforcing steel. A temporary treatment system is attached to the concrete and the applied voltage causes a direct current, which can be up to 1 A/m^2 , to flow through the concrete for typically 4 to 8 weeks. In the ECE system is then removed following completion of the treatment process. Currently, ECE of bridge decks, using a system such as that illustrated in figure 3, requires changes to the traffic pattern during operation. However, ECE of concrete bridge piers do not generally require the rerouting of traffic.

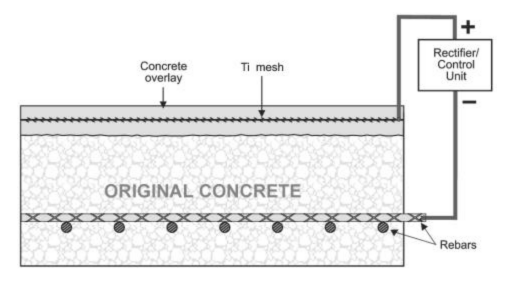


Figure 2. Cathodic protection system for reinforced concrete

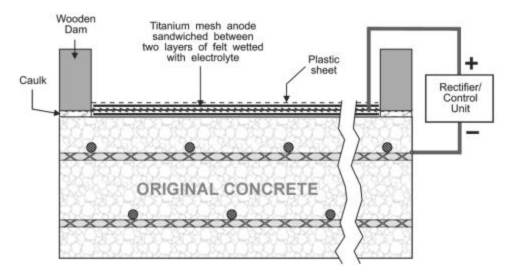


Figure 3. Illustration of ECE setup on the 34th Street Bridge in Arlington, Virginia, USA [17]

Both electrochemical techniques produce hydroxyl ions at the cathode or the rebars, while water is decomposed at the anode. As shown in the following reactions, hydrogen gas can be produced at the cathode, and chlorine evolution and/or acidification of the electrolyte can occur at the anode during ECE. Two possible reactions at the cathode during ECE are: [16]

$$2H_2O + O_2 + 4e^- \rightarrow 4OH^- \tag{2}$$

$$H_2O + e^- \rightarrow H^0_{ADS} + OH^-$$
 (3)

At the anode, ECE can generate the following reactions: [16]

$$2H_2O \rightarrow 4e^- + O_2 + 4H^+$$
 (4)

$$2Cl^{-} \rightarrow 2e^{-} + Cl_{2} \tag{5}$$

In addition, the following chemical reaction can occur in the electrolyte: [16]

$$H_2O + Cl_2 \rightarrow HCl + HClO$$
 (6)

As hydroxyl ions are produced at the cathode the pH adjacent to the steel increases, which is beneficial for the rebar, but these electrochemical reactions can create adverse effects like hydrogen embrittlement or alkali aggregate reaction. In equation 3, " H^{o}_{ADS} " is nascent hydrogen, which could either enter the metal or form hydrogen gas. These issues are discussed further in the section "Effects of Electrochemical Chloride Extraction" on page nine.

Material Requirements for Electrochemical Chloride Extraction

Figure 3 is an illustration of an ECE system using a catalyzed titanium mesh anode, however other anodes have performed satisfactorily. [14, 16, 17, 19-25] Steel mesh anodes cost less than inert catalyzed titanium mesh anode, but some of the steel is consumed during the extraction process and therefore it has a shorter functional life. [20, 26] In addition, some have suggested that the corrosion product from the steel anode can deposit in concrete pores and decrease the chloride extraction efficiency if ECE is being applied on an upward facing horizontal surface. [25]

Potable water has shown favorable results as an electrolyte during ECE. [22] Calcium hydroxide solutions have also been used as an electrolyte in many applications. A benefit of using calcium hydroxide is it reduces the chance of the electrolyte becoming acidic and etching the concrete when a catalyzed titanium anode is used. [17,26] In addition, increasing the alkalinity of the electrolyte reduces the evolution of chlorine gas. [25] A third common electrolyte is lithium borate solution, which is useful when dealing with concrete containing aggregates susceptible to alkali-aggregate or alkali-silica reaction (ASR). [22, 26, 27] The lithium borate solution is actually a mixture of lithium hydroxide and boric acid, which ensures lithium ions are available to penetrate the concrete and reduce or eliminate ASR. [3, 26] This solution is the most expensive of the commonly used electrolytes. [26] Although the electrolytes presently used for ECE do not contain corrosion inhibitors, Asaro, et al., demonstrated in a strategic highway research program (SHRP) study that it is possible to inject inhibitors using a similar setup as ECE. [28]

During ECE it is important to maintain good contact between the electrolyte and the concrete surface to minimize circuit resistance. This has been accomplished using three different methods: sprayed cellulose fiber, synthetic felt mats, and surface-mounted tanks. For vertical surfaces, sprayed cellulose fiber and surface-mounted tanks are generally used. When treating horizontal surfaces, synthetic felt mats are more common. [22, 26]

Conductivity and Electrochemical Chloride Extraction

Work by Christensen, et al., suggested that the binding of ionic species and the increase in alkalinity during the hydration process has a strong influence on the conductivity. [29] During the early stages of hydration in a chloride free cement (≤ 100 hr.), conductivity is dominated by Na⁺, K⁺, Ca²⁺, OH, and SO₄-², [29] As the hydration process continues, Christensen, et al., found that only Na⁺, K⁺, and OH contribute significantly to the conductivity in these samples. [29] With equation 7 and using the ionic conductivity values given in table 2, Banfill calculated the transference values for a mixture containing 0.5 mol/liter sodium hydroxide and 0.5 mol/liter sodium chloride, which are given in table 3.^[30] Based on these values, it was concluded that the current flow from the reinforcing steel toward the anode was composed of 72% hydroxyl ions and 28% chloride ions during electromigration.^[30]

$$t_{j} = \frac{I_{j}}{I_{total}} = \frac{\left|z_{j}\right|c_{j}I_{j}}{\sum \left|z\right|cI} = \frac{FW_{j}}{35.5I_{total}t}$$

$$(7)$$

Where.

= Transference number of species j

= Current due to species j

 $I_{total} = Total current$ $z_j = Charge on specification is the second current of the$ = Charge on species j

= Concentration of species j

Ionic conductivity of j (infinite dilution)

Faraday constant

Mass of species i removed

Table 2. Ionic conductivity values [30]

Positive Ions	Conductivity (ohm ⁻¹ cm ⁻² eq ⁻¹)	Negative Ions	Conductivity (ohm ⁻¹ cm ⁻² eq ⁻¹)
H^{+}	349	OH.	198
Na ⁺	50.1	Cl	75.2
K ⁺	73.5	½ SO ₄ ²⁻	79.8
Ca ²⁺	59.5	½ CO ₃ ²⁻	69.3
		HCO ₃	44.5

Table 3. Calculated transference values for a solution containing 0.5 mol/l NaCl and 0.5 mol/l NaOH [30]

Species	Value
t_{Na+}	0.27
t _{OH-}	0.53
t_{Cl}	0.20

The calculation by Banfill assumed that for every 96,500 coulombs of charge passed, one mole or 35.5 g of chloride ions successfully migrated to the anode. However, after calculating the efficiency of an early SHRP study (6.9-7.8%), Banfill concluded that other negative ions (i.e. OH^- , SO_4^{-2}) and resistive heat generation must account for the efficiency loss. Tritthart demonstrated that the concentration of hydroxyl ions in concrete increased during ECE, and suggested that during ECE the hydroxyl ion influences the rate of removal of the chloride ion. This is because as hydroxyl ions are being produced at the cathode and migrating towards the anode during ECE, these ions will compete with chloride ions as charge carriers. Therefore, based on the calculations by Banfill and the results from Tritthart, it is not surprising that the efficiency of chloride removal would decrease as the treatment progressed. $^{[30,32]}$

Chatterji suggested that although free chloride ions can participate in electrolysis, bound chloride ions would first require exchange with hydroxyl ions before being able to contribute to the conductivity. For the exchange between bound chloride ions and hydroxyl ions to occur, Chatterji proposed that the electrolysis would require sufficient time and an elevated hydroxyl ion concentration. The fact that not all of the chlorides are initially free to migrate could reduce the extraction efficiency of the process. In the work by Elsener, et al., indications of a chemical equilibrium between bound and free chlorides were shown. Others have supported this idea and have even included this as a factor in proposed models. In the kwaba, et al., suggested the following chemical equation would exist for a chloroaluminate compound,

$$CaCl2 + 3CaO·Al2O3 + 10H2O? 3CaO·CaCl2·Al2O3·10H2O$$
(8)

All these studies have led to the development of some general relationships between the various factors and ECE. Table 4 summarizes these relationships, which were determined either experimentally or through modeling. However, cracks in the concrete are not addressed in this table. This is because it was determined that the effect of small cracks (≤ 0.5 mm wide) did not significantly affect the current distribution. Bennett, et al., suggested that even though small cracks appeared to fill with a white precipitate during ECE, it is best if damaged concrete is repaired prior to ECE. In addition, Bennett, et al., indicated that variations in the depth of cover did not greatly influence the current distribution during ECE.

Table 4. Influences of various factors on ECE [31, 37-41]

Factor	Effect
Increase quantity of reinforcing steel	Increases chloride extraction rate
Increase the applied voltage	Increases chloride extraction rate
Higher initial chloride concentrations	Increases chloride extraction rate
Reinforcement mats placed directly over each other **	Increases chloride extraction rate
Increasing temperature above 35°C	Increases chloride extraction rate
Multiple applications of ECE	Increases chloride extraction efficiency
Initial chloride concentration on final chloride concentration	No influence
Potentiostatic vs. galvanostatic operation	No influence
Renewal of anolyte to maintain maximum concentration gradient	No influence
Carbonated layer in front of chlorides being extracted	Decreases chloride extraction rate

^{**} Based on mathematical model

Electrochemical Chloride Extraction Projects

Various studies have demonstrated ECE is a promising bridge restoration alternative to CP for chloride-contaminated concrete bridges. [8,14,16,17,20-23,26,42,43] Table 5 lists the reinforced concrete structures in North America that were treated using ECE. The majority of these structures can be categorized as either bridge piers or decks. In addition, this table includes a summary of the percentage of chlorides removed from selected North American structures using ECE treatment. Unfortunately, many of these results are not reported for the same concrete depths, so a comparison between structures is impossible. However, it is clear that although the amount of chlorides removed from most of these structures was substantial, some of the chlorides remain in the structure following treatment. [3, 19]

Half-cell measurements on some of these treated structures following ECE treatment indicate a low probability of corrosion. Table 6 lists the half-cell measurements that were taken on treated and untreated structures. In addition, all of the structures listed in table 6 are also cited in table 5.

Table 5. ECE treatment on selected North American structures [17,21,31,42-55]

Location	Date	Area Treated	Chloride Removed (%)	Current Efficiency (%)
Hwy #192 Bridge Substructure, Council Bluffs, Iowa		1209 m ²		N/A
Highway 11 Bridge Abutments, North Bay, Ontario	2000	646 m ²	N/A	N/A
Eastern Avenue Bridge #576 Abutments, Washington DC	2000	220 m ²	N/A	N/A
3 rd St. Viaduct, Bridge Substructure, Minot, North Dakota	1999	100 m ²	N/A	N/A
St. Adolphe Bridge Deck, St. Adolphe, Manitoba	1999	14704 m ²	N/A	N/A
S02 of 38061 Substructure, Jackson County, Michigan	1999	109 m ²	N/A	N/A
I-480 Bridge Substructure, Omaha, Nebraska	1999	1400 m ²	N/A	N/A
Burlington Skyway Substructure, Burlington, Ontario	1999	1533 m ²	N/A	N/A
Hwy #192 Bridge Substructure, Council Bluffs, Iowa	1998	463 m ²	N/A	N/A
I-480 Bridge Substructure, Omaha, Nebraska	1998	1525 m ²	74 (at 0-25 mm) 63 (at 50-75 mm)	N/A
St. Adolphe Bridge Deck, St. Adolphe, Manitoba	1998	1115 m ²	84 (at 0-25 mm) 70 (at 25-51 mm)	N/A
Pembina Highway Overpass Structure, Winnipeg, Manitoba	1998	220 m ²		N/A
Industrial Spur Bridge Substructure, Peoria, Illinois	1998	462 m^2	N/A	N/A
Starbuck Bridge Deck, Winnipeg, Manitoba	1997	270 m^2	N/A	N/A
I-395 & Dunwoody Substructure, Minneapolis, Minnesota	1997	225 m^2	N/A	N/A
Carousel Center Parking Deck, Syracuse, New York	1997	100 m ²	N/A	N/A
Islington Ave. Bridge Interceptor Chambers, Toronto, Ontario	1997	180 m ²	N/A	N/A
Burlington Skyway Substructure, Burlington, Ontario	1997	268 m ²	N/A	N/A
Tulls Highway Overpass Deck, Seaford, Delaware	1997	1550 m ²	N/A	N/A
Hwy #6 & #11 Overpass Piers, Regina, Saskatchewan	1995	180 m ²	Up to 80	N/A
5 th Street & I-64 Substructure, Charlottesville, Virginia	1995	488 m ²	27–60 (at 6-19 mm) 13–53 (at 25-38 mm)	9 to 12
Hwy #1 & #6 Overpass Piers, Regina, Saskatchewan	1995	370 m^2	N/A	N/A
Hwy #2 Overpass Piers, Morinville, Alberta	1995	55 m ²	62 -96	N/A
34th Street & I-395 Bridge Deck, Arlington, Virginia	1995	733 m ²	76–82 (at 6-19 mm) 72–32 (at 19-32 mm)	11 to 15
Hwy #11 & #16 Overpass Piers, Saskatoon, Saskatchewan	1994	150 m ²	62-88	N/A
Pier Columns, SHRP, USA	1992	49 m ²	N/A	7 to 13
Abutment Area, SHRP, USA	1992	17 m ²	N/A	12 to 19
Deck Area, SHRP, USA	1991	136 m ²	60 (25 mm from bars)	20
Portage Avenue & Rt. 90 Retaining Wall, Winnipeg, Manitoba	1991	N/A	20 - 76	N/A
Burlington Skyway Pier, Burlington, Ontario	1989	31 m ²	27 (East Face) 59-60 (West Face) 57 (South Face)	11 (East) 32-33 (West) 30 (South)
U.S. Route No. 33 Bridge Deck (ODOT No. UNI-33.1138-R) Marysville, Ohio	1975	18 m^2	31 in 12 hr (at 0-25 mm) 51 in 24 hr (at 0-25 mm) 59 in 12 hr (at 25-51 mm) 70 in 24 hr (at 25-51 mm)	N/A

N/A = Not Available

Table 6. Half-cell potentials on treated and untreated North American structures [43, 44, 49]

ECE Date	Location	Test Date	Half Cell, mV (vs. Cu/CuSO ₄)
1989	Burlington Skyway Pier, Burlington, Ontario	Untreated	0% > -200 96% between -200 and -350
			4% < -350 96% > -200
		Treated	4% between -200 and -350
			0% < -350
1991	Portage Avenue & Rt. 90 Underpass	Untreated	84% < -350
	Retaining Wall, Winnipeg, Manitoba	Treated	100% > -280
1995	Hwy #6 & #11 Overpass Piers, Regina,		49% > -200
	Saskatchewan Un	Untreated	27% between -200 and -350
			24% < -350
			99% > -200
		Treated	1% between -200 and -350
			0% < -350
1997	Starbuck Bridge Deck, Traffic Bearing		6% > -200
	System, Winnipeg, Manitoba	Untreated	75% between -200 and -350
			19% < -350
			96% > -200
		Treated	4% between -200 and -350
			0% < -350

Effects of Electrochemical Chloride Extraction

The beneficial effect of ECE on the reduction of corrosion induced concrete deterioration of a structure after the treatment is important. However, there were concerns about the structural effects of ECE, such as hydrogen evolution at the cathode, bond strength loss between the concrete and reinforcement, and ASR susceptibility around the reinforcement. Many of these issues relate to the high voltage and current densities used during extraction, which changes the chemistry around the reinforcement and redistributes ionic species, thus altering the concrete's properties. [3, 56, 57]

The generation of nascent hydrogen at the cathode is inevitable due to the high voltages used during ECE. [3, 24] If hydrogen is absorbed into the steel, it could lead to hydrogen embrittlement and reduce the fracture toughness of the steel. [15,24] If hydrogen gas is produced, it can increase the local pressure and eventually promote cracking. [15,24,57] Currently, research indicates that hydrogen evolution will not adversely affect the structure, if current density levels are kept at 1 A/m² or less. [3, 23, 25] In addition, the lower strength steels used for reinforcement are not as susceptible to hydrogen embrittlement as high-strength steel. [3, 23] It is not surprising then that ECE is currently not recommended for high-strength steels used in prestressed concrete. [3, 23]

A study by Bennett, et al., suggested that porosity increased in the cement paste adjacent to the reinforcing steel. Using mercury porosimetry, it was determined that a significant increase occurred in the one- to ten-micron pore range following ECE. In addition, the cement adjacent to the top steel mat had undergone softening when compared with concrete extracted from deeper depths. In lekwaba, et al., noted the softening effect when the concrete was exposed to higher current densities (3 A/n^2) , but the effect was insignificant in samples exposed to lower current densities (1 A/m^2) . However, Bertolini, et al., did not observed a statistically significant change in microhardness measurements made near the reinforcing steel after exposing samples for twelve weeks to current densities that ranged from 5 mA/m² to (1 A/m^2) . In addition, Bennett, et al., indicated that even at high current densities (20 A/m^2) , ECE was not detrimental to the compressive strength of the concrete.

Broomfield discusses research that indicates that corrosion product on the surface of the reinforcing steel improves the bond strength with the concrete. [3,25] The elimination of expansive corrosion product during ECE seems to reduce this bond strength. [3] Initially, the force required is greater for the corroded sample, but as the ECE treatment time increases, the pull out load decreases to approximately the same value as the control sample. [3] Others have shown that in addition to removing chlorides, ECE physically changes the concrete. [61,62] However, these effects appear to be minor at the current densities commonly employed during the ECE process.

Bertolini, et al., indicated that ASR could result if ECE was applied to concrete containing susceptible aggregate. This was due to accumulation of alkali metal ions and hydroxyl ions near the reinforcing steel during ECE. However, electrolytes containing lithium ions (i.e. lithium borate electrolyte) have demonstrated the ability to suppress ASR. [3, 23, 59]

Purpose

Currently, the primary focus of the research has been to study how the electrical parameters of the regions between the anode and the cathode change during ECE. It was expected that this approach would provide insight into the decrease in current flow during the early stage of ECE treatment, with which improvements to the efficiency of chloride removal can be made. In addition, the influence of w/c ratio on ECE was investigated using specimens made of several w/c ratios. It was hoped that this might lead to a correlation between the w/c ratio and the time required for chloride extraction.

EXPERIMENTAL METHOD

Specimen Design

Two types of reinforced concrete specimen designs are being used during this portion of the study. Tables 7 and 8 list the basic design features of each type of specimens. These specimens included variations in the method of introducing chlorides into the concrete, cover thickness, and w/c ratios. Tables 9 and 10 list the mix designs for the Type I and II concrete specimens, respectively. After the specimens had cured, a dam was affixed to the top of each specimen to hold the appropriate solutions. The Types I blocks were kept in a controlled laboratory environment, whereas Type II blocks were exposed outside to further simulate field conditions.

As illustrated in figure 4, the Type I specimens were designed with two rows of activated titanium rods embedded in the concrete above the reinforcing steel bar. Each row has four activated titanium rods aligned in a horizontal plane at 1.0 cm below the top concrete surface and 1.0 cm above the rebar. This arrangement allowed for the measurements of the IR drop and voltage differences at selected depths during an ECE experiment. A list of these points is provided in table 11. In addition, these rows of embedded titanium electrodes allowed for measurement of the changes in the concrete resistivity at the two depths, using the four-pin method (ASTM G-57).

Each Type II concrete specimen contained a thermocouple, two titanium-mesh ribbon strips, and a corrosion probe embedded in the concrete, as illustrated in figure 5. The thermocouple was located 3.8 cm from the surface and horizontally centered in the sample. Each titanium mesh ribbon was 1.3 cm wide and 5.1 cm long. The two ribbons were 6.35 cm apart and located 1.0 cm below the surface. The corrosion probe was a graphite reference electrode and a 1.3-cm wide titanium counter electrode, all encased in concrete. Before casting the specimens, the probe was attached to the upper reinforcing steel mat. Table 12 lists the various contact points between which measurements can be made during ECE experiments on these specimens.

Table 13 lists the differences in some of the physical characteristics of Type I and II concrete specimens being used in this study.

Chloride Exposure Method	Height x Length x Width	Cover Thickness	W/C	Number of Block Cast	Blocks Tested
Admixed and Ponding	9.2 cm x 12.7 cm x 30.5 cm	4.4 cm	0.40	3	3
Admixed and Ponding	9.2 cm x 12.7 cm x 30.5 cm	4.4 cm	0.45	3	3
Admixed and Ponding	9.2 cm x 12.7 cm x 30.5 cm	4.4 cm	0.50	3	3
Admixed and Ponding	9.2 cm x 12.7 cm x 30.5 cm	4.4 cm	0.55	3	3
Admixed and Ponding	10.5 cm x 12.7 cm x 30.5 cm	5.7 cm	0.40	3	2
Admixed and Ponding	10.5 cm x 12.7 cm x 30.5 cm	5.7 cm	0.45	3	2
Admixed and Ponding	10.5 cm x 12.7 cm x 30.5 cm	5.7 cm	0.50	3	2
Admixed and Ponding	10.5 cm x 12.7 cm x 30.5 cm	5.7 cm	0.55	3	2
Admixed and Ponding	11.8 cm x 12.7 cm x 30.5 cm	7.0 cm	0.40	3	0
Admixed and Ponding	11.8 cm x 12.7 cm x 30.5 cm	7.0 cm	0.45	3	0
Admixed and Ponding	11.8 cm x 12.7 cm x 30.5 cm	7.0 cm	0.50	3	0

7.0 cm

0.55

3

0

Table 7. Description of Type I concrete test blocks

11.8 cm x 12.7 cm x 30.5 cm

Admixed and Ponding

Table 8. Description of Type II concrete test blocks

Chloride Exposure Method	Height x Length x Width	Cover Thickness	W/C	Number of Block Cast	Blocks Tested
Ponding	17.7 cm x 61.0 cm x 60.8 cm	3.8 cm	0.45	4	1
Ponding	17.7 cm x 61.0 cm x 60.8 cm	3.8 cm	0.50	4	1
Ponding	17.7 cm x 61.0 cm x 60.8 cm	3.8 cm	0.55	4	1
Ponding	17.7 cm x 61.0 cm x 60.8 cm	3.8 cm	0.60	4	1
Ponding	19.0 cm x 61.0 cm x 60.8 cm	5.1 cm	0.45	4	0
Ponding	19.0 cm x 61.0 cm x 60.8 cm	5.1 cm	0.50	4	0
Ponding	19.0 cm x 61.0 cm x 60.8 cm	5.1 cm	0.55	4	0
Ponding	19.0 cm x 61.0 cm x 60.8 cm	5.1 cm	0.60	4	0
Ponding	20.3 cm x 61.0 cm x 60.8 cm	6.4 cm	0.45	4	2
Ponding	20.3 cm x 61.0 cm x 60.8 cm	6.4 cm	0.50	4	2
Ponding	20.3 cm x 61.0 cm x 60.8 cm	6.4 cm	0.55	4	2
Ponding	20.3 cm x 61.0 cm x 60.8 cm	6.4 cm	0.60	4	2

Table 9. Mix design for Type I concrete specimens

W/C	0.40	0.45	0.50	0.55
Cement (Type I/II), kg/m ³	377	377	377	377
Water, kg/m ³	151	169	188	208
Course Aggregate, kg/m ³	898	898	898	898
Fine Aggregate, kg/m ³	886	886	886	886
Cl Added, kg/m ³	5.77	5.81	5.87	5.91
Cl, % by Wt.	0.25	0.25	0.25	0.25

Table 10. Mix design for Type II concrete specimens

W/C	0.45	0.50	0.55	0.60
Cement (Type I/II), kg/m ³	377	331	301	276
Water, kg/m ³	170	166	166	166
Course Aggregate, kg/m ³	1061	1061	1061	1061
Fine Aggregate, kg/m ³	719	766	794	815
Daravair (Air Entrainment), kg/m ³	0.20	0.14	0.12	0.08
Daratard (Set Retarder), kg/m ³	0.75	0.63	0.56	0.52

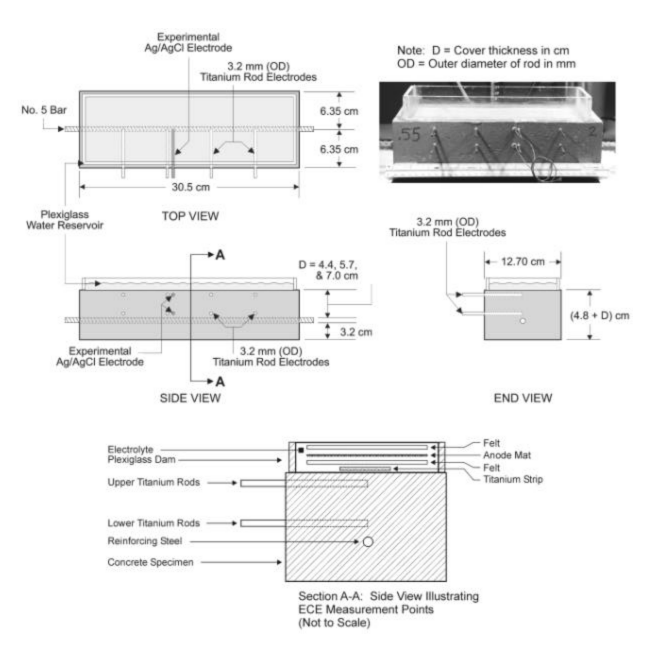


Figure 4. Illustration of a Type I Specimen

Table 11. Description of contact points used to make measurements in Type I concrete test blocks

Region Studied	Description
Anode/Anolyte Ti Strip	Measurement contact points are the anode mat and the a titanium
Anode/Anolyte 11 Surp	strip located in the anolyte.
Anode/Upper Ti Rod	Measurement contact points are the anode mat and a titanium rod
Alloue/Opper 11 Rou	located in the top row of embedded titanium rods.
Anode/Rebar	Measurement contact points are the anode mat and reinforcing steel
Anoue/Rebai	mat.
	Measurement contact points are a titanium strip located in the
Anolyte Ti Strip /Upper Ti Rod	anolyte and a titanium rod located in the top row of embedded
	titanium rods.
Lower Ti Rod/Rebar	Measurement contact points are a titanium rod located in the bottom
Lower II Rod/Rebai	row of embedded titanium rods and the reinforcing steel mat.
	Measurement contact points are a titanium rod located in the top row
Upper/Lower Ti Rod	and a titanium rod located directly below it in the bottom row of
	embedded titanium rods.

Table 12. Description of contact points used to make measurements in Type II concrete test blocks

Region Studied	Description
Anode/Anolyte Ti Strip	Measurement contact points are the anode mat and a titanium strip located in the anolyte.
Anode/Rebar	Measurement contact points are the anode mat and reinforcing steel mat.
Anolyte/Concrete Ti Strip	Measurement contact points are a titanium strip located in the anolyte and a titanium strip embedded in the concrete.
Concrete Ti Strip/Rebar	Measurement contact points are a titanium strip embedded in the concrete and the reinforcing steel mat.

Table 13. ECE comparison between the different specimens

Description	Type I	Type II
Treated Concrete Surface Area	248 cm ² *	3716 cm ² **
Rebar Surface Area	130 cm ² *	4865 cm ² **
Number of Reinforcing Mats	1 (single bar)	2

^{*} Based on the interior dam dimensions (9.5 cm X 26.1 cm)

^{**} Based on the interior dam dimensions (60.96 cm X 60.96 cm)

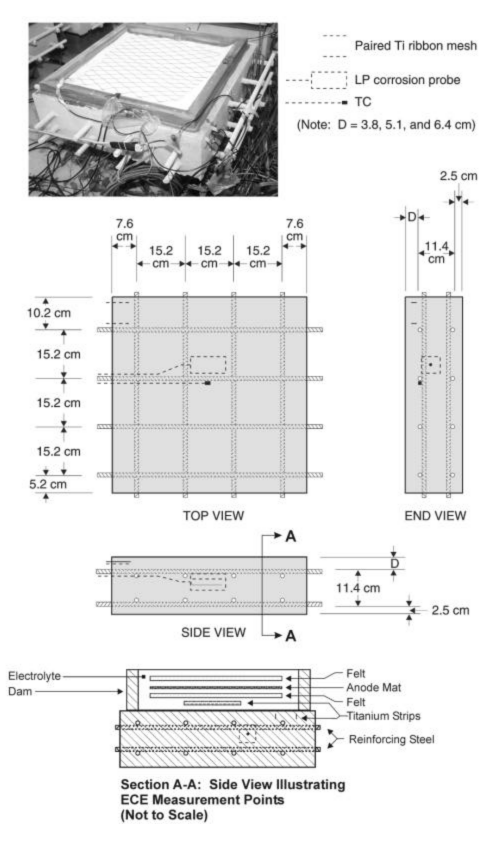


Figure 5. Illustration of a Type II Specimen

Electrochemical Chloride Extraction

Chlorides were extracted from the concrete specimens following the methods employed in previous ECE projects. The ECE parameters used are listed in table 14. In each experiment, a titanium mesh anode and two pieces of felt were cut to fit the inside dimensions of the dam. A piece of felt was placed on the surface of the concrete inside the dam, which was followed by the titanium mesh anode, and finally the titanium mesh was covered by a second piece of felt. The sandwiching of the titanium mesh between the felt ensured the complete wetting of the titanium-mesh anode. The anolyte was carefully added until the solution level inside the dam completely covered the upper felt mat. Either a saturated calcium hydroxide solution or a lime and water solution were used as the anolyte during ECE. A DC power supply was set to operate in constant current mode (1 A/m²) until it reached the maximum voltage output, at which time it would switch from constant current to constant voltage mode. The maximum voltage setting was dependent on the power supply. For the Type II specimens the maximum voltage was 40 V. A maximum of 40 V was also applied to one Type I specimen from each of the four w/c ratios studied, all of which had a cover thickness of 1.7 cm. For all other Type I specimens, the maximum voltage was between 9 V and 15 V. The positive lead from the power supply was attached to the anode and the negative lead to the reinforcing steel mat. To minimize the evolution of chlorine during ECE, the pH was maintained above 10 by adding either calcium hydroxide or lime to the anolyte.

 Description
 Selection

 Anode Material
 Activated titanium mesh

 Anode Contact Material
 Two layers of felt: one above and one below the mesh anode

 Electrolyte
 Saturated calcium hydroxide or lime

 Maximum Current Density (based on concrete surface area)
 1 A/m²

Table 14. ECE parameters

Current and Voltage Measurements

Current and voltage measurements were made using a Tetronix digital multimeter or an IO Tech Logbook data acquisition system. With both instruments, voltage measurements were made directly. The current was calculated using the measured voltage across a resistor of known resistance and Ohm's law.

IR Drop Measurements

The total electrochemical cell voltage is the sum of a series of contributions from individual voltage differences. In simplest terms, these contributions are comprised of a thermodynamic potential difference (E_{thermo}), anodic and cathodic overpotentials (η_a and η_c , respectively), and the voltage drop due to current flow through a resistive solution (IR_{sol}). During ECE, the measured voltage difference (E_{meas}) is represented by equation 9 since the system is being driven. Figure 6 illustrates this relationship graphically.

$$E_{Meas} = E_{Thermo} + \left| \boldsymbol{h}_{a} \right| + \left| \boldsymbol{h}_{c} \right| + IR_{Sol} \tag{9}$$

If the applied current is rapidly interrupted and the change in voltage is quickly recorded, the IR_{sol} component of the total voltage can be determined, as shown in figure 7.^[66] In the galvanostatic case, the solution resistance, R_{sol} , can be easily determined by dividing the measured voltage drop with the current.

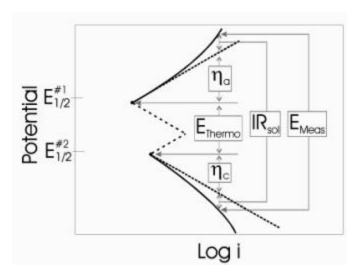


Figure 6. Illustration of voltage components for a driven system [64]

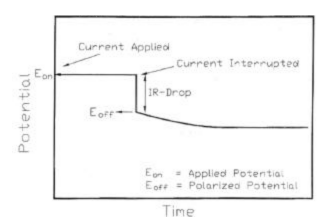


Figure 7. Change in voltage after interruption of applied current [66]

As with any measurement, care must be taken to minimize errors since this could yield misleading results. Therefore, the data acquisition system used must be capable of capturing events that are as short as a few milliseconds. [66] Thompson and Payer suggested the use of an oscilloscope to capture IR-drop events. [66] Moreover, since the current value is required to calculate the solution resistance, fluctuations in the current can significantly impair calculations of the solution resistance. [66]

To guarantee the current interruption were reproducible, a solid-state relay controlled by a timing circuit was used to interrupt the current. Testing confirmed the solid-state relay interrupted the current in less than 6 ns. The timing circuit then maintained the solid-state relay in an open position for 13 ms.

To gather the voltage vs. time data, the IO Tech Logbook data acquisition system was used. To ensure the Logbook system would suffice, a HP 150MHz oscilloscope was used to verify that the Logbook acquisition rates were adequate. This was performed on circuits with known resistance values as well as on concrete test specimens. Following a series of successful comparisons between the two instruments, evaluations of Type I specimens began using the Logbook system. The Logbook acquisition system was set to gather data during periodic interruptions of the ECE process. Although the total current interruptions lasted for only 13 ms, data was gathered before, during, and after the interruption. This data was then used to determine changes in the IR-drop during ECE.

4-Pin Resistivity Measurements

As discussed earlier, the Type I concrete specimens were designed with the intention of making resistivity measurements during ECE. This was performed following ASTM Standard G 57, and using a Nilsson Soil Resistance Meter, Model 400. ^[67] This type of meter induces an AC signal between two outer pins while the voltage drop is measured between two inner pin, which is illustrated in figure 8. ^[67] The output from this meter is in units of ohms (resistance), and therefore if the pins are evenly spaced and the inner pin spacing (a) is known, the resistivity (\mathbf{r}) can be calculated using the following equation. ^[67,68]

$$\mathbf{r} = 2\mathbf{p}aR \tag{10}$$

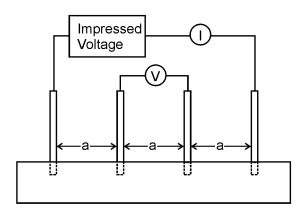


Figure 8. Four pin resistivity test method

Half-Cell Measurements

Half-cell measurements were made following the guidelines set forth in ASTM Standard C 876. Since these measurements were made on concrete surfaces, a damp sponge was used to ensure adequate contact between the half-cell and the concrete surface. Connectors were welded to each reinforcing steel mat, so that each piece of rebar was externally connected to each other. This ensured that the entire mat was conductive. In all cases, saturated Cu/CuSO₄ electrodes (CSE) were used to make measurements against the internal reinforcing steel mat.

Collection of Concrete Samples

To monitor the changes in chloride concentrations in the specimens during ECE, ground concrete samples were collected for chloride analysis from the specimens at different stages of the experiments, in accordance with AASHTO T 260. A sample collection scheme was used on the Type II specimens to minimize any possible interference that the collection of samples during the ECE experiment would impose on the current flow and voltage distribution in the specimen following the collection of concrete samples. Under this scheme, concrete samples were collected by starting on the outer perimeter of a specimen and working inward. An example of the sampling patterns used is shown in figure 9. At each time, three different points were sampled. For the blocks with a cover thickness of 6.4 cm, sampling was performed above a single bar. For the blocks with a cover thickness of 3.8 cm, sampling included the intersection of two bars, above a single bar, and where no bars were present. However, the same approach was not possible for the Type I specimen due to size constraints. In this case, the same technique for sample collection was used, but fewer samples were collected. Samples were collected before and after ECE in all cases. The sample depths were from the surface to 0.6 cm, 0.6 cm to 1.9 cm, 1.9 cm to 3.2 cm, and 3.2 cm to 4.1 cm. All of these depths were above the top reinforcing steel mat.

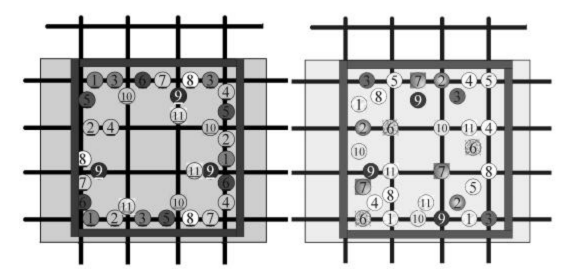


Figure 9. Type II specimen drill pattern for sample collection: Right, for a cover thickness of 6.4 cm; Left, for a cover thickness of 3.8 cm

Potentiometric Titration

The acid-soluble chloride concentrations of the collected ground concrete samples were determined following AASHTO T 260. The analysis followed Method II in this standard, which uses the Gran Plot Method to determine the endpoint of the titration. A silver-ion-selective electrode was use during the titration.

X-Ray Diffraction

Surface residue that formed during ECE was analyzed using x-ray diffraction (XRD). It was anticipated that XRD would help in identifying any crystalline material in the residue. To accomplish this, residue samples were scraped from the surface and ground into a fine powder for analysis. These samples were then placed into the instrument for analysis.

XRD was performed using a Scintag automated diffraction system. During analysis, the applied voltage was 40 KV and the current was 35 mA. A copper target was used for the Ka x-ray source with a nickel filter to reduce undesirable components in the spectrum. The divergence and scatter slits on the source were 2 and 3, respectively. On the detector, the scattering and receiving slits were 1 and 0.5, respectively.

The XRD spectrum was evaluated using the program Diffraction Management System Software, version 1.1. Background subtraction was performed using a boxcar curve fit with a filter width of 1.5 degrees. The program's peak library software was used to compare the unknown sample against known spectra.

X-Ray Photoelectron Spectroscopy

To aid in identifying the composition of the residue that formed during ECE, a Perkin Elmer 560 system adapted for x-ray photoelectron spectroscopy (XPS) was used to analyze powder samples. Unlike XRD, which yields information about the bulk material in a sample, XPS provides surface information. The XPS data would be used to provide additional insight into the elements and compounds on the surface of the residue.

Charging effects during XPS analysis were adjusted for by setting the adventitious carbon line to 284.8 eV. Preliminary peak comparisons were first made against values cited in the literature. After reducing the possibilities, final identification was made against a sample of reagent grade calcium chloride.

RESULTS AND DISCUSSION

Currently, this study has involved a number of specimens to obtain the following results. ECE studies have been completed on 20 of the Type I and 12 of the Type II test blocks. These blocks included different cover depths, which ranged from 3.5 cm to 6.4 cm. To evaluate the affect of the w/c ratio on ECE, test blocks with ratios ranging from 0.40 to 0.60 were designed. Descriptions of the block designs are presented in tables 7 through 12 and illustrated in figures 4 and 5.

Changes in the Current and Voltage during ECE

During ECE, the current was monitored, while internal and external voltage measurements were made. The designations in the following graphs to the different measurement points that were monitored during ECE for the Type I and Type II test blocks are listed in tables 11 and 12, respectively. Figures 4 and 5 can provide additional insight into the connections used to make measurements.

The typical change in voltage during ECE for a set of Type I specimens with different w/c ratios, but the same cover depth is shown in figure 10. The benefit of these specimens was the ability to measure the voltage changes in different layers between the anode and cathode. Under constant voltage conditions, it can be seen that the voltage between the anode and the upper titanium rods increases. At the same time, the voltage between the lower titanium rods and the reinforcing steel is decreasing. In each case, the rate of change of the voltage is greatest initially, i.e., during the first 10 to 15 days, and then the rate of change decreases for the remaining extraction period. All of the specimens evaluated exhibited this type of behavior. Even with larger applied voltages, the voltage difference in the top layer of concrete increased during ECE. A typical example of this is shown for two specimens in figure 11. As the applied voltage to the slab increases under constant current conditions, the voltage across the top layer of concrete increases.

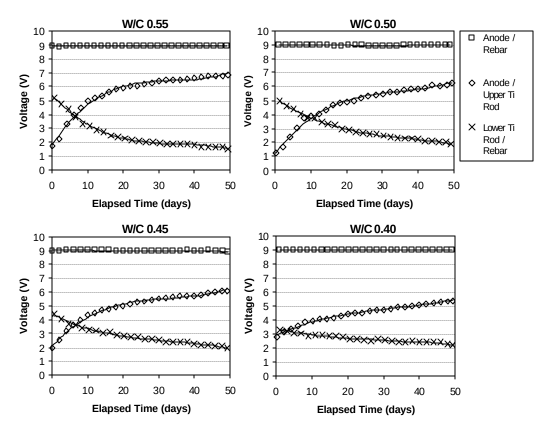


Figure 10. Measured voltage changes during ECE in a set of Type I specimens (Maximum voltage was 9V)

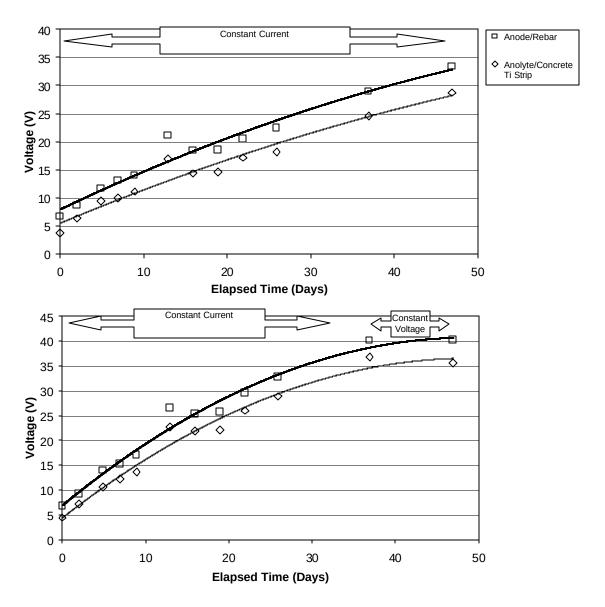


Figure 11. Comparison of the voltages measured between (1) the anode and rebar and (2) between the anolyte and the titanium strip in the concrete. Top: for a 0.55-w/c Type II specimen; Bottom: for a 0.50-w/c Type II specimen (Maximum voltage was 40V)

Influence of Concrete Surface on Voltage and Current

To better understand the influence of the surface-layer concrete on the voltage and current, one of the Type I specimens was subjected to an extended ECE experiment and analysis. A timeline illustrating the sequence of events that took place in this extended experiment is shown in figure 12. First, the specimen was ECE treated or polarized for 26 days and then depolarized for 38 days. The voltage and current data for this 26-day ECE treatment are shown in figure 13a. Then, the specimen was re-energized for 12 hours. The voltage and current data for this first 12-hr polarization are shown in figure 13b. This was followed by removal and storage of the electrolyte used during the ECE test period. The surface of the concrete was then sandblasted and the stored electrolyte was poured back into the reservoir. After allowing twenty hours for the solution to soak into the concrete, ECE was initiated again for another 12 hours and

the same measurements were made (figure 13c). Finally, the sample was depolarize for 36 hours and then re-energized for a final 12-hour duration and a final set of measurements was made (figure 13d).

Comparison of figures 13a and 13b would indicate that the voltages and current of the system at the start of the first 12-hour re-polarization were practically the same as where the system was at the end of the 26-day of polarization. Comparison of figures 13b and 13c would reveal the effect of the sandblasting the surface layer concrete on the voltages and current. It is clear that sandblasting the surface of this specimen increased the current density by over 0.3 A/m². This current density was even greater than the initial current density value (at the beginning of the 26-day treatment). Following sandblasting, the voltage between the anode and the reinforcing steel also switched from constant-voltage mode to constant-current mode, but then returned to a constant-voltage mode. In addition, sandblasting resulted in voltage changes within the different concrete regions, which are on the order of approximately 2 V. This indicates that the surface of the concrete appears to have a significant influence on the voltage and current during ECE. However, additional testing on other specimens will be required to confirm these observations.

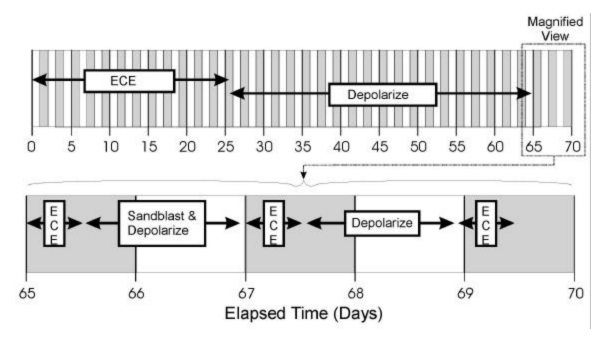


Figure 12. Timeline of concrete surface study

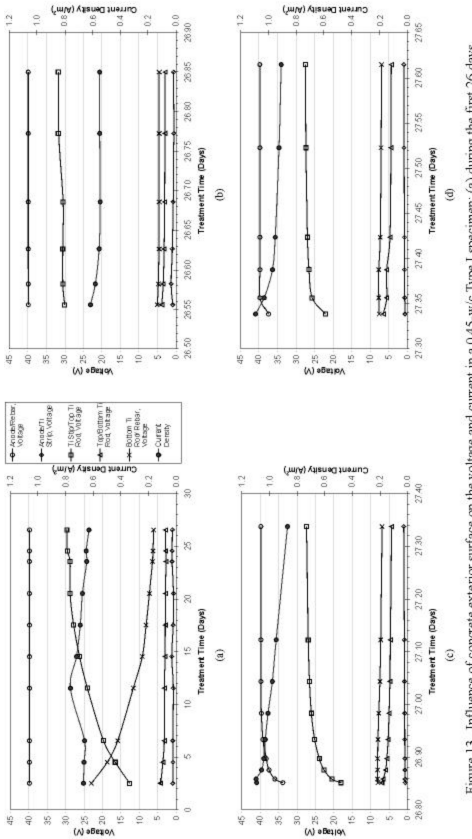


Figure 13. Influence of concrete exterior surface on the voltage and current in a 0.45-w/c Type I specimen: (a) during the first 26 days of ECE; (b) during the first 12 hours of ECE, following 38 days of depolarization; (c) during the second 12 hours of ECE, following depolarization and sandblasting; and (d) during the third 12 hours of ECE, following depolarization without additional sandblasting (Current density based on concrete surface area, Maximum Voltage 40V)

Changes in the Concrete Resistance During ECE

The resistance between different points in the concrete was determined using the IR Drop technique, which was discussed in the earlier section "IR Drop Measurements" on page sixteen. During ECE, the resistance was observed to increase in the region between the anode and the upper layer of titanium rods, as shown in figure 14. In contrast, the solution (concrete) resistance decreased in all other regions during ECE. A comparison, based on w/c ratios, of the resistance between the anode and the upper titanium rods did not indicate an obvious relationship. This pattern was consistent in all of the samples studied.

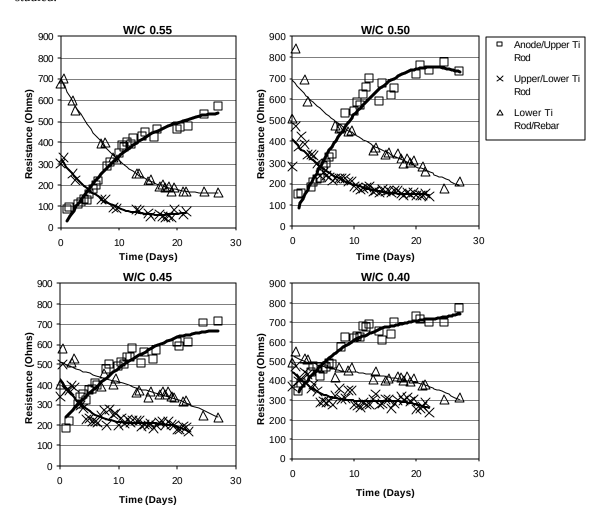


Figure 14. Example showing the change in resistances for a single set of Type I specimens during ECE

Resistivity

It was apparent that, immediately before ECE, the resistivity of the top layer of concrete, as measured by the upper row of four titanium rods, was less than that of the lower concrete that surrounded the lower set of four titanium rods, as shown in figure 15. This difference was observed in the other specimens prior to ECE. It is also apparent in figure 15 that the resistivity of the lower region of the concrete undergoes only a relatively slight to moderate change during ECE. In contrast, the resistivity of the top layer of concrete increased and exceeded that of the lower concrete and then remained at a level that is greater than that of the lower concrete region. However, a clear relationship between w/c ratio and

resistivity for either the upper or lower titanium rod region was not evident. This is shown in figures 16 and 17, respectively.

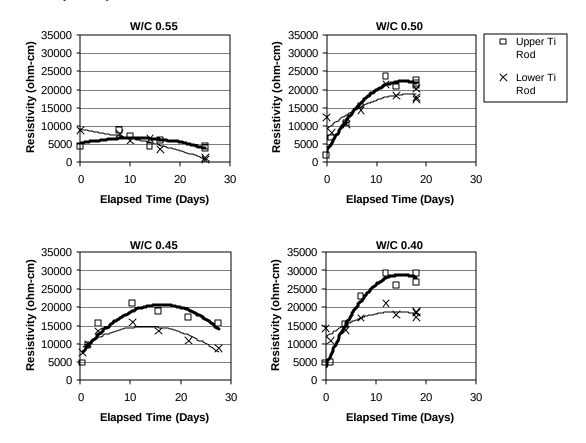


Figure 15. Change in Resistivity for Type I specimens of various w/c ratios (cover thickness of 4.4 cm)

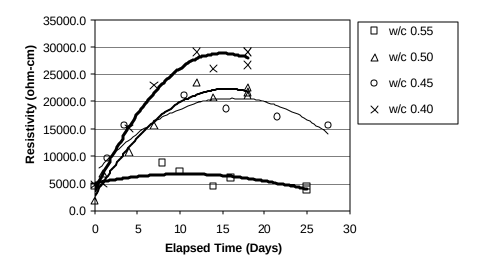


Figure 16. Resistivity change in the upper layer of concrete, for Type I Specimens with various w/c ratios (cover thickness of 4.4 cm)

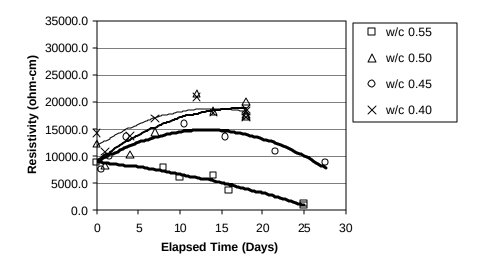


Figure 17. Resistivity change in the lower layer of concrete, for Type I Specimens with various w/c ratios (cover thickness of 4.4 cm)

Chloride Concentration in Concrete with ECE

The previous data have shown that the concrete specimens studied behaved in a stratified manner during ECE. Generally, the resistivity in the upper layers of concrete increased while that at the lower layers decreased. Upon determining the chloride concentration before and after ECE in Type I specimens, it is evident that a large change in chloride concentration occurs near the surface adjacent to the anode, which is shown in figures 18 and 19. (The negative values in figures 18 and 19 indicate a decrease in chloride concentration; conversely, positive values signify an increase in chloride concentration.) It is apparent that significant removal of chlorides during ECE are occurring in the concrete layer near the anode. In contrast, the treated Type II specimens exhibited a more even removal distribution at deeper depths in the concrete, which is shown in figures 20 and 21.

Unlike the Type II specimens, the Type I specimens exhibited a decrease in chloride removal at deeper depths within the concrete and some specimens even display an increase in chloride concentration near the reinforcing steel. This is attributed to two factors: (1) the use of admixed chlorides in the Type I specimens and (2) the difference in the steel surface area in the different specimen types (table 13). It is possible that the admixed chlorides below the bar were drawn upward during ECE. In addition, these results support earlier studies that indicate the importance of available cathode area. The decreased cathode surface area in the Type I specimens appears to decrease the total current flow, which decreased the percentage of chloride removed. In the Type I specimens, 0% - 52% of the chlorides were removed during treatment, whereas 33% - 76% of the chlorides were removed from the Type II specimens.

Cover thickness appears to influence the quantity of chloride removed. However, additional samples must be evaluated to provide statistical validity to the observed trend.

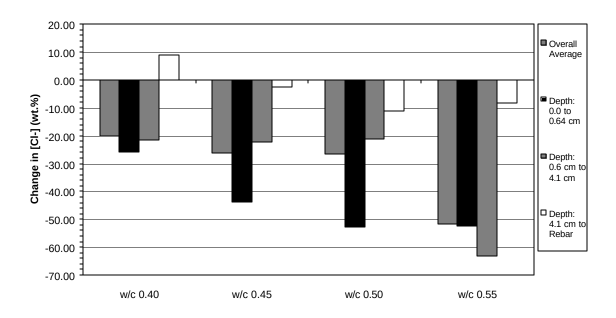


Figure 18. Average change in chloride concentrations due to ECE in Type I specimens with 4.4 cm of concrete cover over rebar

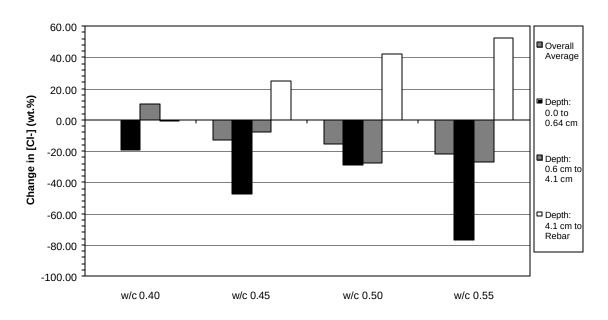


Figure 19. Average change in chloride concentrations due to ECE in Type I specimens with 5.7 cm of concrete cover over rebar

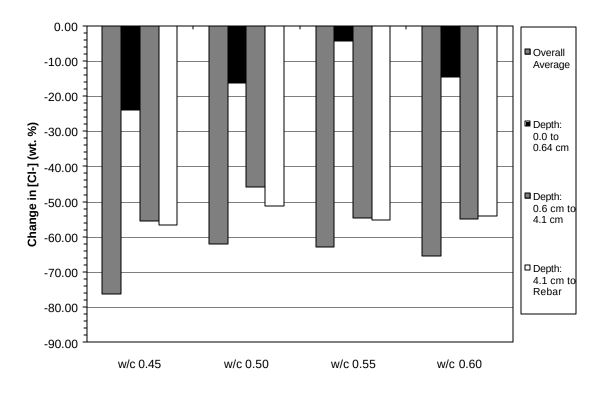


Figure 20. Change in chloride concentrations due to ECE in a single set of Type II specimens with 3.8 cm of concrete cover over rebar

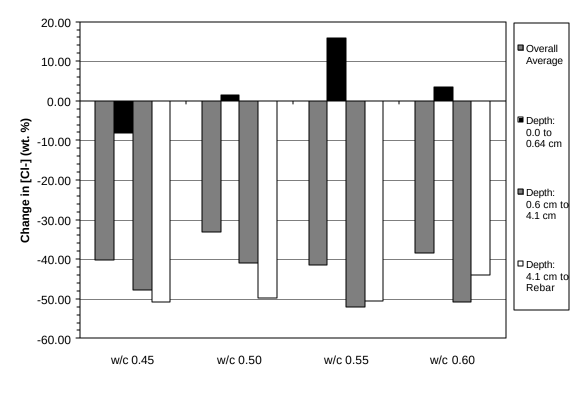


Figure 21. Average change in chloride concentrations due to ECE in Type II specimens with 6.4 cm of concrete cover over rebar

Visual Observations

After completing ECE, each specimen's surface was examined for physical changes. In each case, a tightly adhering substance had formed on the surface. Images of this formation are shown in figure 22. Attempts to remove the unknown material from the surface proved difficult. Upon trying to cleave it, a portion of the concrete was dislodging in addition to the unknown material. This provided a cross section of the interface, which is shown in figure 23. A surface formation following treatment has been found on actual treated bridge decks. As shown in figure 24, the formation parallels the reinforcing steel mat below. It is evident from these photographs that the surface formation creates perpendicular lines that crisscross the older concrete, as well as the recently repaired concrete (areas covered with more white material than their surrounding). Surface formation samples were collected and are currently being analyzed.

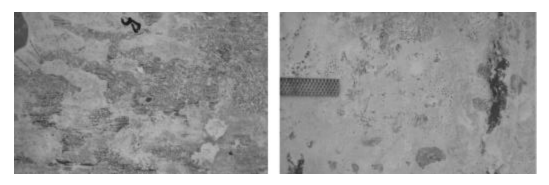


Figure 22. Tightly adhering layer of white material formed on the concrete surface during ECE

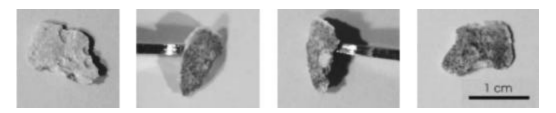


Figure 23. Various views of surface layer that formed on the concrete during ECE: (Left) top view, (Middle) edge views, (Right) bottom view



Figure 24. Layer of white material formed on the concrete surface directly above the reinforcing steel following ECE on an actual bridge deck ^[70]

The increased current flow because of sandblasting the concrete surface, which removed the residue, was discussed earlier. That particular specimen is shown in figure 25. It is interesting to note that

white deposits formed in the exposed pores, which is shown in the bottom photograph in figure 25. As indicated earlier, the study of new samples (shown in figure 27) will add statistical validity to the observed improvement in current flow following sandblasting. It is expected that removal of the tightly adherent surface formation will improve the ECE process.

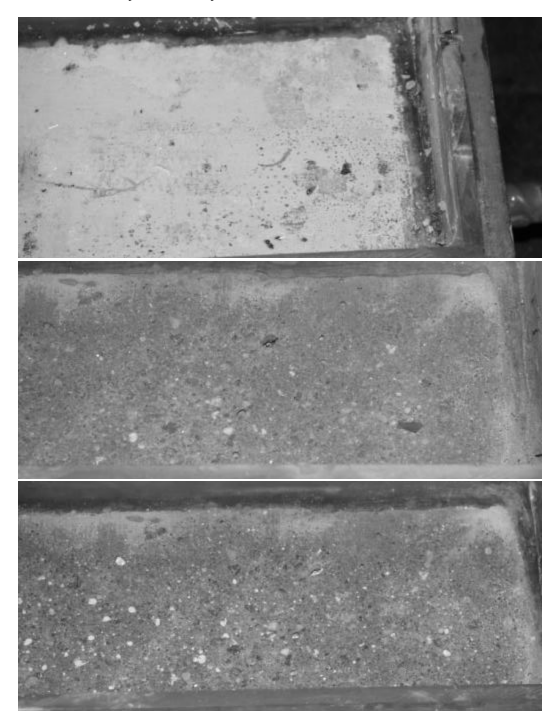


Figure 25. Type I specimen surface appearance: (Top) after ECE but prior to sandblasting, (Middle) after sandblasting but prior to second application of ECE, (Bottom) following a second application of ECE for 12 hours

Surface Deposit Analysis

To identify the white material covering the concrete surface following ECE, which is shown in figures 22 and 23, XRD was performed on two different Type II specimens. Calcium carbonate was clearly identified by XRD to be present in both specimens. Figure 26 is an overlay of a calcium carbonate spectrum over the top of the spectrum for one of the unknown samples. It is apparent that all of the peaks locations in the standard match the peaks locations in the unknown sample. However, the intensities differed, which indicates additional crystalline materials are most likely present in minute amounts in the sample. In addition, not all of the peaks in the unknown sample diffraction pattern are accounted for by the calcium carbonate standard. Based on these observations, calcium carbonate appeared to be the major component in the white residue; however, the presence of other components in trace quantities is evident. The XRD spectrum from the other sample analyzed d isplayed the same characteristics as those found in the sample shown in figure 26. Table 15 lists the intensities and peak positions for the two samples examined.

XPS was also performed on the white material deposited on one Type I specimen and one Type II specimen. The Type I specimen used calcium hydroxide as the electrolyte, whereas industrial lime was used as the electrolyte for the Type II specimen. The XPS data indicated that calcium chloride was present on the surface of both samples. In addition, XPS detected magnesium in the sample exposed to a solution of industrial lime, but magnesium was not present in the sample that used reagent grade calcium hydroxide as the electrolyte. Work is underway to evaluate the significance of these finding through ongoing analysis on additional samples. Table 16 lists the elements and binding energies for the two samples examined.

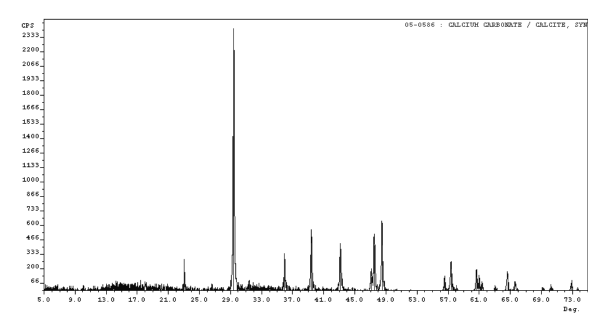


Figure 26. Surface deposit XRD pattern from a Type II specimen comparing the unknown material to calcium carbonate

Table 15. Surface deposit peak data using x-ray diffraction

Unknown #1 S	Unknown #1 Surface Deposit		Unknown #2 Surface Deposit		
2?, Deg	Intensity, CPS	2?, Deg	Intensity, CPS		
13.080	61.000	12.380	78.333		
13.320	54.000	12.980	95.000		
14.040	56.000	13.220	88.333		
14.340	84.000	13.320	71.667		
14.500	56.000	13.580	88.333		
14.800	59.000				
15.120	63.000	13.660 13.849	131.667 81.167		
15.120	63.467	14.140	128.333		
15.760	56.000	14.300	103.333		
15.880	63.000	14.460	80.000		
16.160	56.000	14.680	76.667		
16.340	54.000	15.040	80.000		
16.760	69.000	15.240	68.333		
17.120	71.000	15.460	56.667		
17.340	48.000	15.800	116.667		
17.660	52.000	16.160	56.667		
18.061	59.950	16.460	113.333		
18.700	56.000	16.620	76.667		
19.520	63.000	17.120	75.000		
20.180	59.000	17.440	93.333		
20.900	60.000	17.620	61.667		
23.119	162.533	17.820	106.667		
29.461	2347.350	18.145	213.700		
29.980	49.000	18.320	126.667		
30.400	53.000	18.740	75.000		
31.500	91.000	19.380	61.667		
31.620	56.000	19.680	75.000		
31.920	74.000	20.280	80.000		
32.160	46.000	21.620	65.000		
32.700	64.000	23.160	188.333		
35.993	266.550	27.440	51.667		
36.300	70.000	28.020	80.000		
39.458	553.950	29.020	60.000		
39.700	79.000	29.393	2408.000		
43.201	415.500	29.690	95.000		
47.190	190.050	30.460	63.333		
47.555	498.533	30.620	88.333		
47.900	54.000	31.500	83.333		
48.557	614.350	31.560	103.333		
56.627 57.446	126.150	32.520 33.020	51.667		
60.708	248.483 180.717	33.020	86.667 53.333		
61.051	132.933	34.040	55.000		
61.412	70.233	34.040	81.850		
64.699	162.000	34.300	93.333		
65.644	77.183	36.051	251.367		
70.320	56.000	39.471	405.367		
72.960	93.000	43.226	448.333		
, 2., 500	75.000	47.228	202.267		
		47.585	467.133		
		48.585	488.733		
		56.624	58.000		
		57.436	178.167		
		60.739	137.117		
		61.480	50.000		
		64.714	120.733		
		65.718	80.733		

Table 16. Surface deposit peak data using x-ray photoelectron spectroscopy

Unknown #1, Ca(OH) ₂ electrolyte		Unknown #2, Industrial lime electrolyte	
Element	Binding Energy, eV	Element	Binding Energy,
(Photoelectron Line)	- '	(Photoelectron Line)	eV
C (1s)	285.9	C (1s)	286.3
Ca (2p 3/2)	348.6	Ca (2p 3/2)	349.0
Cl (2p 3/2)	199.1	Cl (2p 3/2)	199.1
O(1s)	533.2	O(1s)	533.4
Mg (2p 3/2)	None Present	Mg (2p 3/2)	51.0
Si (2p)	102.7	Si (2p)	102.9

CONCLUSIONS AND FUTURE WORK

Conclusions

- A clear relationship between w/c ratio and chloride extraction rate was not observed.
- Sandblasting the concrete surface after applying ECE increases the current density at equivalent operating voltages, thus increasing the efficiency of the ECE process.
- The decrease in current flow that occurs typically during the early stage of an ECE is attributable to a significant increase in the electrical resistivity of the surface layer of concrete.
- Preliminary analyses indicate that the resistivity increase is related to the formation of residue on the concrete surface and pores. Analyses of the residue reveal that it contains calcium carbonate, calcium chloride, and other yet unidentified minor components.

Future Work

Future research will begin to focus on two regions, (1) the anolyte/concrete interface and (2) the concrete/steel interface. In addition, the effects of sandblasting and various electrolytes on current efficiency will be investigated. Testing will include previously used nondestructive techniques as well as the destructive evaluation of samples. The rational for destructive evaluation is to ascertain a clearer understanding of changes within the concrete and at the concrete/steel interface. Currently, pH, XRD, and XPS measurements, as well as visual inspection are being planned for the interior portions of the concrete.

The upcoming testing will be performed on existing specimens as well as on new specimens with a different design. It is anticipated that by combining favorable features from prior specimens, this improved specimen design will provide additional insight into the changes occurring within the concrete during ECE. These specimens will contain the monitoring devices used in the Type I specimens, but the cathode area will be increased. This will be achieved by not only increasing the amount of steel in a single layer, but the number of layers will also be increased. In addition, some of the samples will include embedded pH electrodes to evaluate changes in alkalinity during ECE. An illustration of the proposed Type III specimen is shown in figure 27. Furthermore, to provide a baseline for this study, some of the new specimens will not contain chlorides.

Modeling the expected beneficial life of a treated structure will begin during the upcoming research period. This will be done using specimens that were treated during the extraction portion of this study. In addition, the possibility of evaluating previously treated structures is being explored.

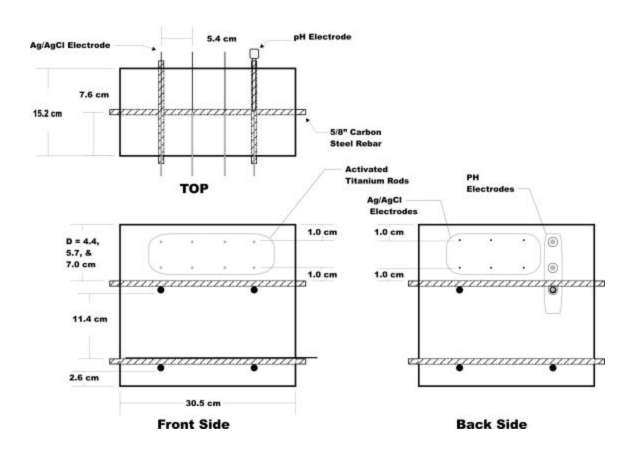


Figure 27. Proposed Type III specimen

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