Experiments were conducted to determine how inhibiting forms of vanadate interact with complex Al alloy microconstituent intermetallics to impart corrosion protection. Cathodic polarization experiments on Al 2024-T3 indicate a strong correlation between inhibition and the presence of tetrahedrally coordinated vanadate. Anodic and cathodic polarization curves were measured on bulk synthesized AlCu, AlCuMg, AlCuFe, and AlCuMgMn in alkaline 0.5 M NaCl solutions with and without 10 mM NaVO3. Vanadate additions generally decreased Ecorr, increased Epit, and decreased the cathodic kinetics of all tested materials. Because of decreased cathodic kinetics, open-circuit potentials (OCPs) were shifted in the active direction in aerated solutions when vanadate was present. This shift pins the OCP just below the observed potting potential for AlCuMg in vanadate solution, effectively preventing breakdown and subsequent support of rapid oxygen reduction by Cu-enriched clusters. Ecorr, Epit, Ecorr – Epit, tcorr, and f at –1.3 V SCE data from polarization experiments were summarized in cumulative distribution plots, and averages are presented in tabulated format. Scanning electron microscopy images of Al 2024-T3 used for 4 h of OCP measurement show that vanadate greatly decreased circumferential trenching in intermetallic particles in both aerated and deaerated solutions. Potentiostatic hold experiments were used to show suppression of AlCuMg dissolution in vanadate solutions.

Manuscript submitted October 6, 2008; revised manuscript received January 5, 2009. Published February 9, 2009.

Aluminum 2024-T3 is a high-strength age-hardened aluminum alloy commonly used in the aerospace industry. Al 2024-T3 contains, by weight percent, 3.8–4.9 Cu, 1.2–1.8 Mg, 0.3–0.9 Mn, and small quantities of Si, Fe, Zn, Cr, and Ti. Alloy additions result in both superior mechanical properties and a heterogeneous microstructure which renders the alloy susceptible to localized corrosion. Appreciable quantities of copper, magnesium, and manganese, added as strengtheners, remain in solid solution. However, through heat-treatments and natural aging, a dispersion of fine Cu and Mg particles and insoluble intermetallic precipitates form within the matrix phase. The main constituent particles in Al 2024-T3 include AlCuMg, AlCuFe, AlCu, and AlCuMgMn. The effect of intermetallic particles on corrosion of aluminum alloys has been widely studied. For Al 2024-T3, it has been found that intermetallics containing Cu are typically noble or become noble to the surrounding aluminum matrix during exposure to many electrolytes, and these particles are capable of supporting rapid cathodic kinetics. Such cathodic particles drive corrosion in the surrounding matrix, leading to pitting and trenching attack morphologies. AlCuMg (S phase) intermetallic particles are of particular interest, because AlCuMg is one of the most abundant intermetallic particles found in Al 2024-T3 and in large part has been found to be responsible for susceptibility of Al 2024-T3 to localized corrosion. The corrosion of AlCuMg is complex; under free-corrosion conditions, the intermetallic is initially anodically polarized by the matrix, leading to selective dissolution of Mg from the intermetallic and nonfaradaic liberation of Cu, which can then be oxidized to form ions that can be reduced on the surrounding matrix. Often what remains of the particle is an enriched Cu remnant, which acts as a local cathode, supporting rapid oxygen reduction and corrosion in the surrounding matrix. Prevention of Mg dissolution from AlCuMg and, as a result, the subsequent formation of local Cu cathodes capable of supporting rapid oxygen reduction could be an effective way to increase the resistance of Al 2024-T3 to localized corrosion.

Historically, chromate-based pigments and coatings have been used successfully to prevent corrosion of aluminum alloys. However, due to environmental and carcinogenic risks associated with chromate use, more "green" alternative inhibitors and coatings have recently received attention. In particular, soluble vanadates, vanadate-based coatings, and inhibitor pigments have been observed to inhibit the corrosion of aluminum alloys and have shown promise as chromate replacements.

Vanadates have a relatively complex aqueous chemistry, dependent on pH, concentration, and ionic strength. This convolutes a straightforward understanding of inhibition. In a simplified description of aqueous vanadate speciation, tetrahedrally coordinated species, metavanadates, and pyrovanadates, predominate in alkaline solutions, octahedrally coordinated species, decavanadates, predominate in acid solutions, and single tetrahedral species exist over a wide pH range at low concentrations. Previous work has shown that the extent of inhibition depends strongly on vanadate speciation, with the greatest inhibition from tetrahedrally coordinated species, which are predominant in alkaline solutions. Tetrahedrally coordinated vanadates have been shown to act primarily through decreased oxygen reduction; however, vanadates have also been observed to be modest anodic inhibitors independent of aeration. Decavanadate ions, which are combinations of 10 octahedrally coordinated vanadate units, are predominant in acidic solutions of appropriate vanadium concentration and have been shown to be poor inhibitors of oxygen reduction. There is evidence that decavanadate increases the cathodic kinetics in acidic NaCl solutions. However, small increases in pH, as found near sites supporting oxygen reduction, can trigger the decomposition of noninhibiting octahedrally coordinated species into inhibiting tetrahedral species, which helps explain corrosion protection observed from pigments containing decavanadate.

Evidence exists that inhibiting tetrahedrally coordinated vanadates suppress the dissolution of AlCuMg intermetallics. Ralston et al. noted suppressed Mg dissolution from Cu–Mg particles exposed to alkaline and mildly acidic 50 mM NaCl solutions with NaVO3 compared to particles exposed to NaVO3-free solutions. Iannuzzi and Frankel used in situ atomic force microscopy scratching to observe that additions as small as 0.1 mM of metavanadate to 0.5 M NaCl suppressed the attack of AlCuMg particles, while corrosion in the surrounding matrix was still observed. Iannuzzi further found that 5 mM of metavanadate prevented transient AlCuMg dissolution, resulting in increased corrosion resistance at open-circuit potential (OCP). The mechanism of suppression of transient dissolution was not clear, but it was speculated that monovanadates on the matrix prevent or displace Cl− adsorption on the surface, which hinders subsequent oxide film breakdown. Generally, it is not certain whether vanadates slow
corrosion of Al 2024-T3 through acting exclusively on Al-CuMg or if suppressed Al-CuMg dissolution is a consequence of overall corrosion inhibition. Although previous work has established that solutions containing predominately tetrahedrally coordinated vanadates prevent or slow Mg dissolution from Al-CuMg, and in turn the formation of Cu-rich cathodes capable of supporting rapid oxygen reduction, the precise relationship between tetrahedral vanadates, Al2CuMg, and the matrix is not currently understood.

Vanadate is a known buffer, and observed inhibition must be rationalized in the context of the effects that buffers have on corrosion. The presence of a buffer can have pronounced effects on the corrosion of Al-Cu alloys.30,31 In unbuffered systems, oxygen reduction results in an increase in alkalinity at cathodic sites, which dissolves the surrounding Al matrix, leading to shallow grooving and trenching around intermetallics and, occasionally, widespread cathodic corrosion across the matrix.30,31 When a buffer is present, local alkalization and associated cathodic corrosion damage modes are suppressed. However, when these modes are suppressed, the cathodic reaction is then available to support penetrating localized corrosion sites such as acid pits and crevices.30,31 This leads to deep and discrete pits but comparatively less mass loss than is observed in unbuffered systems.30

Due to the small size of constituent intermetallic particles present in Al 2024-T3, direct electrochemical testing of different intermetallics in the matrix is not feasible. However, previous work on aluminum alloys using a microcapillary electrode and synthesized “bulk” intermetallics demonstrates that intermetallic-specific electrochemical data can be obtained.2,3,32 The microcell is a modified standard three-electrode setup that uses a thin, glass, silicon-coated capillary connected to an electrolyte reservoir, containing both a reference electrode and counter electrode to contact and allow electrochemical experiments on micrometer-scale-diameter working electrodes.33,34 General details of the microcell setup and a specific description of the microcell used for this work can be found in the literature.3,33,34

By choosing bulk intermetallics that are representative of constituents in Al 2024-T3, pure Al and Cu, and an Al 4% Cu solid solution used as a matrix analog, electrochemical characteristics of specific intermetallic phases can be catalogued and used to rationalize observed behavior of the bulk alloy. The objective of this work is to determine how inhibiting vanadates interact with the matrix and constituent particles of Al 2024-T3. In addition, this work aims to develop a deeper understanding of the suppression of Mg dissolution from Al-CuMg intermetallics in inhibiting vanadate solutions.

Experimental

Solution preparation.— Solutions for all experiments were prepared using reagent-grade chemicals. The NaVO3 for solution preparation was purchased from Fluka Chemika with an assay of ≥98%. Cathodic polarization experiments were conducted in 0.5 M NaCl solutions adjusted to pH 5.1 using HCl with 0.25 and 0.0025 M NaVO3 to show the effect that tetrahedral vanadates have on inhibition compared to octahedral vanadates. However, most experiments were conducted in alkaline 0.5 M NaCl solutions with and without 10 mM NaVO3 to characterize the inhibitive effect tetrahedrally coordinated vanadates have on different constituent intermetallics in Al 2024-T3. The initial as-dissolved 0.5 M NaCl + 10 mM NaVO3 solution was yellow and had a pH of 6.37. As previously mentioned, vanadates have been shown to provide a sufficient measure of solution and species stability. Although the NaVO3 solution remained stable, the pH of NaCl solutions adjusted to approximately pH 9.2 decreased with time. This is likely the result of H2CO3 formation as a consequence of the subsequent proton formation from equilibria involving HCO3− and CO32−, which both have increased solubility in alkaline solutions.35

As a result, care was taken to monitor and measure the pH immediately prior to and during each testing session. Unless specifically stated, the pH of NaCl-only solutions was between 9.05 and 9.20. Additionally, the reservoir, capillary, and tubing of the microcell were frequently flushed with fresh solution throughout experimentation.

NMR.—Vanadates have a complex aqueous speciation depending on both concentration and pH, and as a result, small changes in pH can have dramatic effects on the type and concentration of species in solution. NMR was used to characterize solutions used for cathodic polarization experiments on an Al 2024-T3 sheet in pH 5.1 NaVO3 solutions and microcell polarization experiments in alkaline 10 mM NaVO3 solution, which were expected to take a number of weeks to complete and for which the possibility of solution evolution with time was a concern. After allowing approximately 2 weeks for stabilization, NMR spectra were collected immediately prior to microcell work in vanadate solutions and 10 days later after completion of experimentation. NMR spectra for cathodic polarization experiments in pH 5.1 solutions were collected immediately after the pH of the vanadate solutions was adjusted. A Bruker DPX 400 MHz superconducting magnet was used to collect high-resolution 51V (105.2 MHz) NMR spectra. An indirect detection probe was used with a 90° pulse duration of 10.38 μs. Spectra were collected using 8192 transients, a spectral window of 73,529 Hz, a 0.051 s acquisition time, and a 0.20 s relaxation delay. Each spectrum had the subsequent process parameters applied: 10 kHz line broadening, zero filling (25 K points), and baseline correction. A solution consisting of 20% v/v VOCl3 in C6D6 (δ35V = 0 ppm) was used as an external standard to reference the 51V chemical shifts. Peaks were identified by comparison to literature.36,37

Potentiodynamic polarization using the microcapillary electrode.— To characterize the inhibitive effects of tetrahedrally coordinated vanadates, anodic and cathodic polarization curves were collected on representative bulk versions of intermetallics found in Al 2024-T3 in alkaline 10 mM NaVO3 + 0.5 M NaCl and NaVO3-free 0.5 M NaCl solutions. Samples used for potentiodynamic polarization experiments using the microcell were sourced from previous work and commercial suppliers. The 99.999% Al and 99.9% Cu samples were obtained from Alfa Aesar. The intermetallic samples used in this study were prepared and studied previously: Al-Cu2Fe and Al3Cu2Mn.42 Al 4% Cu,43 Al2CuMg, and Al2Cu.44 Samples were ground in 200 proof ethyl alcohol to 1 μm using SiC grinding papers, followed by polishing using 6 and 1 μm diamond pastes. All electrochemical experiments presented in this paper were made using an Autolab PGSTAT 100 potentiostat in conjunction with General Purpose Electrochemical Systems data-acquisition software. Both anodic and cathodic polarization experiments were preceded by 30 s of OCP measurement and were carried out in aerated solutions using a 0.01 V/s scan rate. Anodic polarization curves were initiated at −0.03 V vs OCP and reversed at either 0.0 V vs OCP or manually at approximately 0.05 V above any observed breakdown. Cathodic polarization experiments were initiated at 0.03 V vs OCP and terminated at −2.0 V saturated calomel electrode (SCE), although capillary tip leaking often resulted in
early termination of the experiment. The contact area used for area normalization of data was estimated from digital images taken after individual experiments.

**Electrochemical experiments on bulk Al 2024-T3 sheet.**— Experiments on bulk Al 2024-T3 electrodes in alkaline 0.5 M NaCl solutions with and without 10 mM NaVO₃ were used to obtain electrochemical data from the actual alloy for comparison to microcell results. Experiments in pH 5.1 NaVO₃ solutions were used to show the inhibiting effect that tetrahedrally coordinated vanadates have on cathodic kinetics compared to octahedrally coordinated vanadates. Further, work on bulk Al 2024-T3 was used to gain insight into Al₂CuMg dissolution. An Al 2024-T3 sheet was used for four different sets of experiments, cathodic polarization, OCP measurement, anodic polarization, and potentiostatic experiments. All samples were polished by hand in 200 proof ethyl alcohol to at least 1 μm in a similar fashion as discussed for microcell sample preparation except for the samples for OCP measurements, which were polished to 1/4 μm using diamond paste, the samples for potentiostatic experiments, which were polished to 1 μm using a diamond suspension, and an automatic polisher rather than diamond paste by hand, and the samples used for cathodic polarization, which were polished to 1200 grit under ethyl alcohol. Experiments on a bulk Al 2024-T3 sheet were carried out using a standard three-electrode setup, which included a SCE reference, a platinum counter electrode mesh, and a 1 cm² exposed working electrode. Cathodic polarization experiments in actively aerated pH 5.1 NaVO₃ solutions were preceded by a 30 min measurement of OCP. The scan was initiated at 0.03 V vs OCP, and a scan rate of 0.5 mV/s was used. OCP was measured for 4 h in actively aerated and deaerated 0.5 M NaCl solutions at approximately pH 9.2 with and without 10 mM NaVO₃ to determine the effect of tetrahedrally coordinated vanadates on OCP with time. For deaerated OCP measurements, solutions were deaerated for 1 h before the electrolyte came in contact with the sample. Anodic polarization curves on the AI 2024-T3 sheet in 0.5 M NaCl solution with 10 mM NaVO₃ solution at approximately pH 9.17 were used to make comparisons between microcell data and data collected from the Al 2024-T3 sheet. Anodic polarization experiments were preceded by a 30 min measurement of OCP. The scan was initiated at −0.03 V vs OCP, and a scan rate of 0.5 mV/s was used with scan reversal at −0.2 V vs SCE. Potentiostatic hold experiments were conducted in 0.5 M NaCl solutions between pH 9.1 and 9.23 with and without 10 mM NaVO₃. These experiments were used to determine if tetrahedral vanadates have an effect on the repassivation of the surface once activated and to show the suppression of Mg dissolution from Al₂CuMg intermetallics. Each 100 mL test solution was deaerated for 30 min prior to experimentation and sample exposure. The samples were held at a conditioning potential of 1 V SCE for 1 s and then held at a specific potential for the next 120 s; potential holds at −1.2, −0.9, −0.8, −0.7, −0.6, and −0.5 V SCE were used.

**Results**

**Inhibition from tetrahedral vanadate species vs octahedral species.**—The effect that tetrahedrally coordinated vanadates have on cathodic kinetics compared to octahedrally coordinated species can be observed through experiments in mildly acidic NaVO₃ solutions. Figure 1 shows the NMR spectra from two different pH 5.1 0.5 M NaCl solutions with (a) 0.0025 and (b) 0.25 M NaVO₃. The subscripts of the peak labels in the figure describes the number of vanadium atoms in each oligomer. For example, V₄ indicates single tetrahedrally coordinated vanadium [VO₄]²⁻, V₅ indicates dimeric vanadium [VO₂(OH)²⁻, VO(OH)]⁻, V₆ indicates trimeric vanadium [VO₃(OH)]⁻, V₇ indicates tetrameric (V₆O₁₇), V₈ is pentameric (V₆O₁₇), and V₉ represents decameric vanadate species [V₁₀O₃₂⁻, V₈O₂₃(OH)²⁻, V₆O₂₃(OH)⁴⁻]. V₈ and V₉ are tetrahedrally coordinated species, and V₁₀ is octahedrally coordinated. Also, the vertical scales of the two spectra in Fig. 1 have been adjusted so that peaks in both spectra can be observed. As a result, the two spectra cannot be compared quantitatively. However, the data do show that the 0.0025 M NaVO₃ solution has a greater proportion of tetrahedrally coordinated species relative to octahedral species compared to the 0.25 M NaVO₃ solution, which contains significantly more octahedral species than tetrahedral species.

Figure 2 shows cathodic polarization curves on Al 2024-T3 in aerated pH 5.1 0.5 M NaCl with 0.25 and 0.0025 M NaVO₃ and without NaVO₃. These experiments show the inverse relationship between NaVO₃ concentration and inhibition of cathodic kinetics at pH 5.1, where the dilute solutions containing relatively more tetrahedrally coordinated vanadate to octahedral vanadates have a larger reduction in cathodic kinetics than more-concentrated solutions with relatively more octahedrally coordinated decavanadate.

**Tetrahedral vanadate species in alkaline electrolytes.**— Small changes in solution pH can have a large effect on vanadate speciation. Concerns that the vanadate test solutions would change over the course of microcell experimentation, with consequences for inhibitor behavior, were addressed using NMR. Figure 3 shows the spectra from two samples of pH 9.17 0.5 M NaCl + 10 mM NaVO₃ test solution taken immediately before microcell experimentation and 10 days later after the conclusion of experimentation in vanadate solutions. The solution did remain stable over the course of experimentation and was found to contain a number of different tetrahedral vanadate species as expected from previous work on vanadate inhibition.9,22 The assignments of V₄ and V₅ are not certain, as two different standards available in the literature leave room for speculative interpretation.26,36 However, definitive assignment of these species is not critical for this work. The solution used for this work contained tetrahedrally coordinated species and predominately V₇; no octahedrally coordinated vanadates were detected.

**Polarization of intermetallics in tetrahedral vanadate solutions.**— Figures 4a-g show sample anodic polarization curves for pure Al, pure Cu, Al₄Cu, Al₃CuMg, Al₃CuFe, and Al₂CuMgN, respectively. The objective of these experiments was to determine the effect of tetrahedral vanadates on the anodic behavio.
ior of different constituent intermetallic particles in Al 2024-T3. Detailed anodic polarization results are discussed below in conjunction with cathodic polarization results for each intermetallic.

Figures 5a-g are a collection of cumulative distribution plots showing $E_{\text{cor}}$, $E_{\text{pit}}$, and $E_{\text{rp}}$ for various intermetallic compounds. $E_{\text{cor}}$ is the corrosion potential, $E_{\text{pit}}$ is the pitting potential, and $E_{\text{rp}}$ is the reversible potential on the reverse scan, which was defined as the potential at the smallest observed current on the reverse scan. Cumulative probability plots are of value because there can be significant variation in the measured characteristic potentials of intermetallic compounds. Knowing this variation is important in the interpretation of corrosion processes. The statistical variation, which would be lost by simply taking averages, can be shown fully in cumulative distribution plots. The corrosion rate was estimated from both anodic and cathodic polarization curves (to be presented later) by extrapolation of linear passive regions and regions of oxygen evolution, respectively, to the intersection with corrosion potential.

Figures 6a-g show sample cathodic polarization curves for pure Al, pure Cu, Al 4% Cu, Al 2Cu, Al 2CuMg, Al 7Cu2Fe, and Al20Cu2Mn3, respectively. These experiments were used to determine the effect of vanadates on cathodic kinetics of intermetallic particles in Al 2024-T3.

Figures 7a-g are a collection of cumulative distribution plots showing corrosion current density ($i_{\text{corr}}$), passivation current density ($i_{\text{pass}}$), and current density at $-1.3 \, V_{\text{SCE}}$ for tested intermetallics and metals. Passivation current density was defined as the current density immediately before breakdown. The current density at $-1.3 \, V_{\text{SCE}}$ was used as a comparative measure of cathodic reduction kinetics, because the potential at this point was below the most active $E_{\text{cor}}$ and this point of the curve allowed a direct comparison of reduction kinetics at potentials where oxygen reduction reactions likely contributed to the cathodic response. Considerable hydrogen evolution reactions may also be present at $-1.3 \, V_{\text{SCE}}$. The following presents results for each tested material.

Pure Al.— The addition of 10 mM NaVO3 to alkaline 0.5 M NaCl results in a shift of corrosion potential to more active potentials relative to NaVO3-free solutions on pure Al. However, NaVO3 also appears to slightly increase the corrosion current density. A small decrease in cathodic kinetics, as seen by a decrease in current density at $-1.3 \, V_{\text{SCE}}$ in vanadate solutions, is largely offset by an increase in anodic kinetics, as seen by an increase in $i_{\text{pass}}$ observed in NaVO3 solutions. NaVO3 increases the breakdown potential of pure Al and does not appear to have an effect on the repassivation potential on the reverse scan.

Pure Cu.— NaVO3 was observed to have little effect on the corrosion potential, corrosion current density, or repassivation potential on the reverse scan of pure Cu. Pure Cu did not demonstrate passive behavior with an observable characteristic breakdown. However, NaVO3 was observed to have an effect on dissolution kinetics. Comparison of current density at $-1.3 \, V_{\text{SCE}}$ shows that NaVO3 solutions slowed cathodic kinetics.

Al 4% Cu.— Al 4% Cu was used as an analog material for the matrix of Al 2024-T3. NaVO3 was observed to shift the corrosion potential to more active potentials, and the corrosion current density was observed to decrease as a result of a decrease in cathodic kinetics. A relatively small increase in $E_{\text{pit}}$ was observed in NaVO3 solutions compared to increases observed on other intermetallic compounds. Vanadate increased the current density observed in the passive region of the anodic polarization scans.

**Figure 2.** (Color online) Cathodic polarization curves on Al 2024-T3 in aerated pH 5.1 0.5 M NaCl with 0.25 and 0.0025 M NaVO3 and without NaVO3. These experiments show an inverse relationship between NaVO3 concentration and inhibition of cathodic kinetics at pH 5.1, which correlates well with a transition from solutions dominated by octahedrally coordinated vanadates to tetrahedral vanadates.

**Figure 3.** NMR spectra showing the presence of tetrahedrally coordinated vanadates ($V_1$, $V_2$, $V_3$, and $V_4$) in pH 9.17 0.5 M NaCl + 10 mM NaVO3 solution used for microcapillary electrochemical experiments (a) immediately prior to experimentation and (b) 10 days later after completion of experiments.
Al\textsubscript{2}Cu, Al\textsubscript{7}Cu\textsubscript{2}Fe, and Al\textsubscript{20}Cu\textsubscript{2}Mn.— These three intermetallics were observed to behave similarly in NaVO\textsubscript{3} solution. All three intermetallics showed a large shift in corrosion potential to more active potentials in NaVO\textsubscript{3} solutions. Al\textsubscript{7}Cu\textsubscript{2}Fe and Al\textsubscript{2}Cu showed a decrease in corrosion current density in NaVO\textsubscript{3} solution, while vanadate had little effect on the corrosion current density of Al\textsubscript{20}Cu\textsubscript{2}Mn\textsubscript{3}. These intermetallics showed an order of magnitude or more decrease in current density at −1.3 V \textsubscript{SCE}, indicating an overall decrease in cathodic kinetics. Further, the breakdown potential of all three intermetallics was observed to increase in NaVO\textsubscript{3}; however, NaVO\textsubscript{3} had little effect on the current magnitude observed in the passive region of the anodic polarization curves.

Al\textsubscript{2}CuMg.— A large degree of variability was observed in the anodic polarization curves of Al\textsubscript{2}CuMg. This variability was most pronounced in the breakdown potential distribution. Breakdown in Al\textsubscript{2}CuMg is believed to be associated with loss in protection conferred by a Cu-enriched layer that forms on the intermetallic surface as the intermetallic dissolves Mg and, to a lesser extent, Al. Two different reproducible behaviors were observed on Al\textsubscript{2}CuMg in 0.5 M NaCl solutions, which are shown in Fig. 4e. Most anodic polarization curves in NaCl-only solutions showed passive behavior up to and above −0.4 V \textsubscript{SCE}. In some cases, the rapid scan rate resulted in polarization of sample intermetallics well beyond potentials necessary for breakdown, as illustrated by one of the NaCl-only curves in Fig. 4e. In these cases, the potential of the plateau after breakdown was used for \(E_{\text{pit}}\) values shown in cumulative probability plots. A small number of polarization curves in NaCl-only solution were observed to have more noble corrosion potentials and more

![Figure 4.](https://example.com/figure4.png)

Figure 4. (Color online) Sample anodic polarization curves obtained using the microcapillary electrode in pH 9.17 0.5 M NaCl solution with and without 10 mM NaVO\textsubscript{3} for (a) pure Al, (b) pure Cu, (c) Al 4% Cu, (d) Al\textsubscript{2}Cu, (e) Al\textsubscript{2}CuMg, (f) Al\textsubscript{7}Cu\textsubscript{2}Fe, and (g) Al\textsubscript{20}Cu\textsubscript{2}Mn\textsubscript{3}.
active breakdowns in relation to the previously discussed $\text{Al}_2\text{CuMg}$ anodic polarization curves. Generally, the addition of $\text{NaVO}_3$ resulted in much more reproducible behavior with less data scatter. For example, $\text{Al}_2\text{CuMg}$ in $\text{NaCl}$ solutions had breakdown potentials scattered from approximately $-0.8$ to $0.2 \text{V}_{\text{SCE}}$. $\text{NaVO}_3$ caused the data distribution to become much tighter, ranging from approximately $-0.7$ to $-0.6 \text{V}_{\text{SCE}}$, which on average is a shift in breakdown potential to more active potentials. Interestingly, the breakdown potential distribution is shifted to lower potentials when vanadate is present in the solution. This behavior is different than that observed for other intermetallic compounds, where breakdown potentials are ennobled by vanadate additions. Additionally, on average, $\text{NaVO}_3$ caused a small decrease in the corrosion potential and corrosion current density; however, larger passive current densities were observed in $\text{NaVO}_3$ solutions. $\text{NaVO}_3$ was observed to decrease the current density at $-1.3 \text{V}_{\text{SCE}}$, an indication of suppressed cathodic kinetics.

In summary, all tested materials showed passive behavior and a breakdown, with the exception of pure Cu. $\text{NaVO}_3$ in approximately pH 9 test solutions shifted the corrosion potential to more active potentials, increased the pitting potential, except for $\text{Al}_2\text{CuMg}$, and decreased the cathodic kinetics. A summary of averaged characteristic potentials and current densities from polarization experiments is found in Table I.

**Figure 5.** (Color online) Cumulative probability plots displaying $E_{\text{corr}}$, $E_{\text{pit}}$, and $E_{\text{rp}}$ of tested materials in pH 9.17 0.5 M NaCl solution with and without 10 mM $\text{NaVO}_3$ for (a) pure Al, (b) pure Cu, (c) Al 4% Cu, (d) Al$_2$Cu, (e) Al$_2$CuMg, (f) Al$_2$Cu$_2$Fe, and (g) Al$_2$yCu$_z$Mn$_3$. The “V” in the legend indicates curves collected in solutions containing $\text{NaVO}_3$. 

In summary, all tested materials showed passive behavior and a breakdown, with the exception of pure Cu. $\text{NaVO}_3$ in approximately pH 9 test solutions shifted the corrosion potential to more active potentials, increased the pitting potential, except for $\text{Al}_2\text{CuMg}$, and decreased the cathodic kinetics. A summary of averaged characteristic potentials and current densities from polarization experiments is found in Table I.

**OCP and scanning electron microscopy images of Al 2024-T3 exposed to tetrahedral vanadate solutions.**—Figure 8 is a plot of the OCP of Al 2024-T3 in aerated and deaerated 0.5 M NaCl solutions with and without 10 mM $\text{NaVO}_3$. The solutions were not intentionally buffered in any way, and as a result the solution pH at initial sample exposure and during OCP measurement varied. Typically, the pH of actively aerated solutions would become more acidic during the measurement. For example, the pH was observed to drop from 9.17 to 6.72 and from 9.19 to 8.85 from the beginning
of OCP measurement until completion 4 h later for aerated solutions without NaVO₃ and with NaVO₃, respectively. In contrast, deaerated solutions would become more alkaline during the hour of deaeration prior to experimentation, and then pH would change little during the 4 h of OCP measurement. Similar to aerated solutions without NaVO₃, deaerated solutions typically were observed to have larger changes in pH compared to solutions with NaVO₃. Vanadates can act as buffers, and this may be important with regard to the mechanism of vanadate solution inhibition. In aerated solutions without NaVO₃, the OCP was observed to initially trend toward more noble potentials while cyclically fluctuating between approximately −0.89 and −0.64 V SCE for the first 20–25 min of measurement before stabilizing to approximately −0.62 V SCE. Although pH was not measured during the experiment, it is suspected the pH was becoming more acidic with time as dissolved CO₂ dissociated to and equilibrated with CO₃²⁻ and HCO₃⁻. In aerated solutions, the addition of NaVO₃ maintained the OCP at values below −0.8 V SCE after 4 h, and the pH remained alkaline. In contrast, in deaerated solutions the addition of NaVO₃ was observed to increase the OCP from approximately −1.100 to −0.85 V SCE after 4 h.

Figures 9a-d show scanning electron microscopy (SEM)/H20849 images of Al2024-T3 in aerated and deaerated 0.5 M NaCl solutions with and without 10 mM NaVO₃ corresponding to the samples used in Fig. 8 after 4 h of OCP measurement. A comparison of samples in aerated solutions with and without vanadate show that in aerated solutions the presence of vanadate greatly decreased attack on and around intermetallic particles, Fig.9a and b. The comparison is not completely direct, as the pH of the NaVO₃-free solution drifted over experimentation to below pH 7. Regardless, the effect of vanadate even if only acting as a buffer is apparent. In particular, in aerated solutions without vanadate, nearly every intermetallic demonstrates attack, and a large number of intermetallics appear to have undergone complete dissolution or undercutting. In addition, a number of small, bright particles, possibly redistributed Cu, are observed on the

**Figure 6.** (Color online) Sample cathodic polarization curves obtained using the microcapillary electrode in pH 9.17 0.5 M NaCl solution with and without 10 mM NaVO₃ for (a) pure Al, (b) pure Cu, (c) Al 4% Cu, (d) Al₄Cu, (e) Al₄CuMg, (f) Al₃Cu₂Fe, and (g) Al₇Cu₂Mn₃.
Table I. Averaged electrochemical data for intermetallics tested in approximately pH 9.17 0.5 M NaCl solutions with and without 10 mM NaVO$_3$. A “NaCl” heading indicates data from NaCl-only solutions, while a “NaCl + V” heading indicates data collected in NaCl solution that contained 10 mM NaVO$_3$.

<table>
<thead>
<tr>
<th>Material</th>
<th>$E_{corr}$ (V SCE)</th>
<th>$E_{pul}$ (V SCE)</th>
<th>$E_{ip}$ (V SCE)</th>
<th>$i_{corr}$ (Amps/cm$^2$)</th>
<th>$i_{pul}$ (Amps/cm$^2$)</th>
<th>$i$ at $-1.3$ V SCE (Amps/cm$^2$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cu</td>
<td>$-0.24$</td>
<td>$-0.24$</td>
<td>$-0.11$</td>
<td>$-0.10$</td>
<td>$1.7 \times 10^{-5}$</td>
<td>$1.9 \times 10^{-5}$</td>
</tr>
<tr>
<td>Al</td>
<td>$-1.01$</td>
<td>$-1.18$</td>
<td>$-0.70$</td>
<td>$0.92$</td>
<td>$-0.91$</td>
<td>$-0.99$</td>
</tr>
<tr>
<td>Al$_8$Cu$_2$Fe</td>
<td>$-0.63$</td>
<td>$-0.93$</td>
<td>$-0.74$</td>
<td>$-0.75$</td>
<td>$5.6 \times 10^{-6}$</td>
<td>$5.7 \times 10^{-5}$</td>
</tr>
<tr>
<td>Al$_8$Cu$_2$</td>
<td>$-0.59$</td>
<td>$-0.91$</td>
<td>$-0.70$</td>
<td>$-0.89$</td>
<td>$5.6 \times 10^{-6}$</td>
<td>$5.7 \times 10^{-5}$</td>
</tr>
<tr>
<td>Al$_8$Cu$_2$Mn$_3$</td>
<td>$-0.68$</td>
<td>$-0.98$</td>
<td>$-0.66$</td>
<td>$-0.69$</td>
<td>$1.2 \times 10^{-5}$</td>
<td>$1.3 \times 10^{-4}$</td>
</tr>
<tr>
<td>Al$_2$CuMg</td>
<td>$-0.92$</td>
<td>$-1.06$</td>
<td>$-0.29$</td>
<td>$-0.60$</td>
<td>$2.2 \times 10^{-5}$</td>
<td>$2.8 \times 10^{-6}$</td>
</tr>
</tbody>
</table>

Figure 7. (Color online) Cumulative probability plots of $i_{corr}$, $i_{pul}$, and $i$ at $-1.3$ V SCE of tested materials in pH 9.17 0.5 M NaCl solution with and without 10 mM NaVO$_3$. A “NaCl” heading indicates data from NaCl-only solutions, while a “NaCl + V” heading indicates data collected in NaCl solution that contained 10 mM NaVO$_3$. The “V” in the legend indicates curves collected in solutions containing NaVO$_3$. 
2024-T3 samples were activated for 1 s at 1 V SCE and then held at more noble potentials at or more positive than −0.8 V SCE exhibited dissolution transients. However, for each curve obtained in NaVO₃ solution, the current response had a lower magnitude than the current response at corresponding potentials obtained in NaCl-only solutions. A comparison of total charge passed in Fig. 11c shows that at all potentials except −0.6 V SCE, vanadate reduced the total charge passed.

**Discussion**

Cathodic inhibition from tetrahedrally coordinated vanadate species.—Figure 2 shows that an inverse relationship exists between NaVO₃ concentration and inhibition of cathodic kinetics at pH 5.1, where dilute solutions perform better than more concentrated solutions. The pH and concentration of these experiments was chosen purposely to be near the transition from solutions dominated by octahedral species (0.25 M NaVO₃) to those dominated by tetrahedral species (0.0025 M NaVO₃). The dilute NaVO₃ solution contains relatively more tetrahedral vanadate as compared to the concentrated NaVO₃ solution, which contains relatively little tetrahedral species compared to octahedral species. At this pH, concentrated NaVO₃ solutions are expected to predominately have decavanadate species, which are octahedrally coordinated. As the concentration of NaVO₃ in solution decreases, tetrahedral species become more prevalent. The increased presence of tetrahedral species correlates with the reduced cathodic kinetics observed in cathodic polarization experiments. This is in agreement with other work, which has suggested that tetrahedrally coordinated vanadates inhibit the corrosion of Al 2024-T3. 18,20-22,27 However, the more concentrated NaVO₃ solution containing predominately octahedral decavanadate appears to be a modest cathodic inhibitor at this pH, in contrast to the findings of previous work, which suggest that decavanadates are reduced and possibly increase corrosion. 18,22 If decavanadate is not responsible for the observed inhibition, then the small concentrations of tetrahedral species in solution are likely responsible.

Suppression of Al₂CuMg breakdown.—In alkaline NaCl solutions, a portion of the total Al₂CuMg intermetallic particle population would be expected to break down at potentials as low as −0.8 V SCE. At the OCP in aerated NaCl solutions, which fluctuates approximately from −0.9 and −0.75 V SCE before the solution begins to become acidic from dissolved atmospheric CO₂ with eventual OCP stabilization near −0.65 V SCE, Al₂CuMg breakdown leads to selective Mg dissolution and formation of Cu-enriched local cathodes. In contrast, Al₂CuMg in alkaline-aerated NaCl with 10 mM NaVO₃ is expected to break down at potentials only above −0.7 V SCE. However, the OCP in NaVO₃ solutions after 4 h was observed to remain below approximately −0.79 V SCE. The shift in alloy OCP to more active potentials is a result of a decrease in cathodic kinetics observed on the matrix and all intermetallic phases, as seen in Fig. 6. Iannuzzi et al. argue that decreased kinetics might be the result of the adsorption of monovanadate to the surface, blocking reactive sites on intermetallic particles and displacing Cl⁻ from the matrix surface. 23 The consequence of decreased cathodic kinetics and the resultant shift in OCP combined with a small increase in Al₂CuMg breakdown potential observed in NaVO₃ solutions, is that Al₂CuMg intermetallics remain “inefficient” cathodes. If Cu-enriched particles resulting from Al₂CuMg dissolution are assumed to act like pure Cu (Fig. 6b), then the rate at which cathodic reactions are supported is greater than compared to rates supported by intact Al₂CuMg in NaVO₃ solutions (Fig. 6c).

Variation in anodic behavior of Al₂CuMg.—As seen in Fig. 4e, two different anodic polarization behaviors were observed in NaCl solutions. In a minority of cases, Eₗₗₗ was observed to be −0.8 V SCE or more noble with breakdown at potentials near −0.6 V SCE or more...
active. Yoon and Buchheit reported that Al$_2$CuMg has an initial OCP in 0.5 M NaCl near −1.2 V SCE, which drifts and becomes stable between −0.86 and −0.76 V SCE after approximately 1000 s. It was claimed that Al$_2$CuMg experienced transient dissolution followed by passivation and stable low-rate dissolution with an OCP near −0.8 V SCE. It seems possible that anodic polarization of Al$_2$CuMg that showed corrosion potentials at approximately −0.8 V SCE had already experienced transient dissolution compared to experiments that showed more active corrosion potentials near the −1.2 V SCE OCP reported by Yoon. Generally, the wide variability in $E_{\text{corr}}$ and $E_{\text{pit}}$ data obtained on Al$_2$CuMg in NaCl could be attributed to samples being at various stages of dissolution prior to polarization, perhaps a result of extreme sensitivity to variation in polishing conditions. In contrast, the behavior of Al$_2$CuMg in NaVO$_3$ solutions showed little variability in electrochemical response, which is possibly evidence that vanadates control the process of dealloying and redistribution of Cu on the surface, resulting in overall suppression of Al$_2$CuMg dissolution. The breakdown potential distribution measured in vanadate solutions is shifted to more active potentials compared to vanadate-free solutions. This is consistent with the idea that vanadates suppress Mg dissolution from the phase. In this case, suppression of Mg dissolution would result in a thinner, less-protective Cu-enriched layer on the surface of the phase that breaks down at lower potentials. Even though breakdown of Al$_2$CuMg is shifted to lower potentials when vanadate is present in solution, the corrosion potential of the alloy remains below the Al$_2$CuMg breakdown potential distribution, and corrosion susceptibility is decreased.

**Vanadate buffering and circumferential attack.**—In this work and in previous work, it has been observed that vanadate solutions act as buffers, requiring more alkali or acid to adjust the pH than...
necessary for equivalent vanadate-free solutions. Particularly, for this work, once vanadate solutions were adjusted to pH values near 9, the solution pH remained stable for months. In contrast, equivalent NaCl solutions adjusted to pH values near 9 would quickly become more acidic as carbonic acid formed from dissolved atmospheric CO2. Leclère et al. have argued that the presence of a buffer or weak acid can suppress alkaline dissolution near cathodic sites resulting from hydroxyl produced by oxygen reduction, which results in large cathodic currents that drive pitting and localized attack more than observed in unbuffered solutions. The production of reducible $H^+$ at sites of local attack may contribute to the increased cathodic currents. A rough estimate of buffering capacity, $\beta$, for both carbonate and vanadate at pH 9.17 is easily calculated using the following simplified equation

$$\beta = 2.303 \frac{K_a C_t [H^+]}{(K_a + [H^+])^2}$$

where $K_a$ is the acid dissociation constant and $C_t$ is the total concentration of acid and base or total buffer concentration. At pH 9.2, the dominant species from atmospheric CO$_2$ in solution are HCO$_3^-$ and CO$_3^{2-}$, which have a $pK_a$ of 10.33 and a total concentration of approximately 0.025 M, which gives a buffering capacity of 0.0035 M/pH unit. If pH 9.17 10 mM NaVO$_3$ is assumed to be dominated by monomeric vanadate, a $pK_a$ of 8.2 is appropriate (other tetrahedral vanadates have $pK_a$ values ranging from 7.92 to 9.1) and gives a buffering capacity of 0.002 M/pH unit. At this pH, the contribution of vanadate as a buffer is small compared to carbonate, while both seem to be relatively ineffective buffers. Figure 9a shows an Al 2024-T3 sample exposed to 0.5 M NaCl without vanadate, which is analogous to a nearly unbuffered solution. The observation of circumferential attack and white corrosion product around intermetallics is consistent with initial alkaline attack from increased pH near cathodic sites, supporting oxygen reduction followed by aggressive pitting and crevice corrosion around intermetallics. In vanadate-free unbuffered deaerated NaCl solutions (Fig. 9c), despite the lack of oxygen, attack and trenches around intermetallic particles are still observed. In contrast, the degree of attack in alkaline deaerated (Fig. 9d), and in particular aerated (Fig. 9b), NaCl solutions that contain NaVO$_3$ is significantly less than observed on corresponding NaVO$_3$-free samples. There is some attack near in-

Figure 10. Duplicate anodic polarization curves of a bulk Al 2024-T3 sheet in aerated 0.5 M NaCl + 10 mM NaVO$_3$ solution at pH 9.17.

![Figure 10](image1.png)

Figure 11. (Color online) Current response of potentiostatic hold experiments on Al 2024-T3 in pH 9.17 0.5 M NaCl (a) with 10 mM NaVO$_3$ and (b) without NaVO$_3$. (c) Summary of total charge passed as a function of potential hold for a and b.
termetallic particles, but complete circumferential attack of intermetallics is not observed. If vanadate were assumed to have an effect as a buffer, this would be consistent with previous observations of attack on Al–Cu alloys in buffered and unbuffered solution, where more trenching is observed around intermetallics in unbuffered solutions as a result of alkalinization, while buffers decrease trenching by neutralizing local cathodic alkalinization. However, it has also been previously observed that total cathodic current is greater in buffered solutions because current is not used up for alkalinization and cathodic attack. This is in contrast to current microcell results, which show decreases in cathodic kinetics in vanadate solutions and Fig. 11c. As a result, the action of vanadate as a cathodic inhibitor seems to dominate any effect from buffering. It may be possible that vanadate acting as a buffer does prevent trenching as observed in Fig. 9b, but there is not a corresponding increase in cathodic reactions in alkaline NaCl solutions.

Conclusions

1. An inverse relationship exists between reduction in cathodic kinetics and NaVO₃ concentration in mildly acidic solutions, where dilute solutions with tetrahedrally coordinated vanadates inhibit cathodic kinetics better than concentrated solutions with octahedrally coordinated vanadates.

2. Tetrahedrally coordinated vanadates inhibit the ability of intermetallic particles and the matrix of Al 2024-T3 to support cathodic reactions in alkaline NaCl solutions.

3. Tetrahedrally coordinated vanadates decrease corrosion potential and generally increase pitting potential.

4. The overall reduction of cathodic kinetics on intermetallics and the matrix of Al 2024-T3 in alkaline vanadate solutions shifts the OCP below the potential required to cause Al₃CuMg breakdown. This may prevent selective Mg dissolution from Al₃CuMg and subsequent formation of Cu-enriched clusters capable of supporting rapid cathodic reaction kinetics.

5. Tetrahedral vanadates decrease the incidence of circumferential attack around intermetallic particles.

6. The inhibition of cathodic reactions provided by tetrahedral vanadates dominates any effect from buffering.

Acknowledgments

Support for portions of this work by Concurrent Technologies Corp., Largo, FL is gratefully acknowledged. The authors also thank Dr. N. Birbilis for intermetallic collection and characterization, Dr. B. N. Padgett for assistance with microelectrochemical cell measurement, Dr. T. L. Young for assistance with NMR measurements, and Dr. G. S. Frankel for discussions on experimental approach.

Ohio State University assisted in meeting the publication costs of this article.

References