



A Galvanic Corrosion Approach to Investigating Chromate Effects on Aluminum Alloy 2024-T3

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The effects of chromate treatment on the corrosion of an aluminum/copper aircraft alloy were examined with a galvanic corrosion apparatus composed of two electrodes and a zero resistance ammeter. Combinations of pure Al, pure Cu, and AA2024-T3 electrodes were immersed in 0.1 M NaCl solution, which was saturated with air, O₂, or argon. Selection of electrode material or differential aeration resulted in partial segregation of corrosion reactions, with one electrode becoming a net cathode and the other a net anode. For the case of an Al/Cu galvanic cell, chromate significantly reduced the observed cathodic current on Cu but had little effect when added to the Al cell. For an AA2024/AA2024 couple, chromate decreased the observed current when present in either the net anode or net cathode compartments. The results indicate that Cr^{VI} in solution, or Cr^{VI} pretreatment, inhibits O₂ reduction on Cu sites. Inhibition is preceded by a spike in the reduction current, implying that Cr^{III} is formed at approximately a monolayer level on Cu. The results for Cr^{VI} effects on the AA2024/AA2024 galvanic couple are consistent with observed polarization curves. Considering the results in their entirety, Cr^{VI} acts as a strong irreversible cathodic inhibitor on Cu and AA2024, but the results do not exclude the action of Cr^{VI} as an anodic inhibitor on AA2024-T3.
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Aluminum alloy 2024-T3 (AA2024-T3) has been well documented to be highly susceptible to corrosion, particularly localized pitting corrosion, and intergranular corrosion. AA2024-T3 is an inhomogeneous alloy (nominal composition Al, 3.8-4.9% Cu, 1.2-1.8% Mg, 0.3-0.9% Mn, 0.5% Fe, 0.5% Si, 0.25% Zn, 0.1% Cr, 0.05% Ti) containing several types of secondary-phase intermetallic particles that are enriched in alloying elements vs. the alloy matrix.¹ Explanations for increased corrosion susceptibility for AA2024-T3 center on the existence of these particles. Intermetallic particles greater than 0.2 μm have been characterized into four groups, accounting for about 85% of particle types, classified as Al₂CuMg (S-phase), Al₆(Cu,Mn,Fe), Al₇Cu₂Fe, and (Al,Cu)₆Mn phases.² The open-circuit potential (OCP) of many of these intermetallic particles is positive (or becomes positive after dealloying) vs. the alloy matrix phase.^{3,4} Subsequently, local galvanic couples occur with the intermetallic sites becoming cathodes. This may explain the observation of pitting of the alloy matrix around these intermetallic particles.^{1,5,6} Recent advances have revealed that S-phase particles dealloy, leaving mostly a copper sponge-like residue.^{1,2,7,8} Al₂CuMg particles have OCP values negative of the alloy matrix and thus are initially anodic.^{4,9} Another study using scanning Kelvin probe force microscopy found the Al₂CuMg particles to be more noble than the matrix in the as-polished condition due to a surface film.^{10,11} Film damage (either by scratching or prolonged exposure to chloride) caused the potential of the particles to become active relative to the alloy matrix and dissolution ensued. Mg and Al dissolution from these particles can result in the above-mentioned porous copper residue that is presumably highly cathodic. Copper from the intermetallic sites has been shown to migrate during the corrosion process and secondary corrosion occurs at sites where it is deposited.^{4,7} Whether this migration results from dissolution or deattachment is not clear.

Chromate conversion coatings (CCCs) have long been used to control corrosion of AA2024-T3 and other metal alloys. However, research has been ongoing to develop alternative corrosion inhibitors due to the toxicity of chromium compounds. A major area of study in this regard is the elucidation of the inhibition mechanism of chromium species, particularly the activity of Cr^{VI} as a cathodic or

anodic inhibitor, or both.^{3,12-19} The mechanism of CCC formation has been examined, revealing the reduction of solution-phase Cr^{VI} to form a Cr^{III}(OH)₃ film and subsequent incorporation of Cr^{VI} into the film as a Cr^{III}-Cr^{VI} mixed oxide.²⁰⁻²³ Furthermore, the release of the incorporated Cr^{VI} species into solution from the CCC has been demonstrated, as has migration to pits or defects.^{14,15} The pH and concentration-dependent release of Cr^{VI} aids in explaining the active inhibitor (or self-healing) properties of CCCs. A fundamental question that needs to be answered is whether chromium acts as an inhibitor of anodic and/or cathodic processes taking place during corrosion of the alloy. Anodic inhibition mechanisms of chromate have been studied extensively. Initial ideas of CCC inhibition centered on the formation of a Cr^{III} film that sealed the alloy surface from the corrosive electrolyte.^{21,24} In this model, cracks or holes formed in the film are plugged by Cr₂O₃ [or Cr(OH)₃] as a result of the reduction of Cr^{VI} from the CCC. Accumulation of a Cr^{VI}-Al^{III} mixed oxide has been observed in actively growing pits but reduction to Cr^{III} was not reported.²⁵ Others have presented research on pure aluminum indicating that CCC formation results in a positively charged surface that effectively repels aggressive chloride anions.²⁶⁻²⁸ However, solution-phase Cr^{VI} was not observed to reduce aluminum dissolution according to one study.¹⁶ The possibility of cathodic inhibition by chromium compounds has not been studied thoroughly for aluminum alloys. However, thin films of Cr^{III} formed on platinum and gold electrodes from reduction of Cr^{VI} effectively inhibited several reduction reactions (including oxygen).^{29,30} Sehgal *et al.* showed that the attack of AA2024-T3 at open circuit was eliminated by the introduction of a small amount of dichromate into the chloride solution, whereas large amounts of dichromate were needed to have a small effect at an applied potential just above the open-circuit value.¹⁹ This result implied strong cathodic inhibition by Cr^{VI}.

Difficulty in studying the corrosion of AA2024-T3 and its inhibition arises due to multiple reactions taking place in the same nominal surface area. Previous studies have utilized current density and potential mapping to study local phenomena on AA2024-T3.^{10,11,31} Additional understanding of the inhibition mechanism might be available by partial or complete segregation of the anodic and cathodic reactions. By employing separate coupled electrodes, galvanic currents may be studied as related to the individual reactions. Furthermore, a galvanic coupling experiment in which a segregated anode and cathode are connected but allowed to

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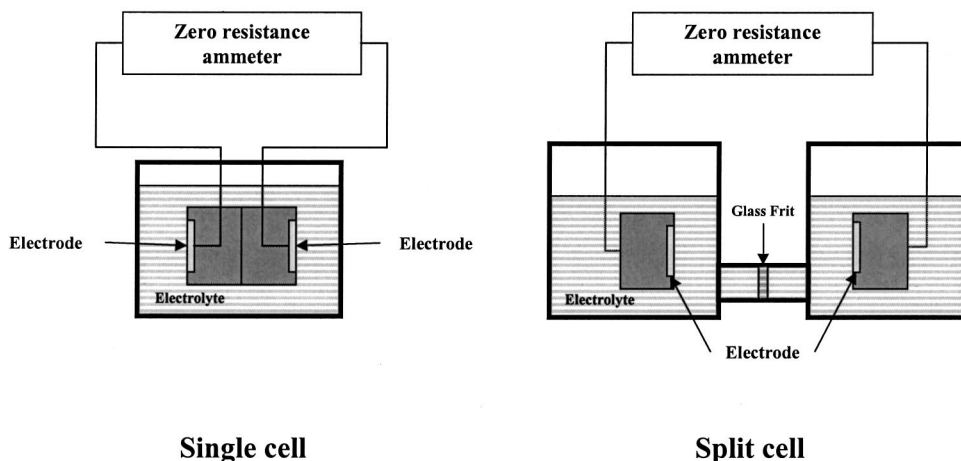


Figure 1. Schematics of single- and split-cell apparatus. Solution compartments could be saturated with Ar, O₂, or air, and Al, Cu, or AA2024-T3 electrodes were mounted in epoxy resin.

float without an externally applied potential approximates the mixed potential relevant to field conditions. Unlike the polarization curves often used to study inhibition, a galvanic coupling experiment does not impose an externally applied potential, which may represent a possibly large current source to drive corrosion. Major differences in chromate effects on AA2024 corrosion have been reported when foil penetration experiments at open circuit were compared to potentiostatic experiments involving an imposed potential.^{18,19} Previously, Liao and Wei used a segregated anode and cathode to establish relationships between the current density and cathode:anode area ratio for aluminum coupled to model alloys.³² The general approach involves two metal samples in separate compartments, with one sample acting as a net cathode and the other a net anode. When one electrode is pure Cu and the other pure Al, the cathodic and anodic reactions are largely segregated, and the observed current mimics the corrosion current in Al/Cu alloys. However, localized corrosion of Al or Al alloys in the form of pitting or grain boundary attack always involves some amount of local hydrogen evolution as a result of the low potential at the active site. The amount of hydrogen evolution has been measured to be about 15% of the anodic current.^{16,33} AA2024-T3/copper cells allow the cathodic reaction to proceed on both electrodes while anodic reactions occur mainly on AA2024. When both electrodes are made from AA2024, the cathodic reaction may be localized preferentially at one electrode by purging one side with oxygen and the other with argon. In the current work, various combinations of electrodes were studied in either a single-cell or split-cell configuration shown in Fig. 1. The split cell employs a porous glass frit that allows ion migration but restricts bulk mixing of the two solutions. The environments of the two chambers in the split cell may be altered independently, thus allowing for more selective control of the anodic and cathodic reactions. The galvanic current between the electrodes is measured with a potentiostat-based zero-resistance-ammeter (ZRA) similar to the continuous reading ZRA configuration reported by Jones^{34,35} and Devay *et al.*³⁶ The approach leads to an admittedly imperfect separation of anodic and cathodic reactions but does permit observations of Cr^{VI} effects on both processes.

Experimental

Barnstead NanoPure™ water with a resistivity of 18 MΩ cm was used for rinsing and solution preparation in all cases. Reagent-grade K₂Cr₂O₇ (Alfa Aesar) and Alodine™ 1200S powder (Henkel Corp.) were used to make Cr^{VI} solutions. Reagent-grade HNO₃ (Fisher Scientific) was used for pH adjustment. Electrodes were made from commercial Al (99.999%, 0.5 mm thick sheet, Alfa Aesar), Cu (99.99% 1.0 mm thick sheet, Alfa Aesar), and Cu (99.99%, 5 mm diam rod, Goodfellow Metals), and aluminum alloy 2024-T3 (0.025 in. thick sheet, ALCOA). Sheet samples were cut to 1 cm² in all cases, attached to copper wire with silver epoxy resin (SPI Supplies/

Structure Probe, Inc.), and embedded in epoxy (Buelher). They were then mechanically polished with successively finer Si:C papers (Buelher: 240, 400, 600, 800, 1200 grit), rinsed with water, and dried under a stream of hot air.

Sample pretreatment with Cr^{VI} solutions was performed in some experiments as noted in the text. For Alodine™ 1200S pretreatment the polished sample was immersed in Alodine™ 1200S pretreatment (7.6314 g/L with HNO₃ added to pH 1.25) for 5 min, rinsed with water, allowed to air-dry for 2 h, and used immediately. For Cr₂O₇²⁻ pretreatment the polished sample was immersed in 0.4 M K₂Cr₂O₇ for 2 h, rinsed with water, allowed to air-dry for 2 h, and used immediately.

Electrochemical measurements were performed using a Gamry Instruments PC3/300 potentiostat/galvanostat/ZRA with Framework™ (version 3.11) and DC105™ dc corrosion measurement software. Galvanic current experiments were performed in either single-cell or split-cell configuration as shown in Fig. 1. Galvanic current between the two electrodes and potential of the system was measured using the Gamry hardware in ZRA configuration. Potentiodynamic polarization experiments were performed using a standard three-electrode electrochemical cell. Experiments utilized Ag/AgCl as the reference electrode (all potentials quoted are with respect to this reference), Pt wire as the counter electrode, and 0.1 M NaCl as the electrolyte. Compressed O₂ and Ar (prepurified) were used for aeration/deaeration studies. Solution additions were made using 0.05 M K₂Cr₂O₇ [0.1 M total Cr^{VI}] in 0.1 M NaCl as noted in the text and figures. The pH of NaCl solutions preceding Cr^{VI} addition was uncontrolled, but was typically 5.5-6.0, and decreased by less than 1 unit upon addition of Cr^{VI}. Injection experiments with controlled pH³⁷ showed only minor differences in galvanic response for pH 4-6.

Results

The “single” cell (Fig. 1) with pure Al and Cu electrodes was examined initially, since they should provide the greatest segregation of anodic and cathodic reactions. The effects of convection, aeration, and inhibition by Cr^{VI} addition on the current for the Al/Cu cell are shown in Fig. 2. A stable current of about -30 μA and mixed potential of about -650 mV was initially established under quiescent conditions. Negative current refers to electron flow from aluminum to copper. Solution stirring and oxygen saturation caused an increase in the current by an order of magnitude while the potential increased to about -550 mV. Deaeration of the solution with argon caused the current to decrease to a level similar to that of the quiescent solution even with solution stirring, while the potential decreased to about -750 mV. Reintroducing oxygen to the system returned the current and potential to previous levels. Addition of K₂Cr₂O₇ (to a total of 50 mM Cr^{VI}) to the system resulted in an

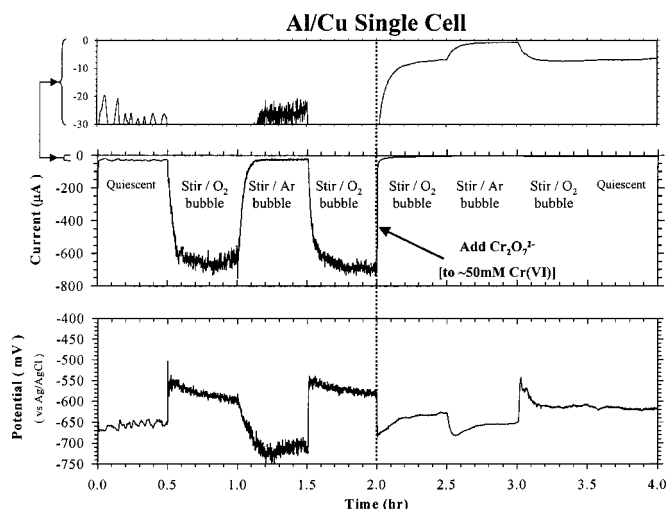


Figure 2. Current and potential vs. time plots for single cell containing pure Al (1 cm²) and pure Cu (1 cm²) electrodes, saturated with Ar or O₂ as indicated. Stirring by gas bubbles was rapid except when “quiescent”. At $t = 2.0$ h, K₂Cr₂O₇ solution was added to bring the total [Cr^{VI}] to 50 mM.

immediate reduction in the galvanic current. Even with oxygen saturation and stirring, the current was below that of the quiescent or deaerated solution prior to Cr^{VI} addition. Closer examination of the current reveals that the same trends occurred for aeration/deaeration and stirring as observed for the uninhibited system but at greatly reduced current levels. The potential decreased after Cr^{VI} was added but gradually returned to -650 mV. After changing from Ar to O₂ saturation or back, the potential slowly returned to ~ -650 mV. In addition, both the current and potential exhibited less noise after Cr^{VI} addition.

The effect of electrode pretreatment with Cr^{VI} was studied using a quiescent single-cell aluminum-copper couple. Experiments were run in three configurations. Either the copper or the aluminum electrode was pretreated with Cr^{VI} with the other being polished but not pretreated. These results were compared to a control experiment in which neither of the electrodes was pretreated. Results of Alodine™ 1200S pretreatment are presented in Fig. 3. Pretreatment of the copper electrode with Alodine 1200S resulted in an initial galvanic cur-

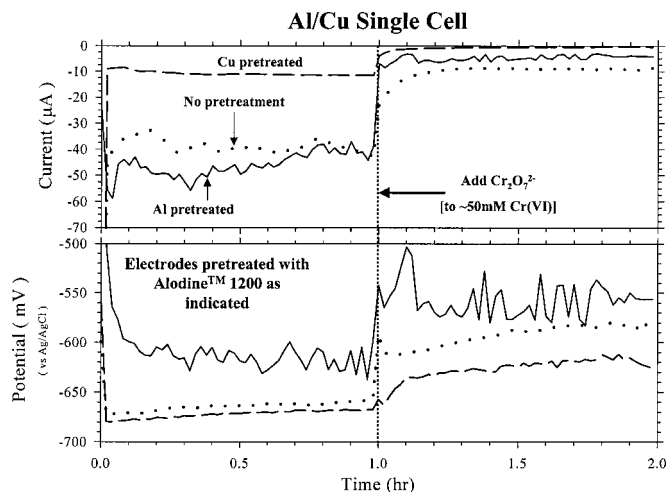


Figure 3. Behavior of the Al/Cu single cell following pretreatment of either electrode in Alodine 1200S for 5 min. The cell solution was air-saturated. At $t = 1.0$ h, K₂Cr₂O₇ was added and stirred to yield a final [Cr^{VI}] of 50 mM, after which the solution was quiescent.

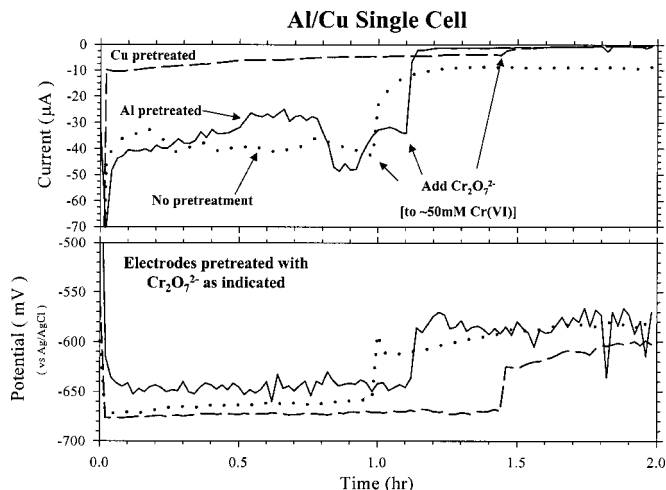


Figure 4. Same as Fig. 3, but samples pretreated by immersion in 0.40 M K₂Cr₂O₇ for 2 h. K₂Cr₂O₇ was added to the solution at the times indicated.

rent significantly lower than the case for pretreatment of the aluminum electrode. Currents after Alodine pretreatment of the aluminum electrode were similar to the control, in which neither electrode was exposed to Cr^{VI}. When K₂Cr₂O₇ was added to the solution to a final concentration of 50 mM Cr^{VI}, the galvanic current decreased in all cases but by varying amounts. Pretreatment in 0.4 M K₂Cr₂O₇ instead of Alodine 1200S yielded very similar behavior, shown in Fig. 4.

Split-cell experiments using an aluminum-copper couple were employed to further examine the individual anodic and cathodic reactions and current inhibition. The galvanic current and potential were monitored for aluminum-copper couples under varying conditions of mass transport and aeration. Employment of the split cell allowed stirring and aeration/deaeration to be studied in either the anodic or cathodic reaction chamber. The separation of anodic and cathodic reactions is illustrated in Fig. 5 for the Al/Cu split cell. Alternation of Ar and O₂ saturation in the copper side caused large variations in current, with a negative current again referring to electron flow from aluminum to copper. The same alternation of Ar and O₂ on the Al side had no observable effect, indicating that O₂ reduction was not active on aluminum, presumably due to the passive

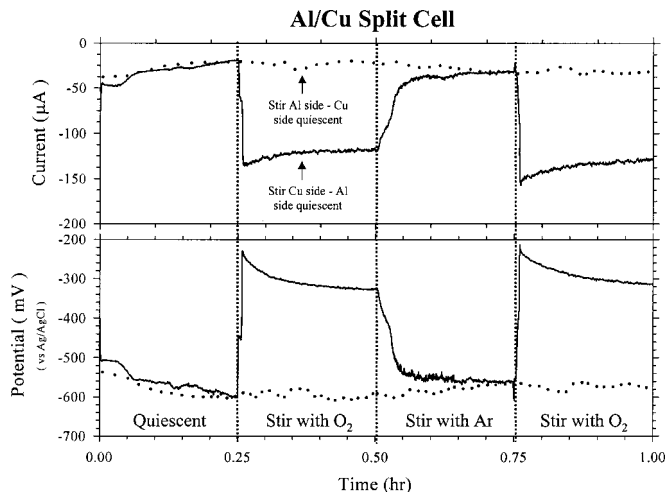


Figure 5. Current and potential behavior of split cell containing Al and Cu electrodes with O₂ bubbling of either the (· · · ·) Al or (—) Cu compartment as indicated. In the first (left) panel, both compartments were quiescent.

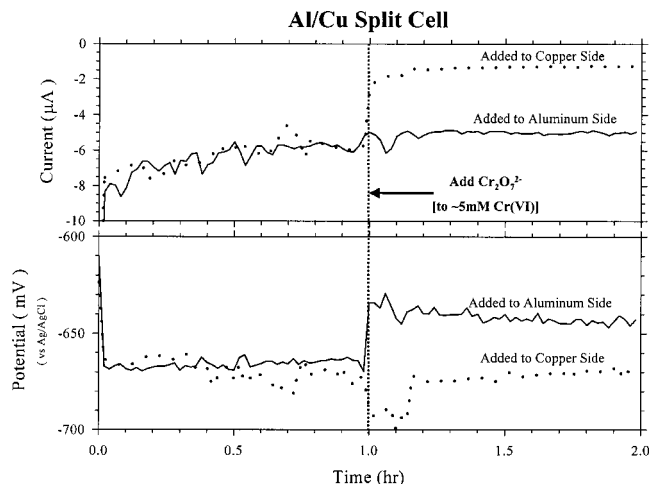


Figure 6. Effect of Cr^{VI} addition (to 5 M total Cr^{VI}) to either the Al or Cu compartments of the split cell described in Fig. 5. Both compartments were air-saturated but quiescent throughout the experiment, except for a brief mixing period after Cr^{VI} addition. For this figure only, the Cu area was 0.05 cm² and the Al remained at 1 cm².

oxide film. The effect of solution Cr^{VI} on the cathodic and anodic reactions of an aluminum-copper galvanic couple using the split-cell configuration are shown in Fig. 6. After 1 h of galvanic corrosion under quiescent conditions, K₂Cr₂O₇ was added (to ~5 mM Cr^{VI}) to either the aluminum or copper half-cell. The galvanic current was inhibited when Cr^{VI} was added to the copper cell, but when Cr^{VI} was added to the aluminum cell the current remained essentially constant. The experiment was repeated with fresh electrodes and aerated solutions, with incremental addition of K₂Cr₂O₇ to the Cu compartment. After stabilization of the current to -58 µA, the addition of Cr₂O₇²⁻ to yield total Cr^{VI} concentrations of 1, 10, and 25 mM decreased the current to -18, -6, and -6 µA, respectively. For the same additions, the potential decreased from -575 mV to -650, -680, and -680 mV.

Figure 7 compares current and potential profiles for a Cu cathode in aerated solution and either Al or AA2024-T3 in quiescent, aerated 0.1 M NaCl. After 30 min of galvanic corrosion, K₂Cr₂O₇ was added (to ~5 mM Cr^{VI}) to the aluminum or alloy chamber of the

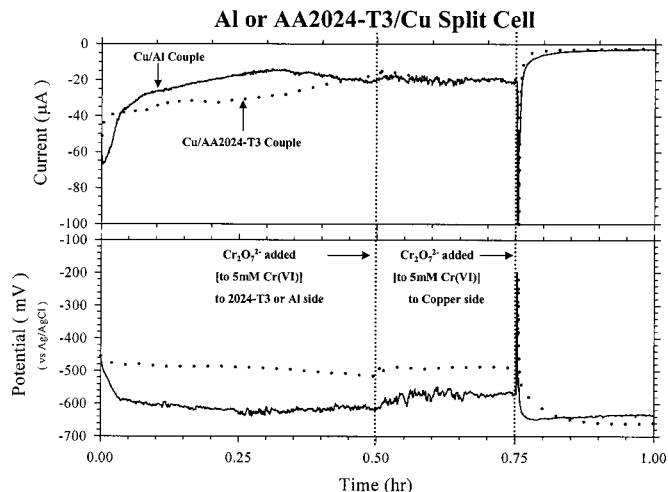


Figure 7. Behavior of split cell containing a Cu electrode in one compartment and either Al or AA2024-T3 in the other. Both compartments were aerated but not agitated. K₂Cr₂O₇ was added as indicated to a total Cr^{VI} of 5 mM.

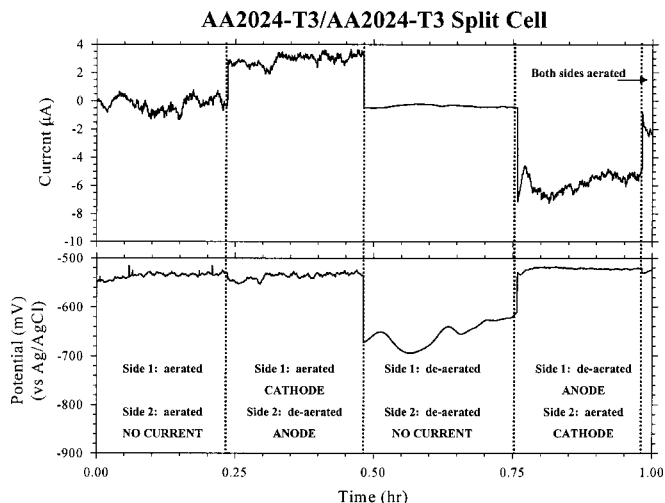


Figure 8. Behavior of split cell with AA2024 electrodes in both compartments, but with aeration of one, both, or neither compartment as indicated.

split cell. Fifteen minutes later K₂Cr₂O₇ was added to the copper-containing chamber (to ~5 mM Cr^{VI}). The results were similar when the copper cathode was coupled to either aluminum or AA2024-T3. The galvanic current plots for either an Al or AA2024 anode track each other for the entire experiment. Little change in current was observed for either couple when Cr^{VI} was added to the anodic chamber. Predictably, the current dropped dramatically when Cr^{VI} was introduced to the copper-containing chamber. The mixed potential of the copper/AA2024-T3 couple was initially higher than the copper/aluminum couple, as expected, since the measured OCP for AA2024-T3 is higher than for pure aluminum. A large current spike was observed immediately upon addition of the Cr^{VI} to the Cu chamber before the current decreased below initial values. The transient was not observed when Cr^{VI} was added to the AA2024 or Al chamber. The spike was not observable in previous experiments (Fig. 2-6), because data acquisition was paused during Cr^{VI} addition.

With AA2024 in both sides of the split cell, galvanic corrosion current may be generated by saturating one side with air and the other with Ar. The aerated side becomes a net cathode, and the Ar side becomes a net anode. This approach is similar to differential aeration cells reported previously to study galvanic currents caused by concentration differences.³⁵ Figure 8 demonstrates the ability to cycle the chambers between being net anodes and net cathodes by air or Ar saturation. Initially both chambers of the split cell were aerated and the resulting current fluctuated around zero under quiescent conditions. Presumably this fluctuation was caused by local corrosion events on one electrode momentarily polarizing it either positive or negative of the other electrode. Argon saturation of side 2 of the split cell yielded positive current (e⁻ flow toward the aerated side), and deaeration of both sides returned the current to near zero. Since both sides were deaerated, local corrosion events were minimized and the resulting “zero” current was much smoother than with both sides aerated. The mixed potential decreased as would be expected with lower oxygen concentration. As air was reintroduced into side 2, the current became negative as electrons flowed toward the aerated electrode, and the potential returned to the range of -525 to -550 mV. The mixed potential was in this range when either or both sides were aerated and only decreased when both sides were deaerated. Inspection of both electrodes in a 2024/2024 split cell after 6 h of corrosion with differential aeration using light microscopy revealed pits on both the deaerated and aerated, with the pitting and damage being noticeably greater on the aerated side.

Figures 9 and 10 show the effect of Cr^{VI} addition on the deaerated and aerated chambers of the split cell. In both experiments a

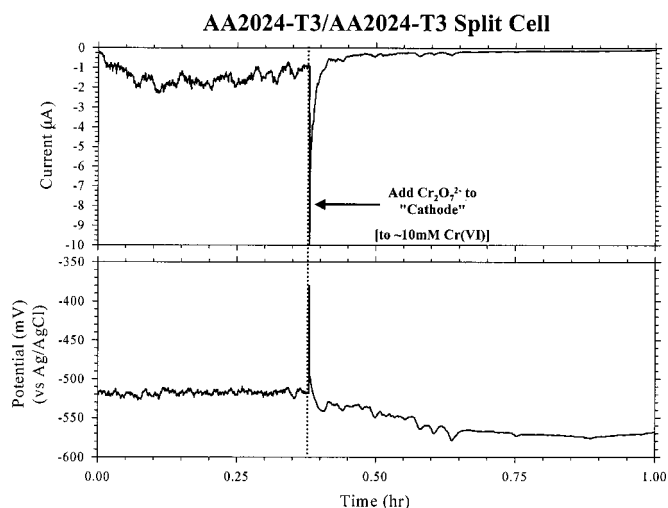


Figure 9. Effect of Cr^{VI} addition to aerated side ("cathode") of a split cell containing AA2024 electrodes on both sides. The opposite side was Ar-saturated, but neither side was stirred except immediately after Cr^{VI} addition.

current is generated by deaeration of one side as shown in Fig. 8. Addition of Cr^{VI} (to ~ 10 mM) to either the aerated (Fig. 9) or deaerated (Fig. 10) chambers caused a decrease in galvanic current, although a current spike was observed only for addition to the aerated chamber. Microscopy of both electrodes after 6 h of corrosion showed that Cr^{VI} addition significantly reduced corrosion damage when added to either the deaerated or aerated side.

Figures 11 and 12 show polarization curves for Cu and AA2024 for conditions relevant to the split-cell experiments. Figure 11 shows the effects of Cr^{VI} addition to aerated chloride solutions on the polarization curves for Cu and AA2024. The corrosion potential for AA2024 in aerated chloride solution with no Cr^{VI} was essentially pinned at the breakdown potential due to the relative nonpolarizability of the localized corrosion reaction. In the Cr^{VI} -containing solution, the OCP was lower as a result of cathodic inhibition, and the breakdown potential was higher. In the aerated Cr^{VI} -free chloride solution, the 1 cm^2 Cu electrode exhibited a mass-transport-limited O_2 reduction current of $40 \mu\text{A}/\text{cm}^2$ over a wide potential range between -0.30 and $-0.65 \text{ V vs. Ag/AgCl}$. In the Cr^{VI} -containing solution, the rate of the cathodic reaction on Cu was two to three

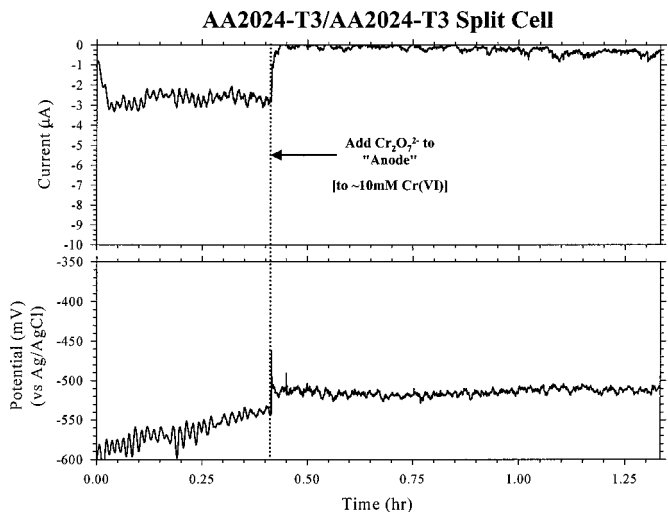


Figure 10. Same as Fig. 9 but with Cr^{VI} addition to the deaerated chamber ("anode").

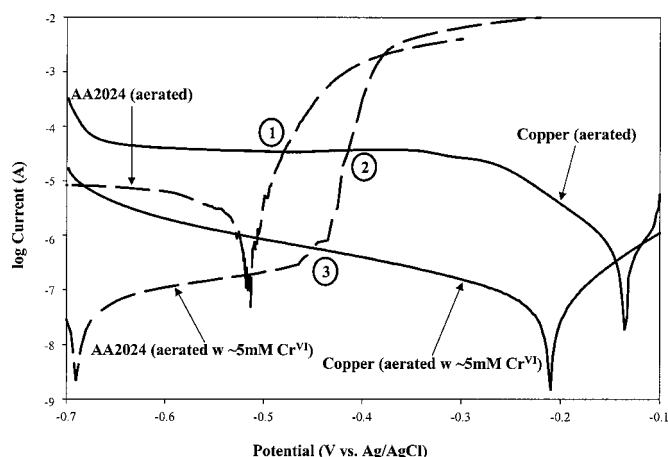


Figure 11. Potentiodynamic polarization curves for Cu and AA2024 electrodes in 0.1 M NaCl for the conditions shown. All electrode areas were 1 cm^2 . In most cases, samples were held at open circuit for 10 min prior to initiating a scan at 1 mV/s in the positive direction from a potential below the OCP. The exception was aerated AA2024, which was initiated at the OCP and scanned in the positive or negative direction using different samples. Labeled intersection points are described in the text.

orders of magnitude lower. The rate of the cathodic reaction on AA2024 in the aerated Cr^{VI} -free solution was also lower than that for Cu in the same solution, and even lower in the Cr^{VI} -containing solution.

Figure 12 shows the polarization curves relevant to the 2024/2024 split-cell experiments of Fig. 9 and 10. The two curves for 2024 in aerated solutions are reproduced from Fig. 11, and the curves for AA2024 in deaerated solution are added. Deaeration or Cr^{VI} addition resulted in a decrease in the cathodic current. The addition of Cr^{VI} had less effect on the cathodic portion of the curve for AA2024 in the absence of oxygen. The passive current densities were decreased and the breakdown potentials slightly increased by the addition of Cr^{VI} .

Discussion

The objective of partial or complete separation of anodic and cathodic corrosion reactions was achieved with either the single-cell design with different electrode materials or the split cell with differential aeration. This separation permitted the localization of inhibitor effects and insights into the inhibition mechanism. However, spatial

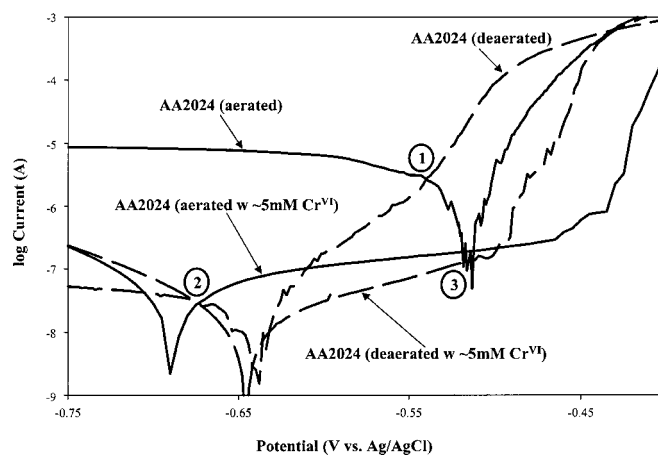


Figure 12. Potentiodynamic polarization curves for AA2024 electrodes in 0.1 M NaCl for the conditions shown. Conditions the same as those for Fig. 11.

separation of the net anode and net cathode necessarily decreases chemical interactions between surface sites, which might occur in real systems when anodic and cathodic regions are in close proximity. For example, a pH change occurring at a local cathode may influence the corrosion reactions at an adjacent anode. In order to localize inhibitor effects to the anode or cathode, these chemical interactions must be neglected. With that caveat in mind, several useful observations are available from the current results and are considered in turn.

The separation of anodic and cathodic reaction for the case of pure Cu and pure Al electrodes is apparent for both the single cell (Fig. 2) and split cell (Fig. 5) and results in the observed net corrosion current. For the Al/Cu case, the observed current was controlled by the availability of O₂ at the Cu electrode and was increased by faster mass transport. Cr^{VI} in solution significantly decreased the current but only if it could interact with the Cu electrode. In the split cell, Cr^{VI} addition to the Al side had no effect on the current, although it increased the potential by about 20 mV (Fig. 6). Pretreatment of Cu or Al electrodes with either Alodine or Cr₂O₇²⁻ solution (Fig. 3) had similar effects to addition of Cr^{VI} on the galvanic corrosion cell(s). Pretreatment of Cu significantly suppressed the corrosion current, while pretreatment of Al had no effect on current and a small positive effect on potential. Taken together, the Al/Cu experiments indicate that the observed galvanic current is limited by O₂ reduction and that Cr^{VI} reduces the rate of O₂ reduction. At least for pure Al, Cr^{VI} had no effect on the anodic reaction rate. The observations on the Al/Cu system serve as important background information for the more realistic AA2024/Cu and AA2024/AA2024 combinations.

Separation of anodic and cathodic reactions is quite clear for the case of the Cu/Al couple, but cells with AA2024-T3 as one or both electrodes are likely to exhibit local galvanic couples between anodic and cathodic sites on AA2024. Removal of O₂ from one or both half-cells suppresses the cathodic reactions, and the observed galvanic current should be significantly larger than the local current. Based on the observed galvanic current, we identify electrodes as "net cathodes" or "net anodes". The fact that the net cathode shows more visual damage than the net anode in the AA2024/AA2024 split cell in the absence of Cr^{VI} indicates that local reactions are significant in this case. The greater damage to the cathode may result from the generation of hydroxide ions near local cathodes, which then etch aluminum near the local cathode. Keeping the issue of imperfect separation in mind, the comparison of the Cu/Al and Cu/AA2024 cells in Fig. 7 shows that the alloy behaved similarly to Al when acting as a net anode. Cr^{VI} addition to the Al or AA2024 chamber had minor effects on galvanic current, while Cr^{VI} inhibited the net cathodic reaction on Cu in both cases. The current spike observed upon Cr^{VI} addition to the Cu chamber implies a transient reduction of Cr^{VI}, and the area of the spike corresponds to approximately 2×10^{-9} mol Cr^{VI} per cm² of geometric copper area. No current spike was observed upon the addition of a NaCl solution containing no Cr^{VI}, indicating that the spike in fact resulted from the reduction of Cr^{VI}. With one prominent exception, the results for the AA2024/AA2024 split cell shown in Fig. 9 and 10 are consistent with those observed for Al/Cu and Cu/AA2024. Cr^{VI} inhibits the observed current when added to the aerated side of the AA2024/AA2024 split cell (Fig. 9), similar to what was observed when it was added to the Cu side of the Cu/AA2024 split cell. This again implies a cathodic inhibition mechanism for Cr^{VI}. However, Cr^{VI} also reduces the current when added to the deaerated or "anodic" side of the AA2024/AA2024 split cell, Fig. 10. The apparent activity of Cr^{VI} as an anodic inhibitor, which is implied by Fig. 10, is at odds with Fig. 7, in which no effect was observed on the anode when the cathode was Cu instead of AA2024. It also seems to contradict other results indicating that Cr^{VI} is not an effective inhibitor for pit growth.^{18,19}

An understanding of this behavior is provided by close examination of the polarization curves in Fig. 11 and 12. With a large Cu cathode, the observed corrosion current is determined by the mass-transfer-limited reduction of O₂ on Cu. The observed current and potential in the split cell (Fig. 7) are approximately equal to those predicted from Fig. 11 at the point marked "1". Under this condition, the Cu cathode polarized the AA2024 anode into a potential range well above the breakdown potential where localized corrosion occurred at a relatively high rate. In the galvanic corrosion condition of the split cell, the rate of the localized attack was limited by the combined cathodic reactions on the Cu and AA2024 surfaces. The breakdown potential for AA2024 is slightly higher in the aerated Cr^{VI}-containing solution. However, when connected to the aerated Cu cathode in the split cell the intersection (point 2 in Fig. 11) was still well above the breakdown potential, and the rate was still dictated by the rate of oxygen reduction on Cu, which remained at the mass-transport-limited rate. There was little change in the current and a small positive shift in potential occurred when Cr^{VI} was added to the AA2024 side, as observed in Fig. 7. The rate of the cathodic reaction on Cu was considerably lower in the Cr^{VI}-containing solution, which limited the galvanic current when Cu was connected to AA2024 in a solution either with or without Cr^{VI}. The shift from point 2 to 3 in Fig. 11 represents the situation when Cr^{VI} was added to the Cu side in Fig. 7.

The effect of Cr^{VI} in the AA2024/AA2024 split-cell experiments of Fig. 9 and 10 can be understood with the help of the polarization curves in Fig. 12. The addition of 5 mM Cr^{VI} increased the pitting potential of AA2024 in deaerated 0.1 M NaCl by a small amount and decreased the passive current density. The increase in pitting potential was somewhat larger in the aerated solution. These observations are evidence of a form of anodic inhibition. The polarization curve for AA2024 in the deaerated chloride solution does not show a very sharp breakdown, and galvanic coupling with an equal area of AA2024 in aerated solution brings the potential close to the inflection point in the polarization curve (point 1 in Fig. 12). When Cr^{VI} was added to the aerated (cathodic) side of the split cell, the potential and current decreased, as shown in Fig. 9. The polarization curves predict that the potential should decrease to point 2 and that the current should decrease to the extent that the electrode polarity reverses, *i.e.*, the AA2024 sample in the aerated chloride +Cr^{VI} solution should become the anode. This extent of inhibition was not observed in the split-cell experiment, but a strong cathodic inhibition was evident.

The polarization curves in Fig. 12 also predict that the steady-state point should have moved from point 1 to point 3 in Fig. 12 when Cr^{VI} was added to the deaerated side of the AA2024/AA2024 split cell. This potential is still close to the breakdown potential of both electrodes, but the passive current density in the deaerated chloride +Cr^{VI} solution is lower than in the solution with no Cr^{VI}, and is about equal to the corrosion rate in the aerated chloride solution. The net current passing between the electrodes should be much lower and the potential should increase upon addition of Cr^{VI}, and that is in fact what was observed in Fig. 10. This observation does not contradict the prior reports that dichromate does not inhibit pit growth if one considers that sustained pit growth was not possible at point 1 in Fig. 12 where AA2024 in aerated and deaerated solutions were coupled. The decrease in current reflects a decrease in the passive current density with the addition of Cr^{VI}.

The activity of Cr^{VI} as both an anodic and cathodic inhibitor has been reported previously by several groups,^{3,12-19} and there is continuing controversy about which action, if any, is dominant in corrosion protection of AA2024-T3. Ilevbare and Scully have reported that Cr^{VI} inhibits dealloying of s-phase particles, thus reducing formation of Cu-rich cathodic sites.^{3,12,13} In the current work, however, the AA2024-T3 was exposed to NaCl solution for approximately 1 h before Cr^{VI} injection (during which dealloying presumably occurred), and Cr^{VI} still inhibited the galvanic current. Although Cr^{VI} may inhibit dealloying in field applications, it also acts as a cathodic

inhibitor on cathodic sites formed by dealloying or present initially. Furthermore, Cr^{VI} inhibited O₂ reduction on pure Cu, which represents the upper limit of Cu dealloying and redistribution. The current results indicate that Cr^{VI} does act as an anodic inhibitor, in particular by inhibiting the initiation of localized corrosion rather than growth. However, they also establish that Cr^{VI} is a strong and irreversible cathodic inhibitor, which can play a dominant role in the stabilization of localized attack.

Preliminary examination of the size and nature of the current spikes observed upon Cr^{VI} addition to the cathode compartment indicate that the spike area is approximately proportional to the area of exposed copper. In addition, the spike area of approximately 10⁻⁴ C/cm² of copper is consistent with the deposition of roughly a monolayer of Cr^{III} oxyhydroxide on exposed metallic copper. The deposition of Cr^{III} on Cu and AA2024-T3 and its action to inhibit O₂ reduction is the subject of a future report.

Conclusions

To better understand corrosion and its inhibition for aluminum alloy 2024-T3, an approach was taken wherein the individual anodic and cathodic reactions were partially separated and studied using various galvanic couples. The use of a split-cell configuration allows segregation of anodic and cathodic reactions and consideration of the effect of chromate on each reaction independently.

Specific conclusions are

1. Partial or complete segregation of anodic and cathodic reactions was achieved either by choice of electrode material or by differential aeration.
2. Dilute chromate solution or pretreatment with Cr^{VI} solution greatly decreased the O₂ reduction rate on both Cu and AA2024-T3.
3. Cathodic inhibition on Cu surfaces by Cr^{VI} solution was accompanied by a transient reduction current equivalent to generation of approximately a monolayer of Cr^{III} oxyhydroxide.
4. Anodic inhibition by Cr^{VI} was observed for the AA2024/AA2024 galvanic couple but not for AA2024/Cu or Al/Cu couples. This apparent discrepancy is consistent with observed polarization curves for AA2024-T3.

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