Journal of The Electrochemical Society, **155** (3) C117-C126 (2008) 0013-4651/2008/155(3)/C117/10/\$23.00 © The Electrochemical Society

C117



Investigation and Discussion of Characteristics for Intermetallic Phases Common to Aluminum Alloys as a Function of Solution pH

N. Birbilis*,a,z and R. G. Buchheit*

Fontana Corrosion Center, Department of Materials Science and Engineering, The Ohio State University, Columbus. Ohio 43210. USA

This paper presents results for corrosion potentials, pitting potentials, and electrochemical characteristics for intermetallic particles commonly present in high strength aluminum-based alloys, for tests conducted in a 0.1 M NaCl solution of varying pH via the use of a microcapillary electrochemical cell. The intermetallics investigated were Mg_2Si , $MgZn_2$, Al_7Cu_2Fe , Al_2Cu , Al_2CuMg , and Al_3Fe . Elaboration of the results reveals that the electrochemical behavior of such compounds varies markedly with pH, with attendant ramifications for localized corrosion and protection in Al alloys. Examples of this are shown for AA7075-T651, where it is shown that the localized corrosion morphology is drastically different upon the bulk alloy depending on the pH of the test environment. A stochastic pitting is observed at an acid pH, near-neutral conditions result in a deterministic-type pitting, and a general corrosion is observed at an alkaline pH.

© 2008 The Electrochemical Society. [DOI: 10.1149/1.2829897] All rights reserved.

Manuscript submitted October 2, 2007; revised manuscript received November 16, 2007. Available electronically January 22, 2008.

In a recent review, we presented the electrochemical characteristics for a number of intermetallic particles and phases common to a large range of engineering aluminum (Al) alloys. In that case, the results were presented in a survey format as a function of chloride concentration in near-neutral solution.

In the case of "real" corrosion of Al alloys, not only will the environments to which they are exposed vary in pH (viz., acid rain, concentrated solution in aircraft lap joints, etc.), but gradients will also exist, often sharp, of pH across the alloy surface. ^{2,3} As ought to be expected, the electrochemical response and activity of most metals and alloys will vary rather markedly with the solution pH. ⁴ This will have attendant ramifications in the evolution and rate, and possibly the morphology, of the corrosion that develops on bulk Al alloys that contain phases and intermetallics of different composition (and hence electrochemical properties).

In the case of Al alloys in near-neutral bulk solutions, it is well established that pitting is influenced by intermetallic particles that exhibit different surface film characteristics to the matrix, and may be either anodic or cathodic relative to the matrix. ^{1-3,5-9} In such cases, two main types of pit morphologies are observed. So-called "circumferential" pits appear as a ring of attack around a more or less intact particle or particle colony, with the attack appearing to be mainly in the matrix phase (often termed "trenching" ^{2,3,10}). The second morphology is the apparent selective dissolution of an intermetallic particle. Pits of this type can be deep and may contain remnants of the particle in them. ³ In contrast, comparatively little is known about bulk alloy corrosion morphology across the pH range.

The microstructures developed in high strength aluminum alloys are complex and incorporate a combination of equilibrium and non-equilibrium phases. In this study, we focused on the electrochemical properties of the synthesized intermetallics MgZn₂, Al₇Cu₂Fe, Al₂CuMg, Al₂Cu, Mg₂Si, and Al₃Fe, which have been quantitatively noted as being prominent in 7xxx series alloys. ¹¹⁻¹⁴ This facilitates a focused discussion of the results in a practical context; however, it is emphasized that a number of the above-mentioned intermetallics can be found in several other common cast or wrought alloys (e.g., 2024, 6061). ^{2.5,10,11,15} The aim of this work is to contribute to a detailed characterization of the electrochemical behavior of the intermetallic compounds in Al alloys, presenting previously unreported information, while attempting to provide a more quantitative description of mechanistic aspects of corrosion phenomena in

* Electrochemical Society Active Member.

commercial Al alloys by determining reaction kinetics phase-byphase and upon well-defined particle types. It may be considered supplementary or additional to the work given in Ref. 1, while it can also be interpreted in the context of development of predictive models.

Experimental

Materials and materials characterization.— Electrochemical testing of intermetallic compounds synthesized in bulk form is now relatively common in the study of localized corrosion phenomena (Ref. 1 and references therein). In this case intermetallic particles were synthesized by combining the necessary proportions of constituent pure metals (supplied by Alfa-Aesar), the methodology and approach of which were covered in detail in Ref. 1. Fabrication was carried out via either induction or arc melting. Subsequent identification of intermetallics was done via scanning electron microscopy (SEM) (Philips XL-30 FEG-ESEM) enabled with quantitative energy dispersive X-ray spectroscopy (EDXS) and the facility to generate backscattered electron Kikuchi patterns. In the present study, the microscopy was also carried out using the Philips XL-30.

Electrochemical characterization.— Electrochemical testing was carried out using a microcell method, as outlined previously in Ref. 1 and 16-18. In this method, the working electrode area is defined by the area of metal that comes into contact with the opening of a microcapillary filled with an electrolyte, while containing a small-wire counter electrode and an electrolytic contact with a saturated calomel reference electrode. There are several unique aspects of microcell testing (viz., the effect of capillary diameter, solution resistance (iR) effects, and current limitations) that we have covered in detail elsewhere ¹⁹ and need to be considered in the execution of such testing. In cases where the bulk alloy was tested (AA7075-T651 supplied by Alcoa), the electrode area was 1 cm² and the testing was carried out using a conventional "flat cell" (PAR).

Potentiodynamic polarization was carried out at a scan rate of $0.01~V~s^{-1}$ using an Autolab PGSTAT 100. The tests reported herein should also be considered to have been carried out in aerated conditions, because no special effort was made to exclude oxygen from the capillary or solution. Each test was repeated at least six times and the average values are reported herein. Measurements were performed in 0.1 M NaCl at pH = 2.5, 6, 10, and 12.5. Temperature is considered to be constant at $20^{\circ}C$.

Results and Discussion

Potentiodynamic polarization response of intermetallic particles as a function of pH.— Figures 1-6 summarize the potentiodynamic polarization response of Al₇Cu₂Fe, Al₂Cu, Al₃Fe, MgZn₂, Mg₂Si,

^a Present address: Department of Materials Engineering, Monash University, Vic. 3800, Australia.

^z E-mail: birbilis.1@osu.edu

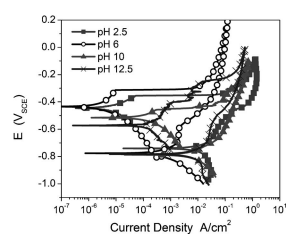


Figure 1. Potentiodynamic polarization diagrams for Al_7Cu_2Fe collected in 0.1 M NaCl at pH 2.5, 6, 10, and 12.5.

and Al₂CuMg as a function of pH. The reported values for both $E_{\rm corr}$ and $E_{\rm pit}$ nominally varied within a window; however, a representative polarization curve at each pH value is seen (in Fig. 1-6) from a group of at least six replicates. The distribution in the measured $E_{\rm corr}$ and $E_{\rm pit}$ is reported in Fig. 7-12. A more detailed discussion of these results is given in the following section.

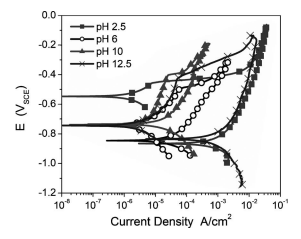


Figure 2. Potentiodynamic polarization diagrams for Al_2Cu collected in 0.1 M NaCl at pH 2.5, 6, 10, and 12.5.

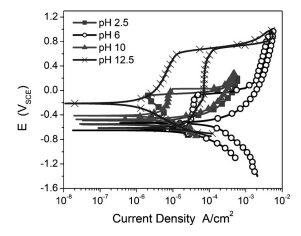


Figure 3. Potentiodynamic polarization diagrams for Al_3Fe collected in 0.1 M NaCl at pH 2.5, 6, 10, and 12.5.

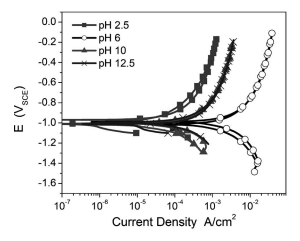


Figure 4. Potentiodynamic polarization diagrams for $MgZn_2$ collected in 0.1 M NaCl at pH 2.5, 6, 10, and 12.5.

 Al_7Cu_2Fe .— Figure 1 indicates that Al_7Cu_2Fe reveals a characteristic value for $E_{\rm pit}$ over the range of pH values investigated. This indicates that over the range of pH 2.5–6 this compound is "spontaneously passive" and does not freely corrode in a 0.1 M NaCl solution. The term spontaneously passive refers to a condition whereby (anodic) polarization above $E_{\rm corr}$ reveals a typical passive region. We observe that Al_7Cu_2Fe displays a clear "breakdown"

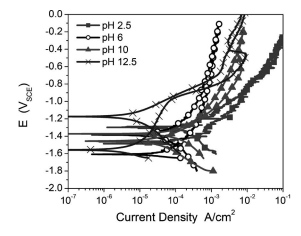


Figure 5. Potentiodynamic polarization diagrams for Mg_2Si collected in 0.1 M NaCl at pH 2.5, 6, 10, and 12.5.

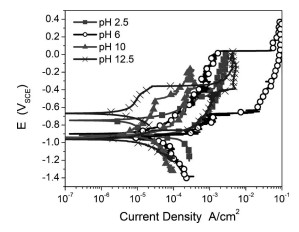


Figure 6. Potentiodynamic polarization diagrams for Al_2CuMg collected in 0.1 M NaCl at pH 2.5, 6, 10, and 12.5.

80

80

100

100

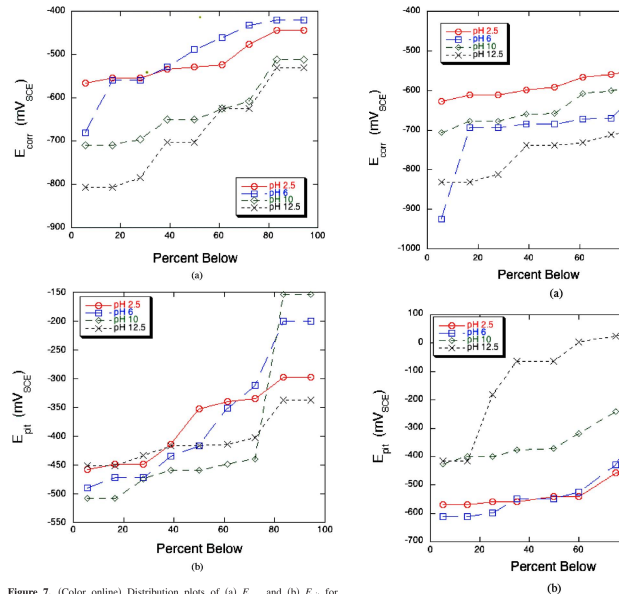


Figure 7. (Color online) Distribution plots of (a) $E_{\rm corr}$ and (b) $E_{\rm pit}$ for Al₇Cu₂Fe collected in 0.1 M NaCl at pH 2.5, 6, 10, and 12.5.

Figure 8. (Color online) Distribution plots of (a) $E_{\rm corr}$ and (b) $E_{\rm pit}$ for Al $_2$ Cu collected in 0.1 M NaCl at pH 2.5, 6, 10, and 12.5.

process and hence a distinct $E_{\rm pit}$ value also. In the case of pH 10 and 12.5, however, we note that prior to breakdown, anodic current values are nominally high in the vicinity of 100 μ A/cm². This is also concomitant with the nominally high self-dissolution rate (i.e., the i at $E_{\rm corr}$). This phenomenon and its likely origin is explained somewhat further in the ensuing discussion, while cathodic polarization data are also given. An apparent trend in the results obtained for this compound is evident, whereby overall the value for $E_{\rm corr}$ nominally decreases with increasing the pH; however, the $E_{\rm pit}$ values slightly increase with the pH as seen in Fig. 7. We also note from Fig. 1 that the repassivation potential for Al₇Cu₂Fe is pH independent and comparatively low at \sim -800 mV_{SCE}.

 Al_2Cu .— Figure 2 shows the electrochemical response of Al_2Cu , revealing again that this compound is spontaneously passive and does not freely corrode in NaCl solution over the range of pH tested (2.5–12). At the lowest pH (most acid), the $E_{\rm corr}$ value is some 200 mV more positive than the values observed at higher pH (alkaline), which is noteworthy. The results for testing at a pH 6, 10, and 12.5 are similar in their nature in that Al_2Cu reveals a well-defined window of passivity for potentials broadly between about -700 and -400 mV_{SCE}. Figure 8b reveals a marked increase in the $E_{\rm pit}$ of

 Al_2Cu in an alkaline pH. Again, we can note from Fig. 2 that the repassivation potential of Al_2Cu is pH independent and confined to a value in the vicinity of $\sim\!-900\,$ mV $_{SCE}.$

 Al_3Fe .— Figure 3 also shows that Al_3Fe displays well-defined regions of passivity prior to breakdown, with an attendant low self-dissolution rate. With the exception of testing at the highest pH, the response of the compound is nominally similar, with an average passive window (viz., $E_{\rm corr} - E_{\rm pit}$) of about 400 mV. However, the passive current density at a pH 6 is nominally about a factor of 5 greater than that at a pH 2.5 or 10 (Fig. 3). Upon increasing the pH we concomitantly observe a disproportionate widening of the passive range to values in the vicinity of half a volt (apparent in Fig. 9b), indicating an apparent thermodynamic stability in the environment tested and at an elevated pH. This strong pH effect is probably due to the insolubility of Fe oxides under increasingly alkaline oxidizing conditions. We note that the behavior of Al_3Fe varied markedly from that of Al_7Cu_2Fe , which also contains Cu.

 $MgZn_2$.— In the case of $MgZn_2$, as seen in Fig. 4, we observe a general response that does not reveal any passivity; hence, corrosion occurs freely above the respective $E_{\rm corr}$. In such cases, the corrosion

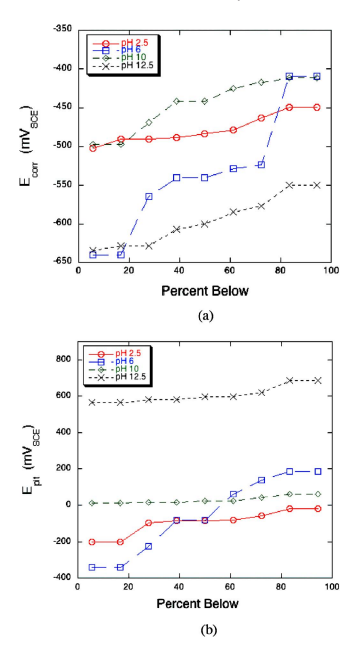


Figure 9. (Color online) Distribution plots of (a) E_{corr} and (b) E_{pit} for Al₃Fe collected in 0.1 M NaCl at pH 2.5, 6, 10, and 12.5.

current gradually increases as potentials more anodic than E_{corr} are realized. The currents sustained by MgZn₂ at potentials separated from $E_{\rm corr}$ by more than about 50 mV are comparatively large, realizing several hundreds of $\mu A/cm^2$. Figure 10 reveals that, across the range of pH tested, the $E_{\rm corr}$ of MgZn₂ is nominally confined between -1000 and -1050 mV_{SCE}. This insensitivity to pH is unique among the phases presented herein. While Mg(OH)2 solubility is low at an elevated pH, the results here tend to support that, into the range of alkaline pH, the electrochemical response of MgZn₂ appears to be dominated by the electrochemical behavior of Zn. Mg₂Si.— Figure 5 reveals the response of Mg₂Si. In an acidic solution, concomitant with the most noble $E_{\rm corr}$ (Fig. 11a), Mg₂Si shows high rates of dissolution with no evidence of passivity and dissolution rates of tens of mA/cm² for potentials greater than nominally about $-1.3\,$ V $_{SCE}$. No apparent passivity or high rates of dissolution are also seen at pH 6 and 9. However, in the case of pH 12.5, a very reproducible window of passivity is observed over a range of about half a volt from E_{corr} , prior to a breakdown process being observed.

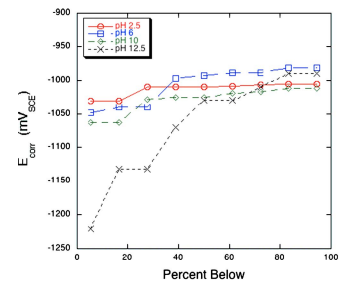


Figure 10. (Color online) Distribution plots of $E_{\rm corr}$ for MgZn₂ collected in 0.1 M NaCl at pH 2.5, 6, 10, and 12.5.

This phenomenon suggests that at such an elevated pH Mg_2Si is spontaneously passive, with an associated decrease in the dissolution rate by about two orders of magnitude over a wide range of potentials. The $E_{\rm pit}$ at an elevated pH is reproducible within a well-defined window as seen in Fig. 11b. This passivity at an elevated pH is the likely result of Mg stability at an elevated pH, suggesting the response of Mg_2Si is dominated by Mg. Figure 5 also reveals that the repassivation behavior varies as a function of the pH, suggesting a selective incongruent dissolution of Si (discussed further below).

Al₂CuMg.— As previously noted, the electrochemical response of S-phase (Al₂CuMg) is complex and often difficult to reproduce within a small error band, ^{20,21} as also evidenced by Fig. 12. This is owing largely to the tendency of the compound to undergo a "dealloying" process that involves the selective dissolution of Mg and Al, with the associated enrichment of Cu at the compound surface. The response of the compound (electrochemically) will therefore ultimately be heavily influenced by the Cu-rich surface layer. The results in Fig. 6 indicate the typical electrochemical response observed in this testing. With the exception of the highest pH tested, $E_{\rm corr}$ values were nominally in the general vicinity of about -850 to -900 V_{SCE}. In the case of pH 2.5 and 6, what is nominally observed is an apparent range of passivity prior to a breakdown process. It is noted, however, that the current density corresponding with this range is rather high at $> 100 \mu A/cm^2$, indicating that the system may not be "passive" at these potentials. This has been previously observed and it was hypothesized that the high current densities realized prior to the apparent breakdown are owing to the dealloying (i.e., a dissolution) process, with the ultimate observed breakdown corresponding to the activation of an essentially pure Cu-rich surface layer. Further information regarding this hypothesis and its observation is not given in this paper; however, we do present the polarization curves as collected in order to convey the information herein as a survey of the results. As the pH increases, the apparent passive potential decreases to lower currents, suggesting that the activity of the compound decreases with pH. At pH 12.5, this is concomitant with an increase in $E_{\rm corr}$ to more noble values, suggesting that the activity of the compound decreases with increasing the pH. The repassivation potential is independent of the solution pH.

In contrast to prior work, the net effect of increasing the pH is not as straightforward as we observed for increasing the NaCl concentration. In the case of NaCl variation, we nominally observed a decrease in $E_{\rm corr}$ (to less noble values) concomitant with a

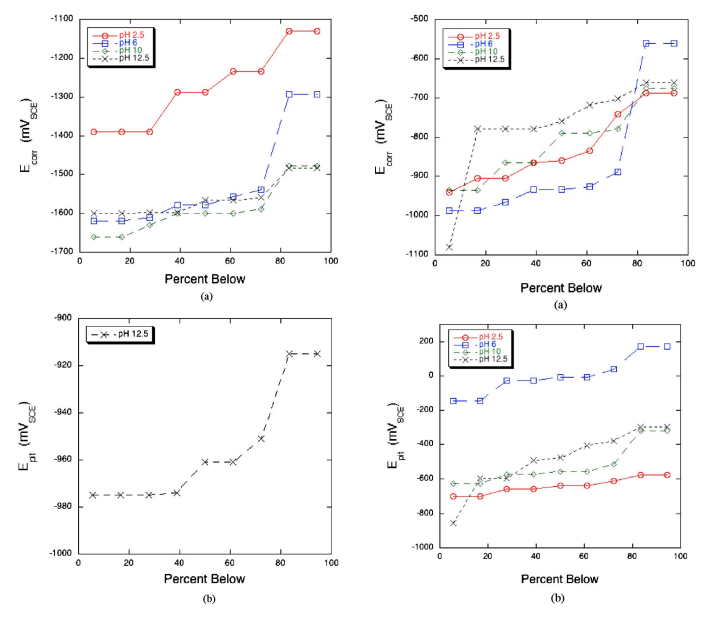


Figure 11. (Color online) Distribution plots of (a) $E_{\rm corr}$ and (b) $E_{\rm pit}$ for Mg₂Si collected in 0.1 M NaCl at pH 2.5, 6, 10, and 12.5.

Figure 12. (Color online) Distribution plots of (a) $E_{\rm corr}$ and (b) $E_{\rm pit}$ for Al₂CuMg collected in 0.1 M NaCl at pH 2.5, 6, 10, and 12.5.

decrease in $E_{\rm pit}$ as the NaCl concentration was increased from 0.01 to 0.6 M. In the case of altering the pH, the associated thermodynamics of each compound will vary depending on whether the solution is acidic, neutral, or basic; to an extent analogous to where the regions of stability may be found on a Pourbaix diagram. Consequently, not only is it difficult to predict the change in $E_{\rm corr}$ of the compounds with environment, but the change in $E_{\rm corr}$ reflecting thermodynamics will not intuitively correspond to the observed kinetics of the system in response to polarization. This is true for all the compounds tested here, reflecting the importance of polarization testing. We note and emphasize that there is a dispersion in the results presented in Fig. 1-6; therefore, the statistical distributions are given in Fig. 7-12. This serves to highlight that characteristic electrochemical parameters do vary within an envelope, while dispersion itself is a characteristic of intermetallic behavior.

Cathodic polarization response of intermetallic particles as a function of pH.— In order to more specifically determine the electrochemical characteristics of intermetallics typically noted as being "cathodic" to the matrix of common aluminum alloys [broadly de-

fined as $E_{\rm corr}({\rm intermetallic}) > E_{\rm corr}({\rm alloy})]$, cathodic polarization curves were conducted upon Al $_7$ Cu $_2$ Fe, Al $_2$ Cu, and Al $_3$ Fe, all of which nominally display an $E_{\rm corr}$ more positive than -700 mV $_{\rm SCE}$.

Figure 13 indicates the cathodic polarization characteristics of Al_7Cu_2Fe , revealing that this compound is capable of sustaining relatively large cathodic current densities over the potential range tested. The response at pH 2.5 and 6 is nominally very similar, with a mean $E_{\rm corr}$ in the vicinity of $-550~{\rm mV_{SCE}}$ and the ability to sustain cathodic currents in the range of several hundred $\mu A/{\rm cm}^2$ at potentials $<-800~{\rm mV_{SCE}}$. As the pH was increased, the $E_{\rm corr}$ became more negative by about 100 mV, with a concomitant increase in the ability to sustain a cathodic current. Rates as high as several mA/cm² were typically observed over the potential range tested. This result may not have been predicted by analysis based on potential differences alone; however, the kinetics as determined from polarization testing shows enhanced rates of reaction, revealing $E_{\rm corr}$ is not a satisfactory parameter to interpret, or even serve as a guide to, kinetics.

Figure 14 reveals the cathodic polarization response of Al₂Cu.

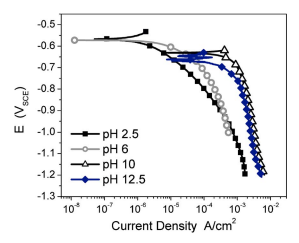


Figure 13. (Color online) Cathodic polarization diagrams for Al_7Cu_2Fe collected in 0.1 M NaCl at pH 2.5, 6, 10, and 12.5.

Within a rather well-defined band and neglecting variations in the $E_{\rm corr}$ values, the cathodic response is rather similar for ${\rm Al}_2{\rm Cu}$ independent of the solution pH. The values of the cathodic current are nominally somewhat below those of ${\rm Al}_2{\rm Cu}_2{\rm Fe}$ for similar potentials at pH 2.5 and 6, and significantly lower at basic pH. These results correlate with those of Scully et al. 7 for ${\rm Al}_2{\rm Cu}$, whereby $E_{\rm corr}$ was found to decrease with increasing the pH; however, in that case the testing was conducted in an inert (Cl $^-$ -free) solution.

In Fig. 15 we see the cathodic polarization curves for Al_3Fe . Under acidic conditions, the cathodic current sustained is nominally about an order of magnitude lower than the values observed at a neutral or basic pH. The response is very similar at pH 6, 10, and 12.5 over a wide range of potentials. With the exception of pH 2.5, Al_3Fe appears to sustain somewhat greater cathodic currents than Al_2Cu , but somewhat lower currents than Al_7Cu_2Fe . Again, the electrochemical response of Al_3Fe as determined herein was not necessarily obvious based on E_{corr} measurements alone, as was previously reported, 15 further highlighting the importance of the presentation of kinetic polarization data such as those given here.

Comments regarding electrochemical behavior of intermetallic particles as a function of pH.— The observed differences in the electrochemical properties for the various compounds tested reflect the electrochemical heterogeneity that may be found in many commercial Al alloys. We note that the electrochemical response varies greatly depending on both the intermetallic composition and the solution pH.

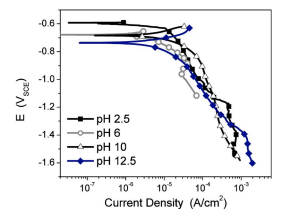


Figure 14. (Color online) Cathodic polarization diagrams for Al_2Cu collected in 0.1 M NaCl at pH 2.5, 6, 10, and 12.5.

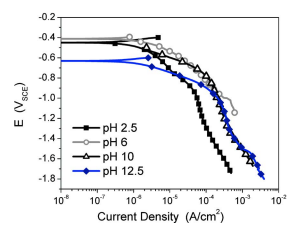


Figure 15. (Color online) Cathodic polarization diagrams for Al₃Fe collected in 0.1 M NaCl at pH 2.5, 6, 10, and 12.5.

The results in Fig. 1-3 indicate that Al₇Cu₂Fe, Al₂Cu, and Al₃Fe reveal a characteristic breakdown potential, hence capable of maintaining some form of surface film, over the range of pH tested. This suggests that such compounds are stable in NaCl environments. It is noted that, as the pH increased, the cathodic current sustained increased in the case of Al₇Cu₂Fe and Al₃Fe. In the case of the former, the cathodic current density rapidly approached the mA/cm² range. In the context of localized corrosion of Al alloys in aerated conditions, this is significant. For example, if such cathodic intermetallics are the preferred sites for oxygen reduction, then the associated hydroxyl ion generation should cause a local pH increase (tending toward alkaline conditions) at the intermetallic surface for a nominally neutral pH bulk solution. In the case where cathodic efficiency increases with the pH, such as the Fe-containing intermetallics, then their potency as local cathodes ought to increase as localized corrosion proceeds. The precise reason why the Fe-containing intermetallics increase in cathodic efficiency is not specifically studied here by way of surface analysis or chemistry, but is posited to be as the result of Fe stability at an elevated pH. For example, Al₇Cu₂Fe compared to Al₂Cu indicates a greater cathodic efficiency, linked to the presence of Fe. Similarly, Al₃Fe shows a greater cathodic efficiency than Al₂Cu.

The work herein has highlighted that Al₇Cu₂Fe, Al₃Fe, and Al₂Cu are all capable of functioning as potent local cathodes, owing to the appreciable cathodic currents sustained over a wide range of potentials typical of Al alloys^{2,12} across the pH range tested. The traditional belief is that the associated local cloud of alkaline pH that emanates from a cathodic particle (owing to hydroxyl ion generation from oxygen reduction) is principally responsible for corrosion of the adjacent matrix in near-neutral bulk solutions, because Al and its solid solutions nominally dissolve in alkaline conditions. ^{22,23} Recent calculations and detailed examinations of peripheral pitting, however,³ indicated that it does not appear likely that the local alkalinity alone causes a significant peripheral dissolution. The details of such work are given in the respective papers, emphasizing that the dominant mechanism for peripheral pitting may be largely due to charge transfer, which, based on the work presented herein, could be possibly amplified by enhanced oxygen reduction kinetics upon the cathodic intermetallics themselves (as the local pH on the intermetallic surface increases) to support self-sustaining acidic corrosion pits. Evidence for anodic dissolution within the peripheral pits is provided by the hydrogen bubble evolution in direct observation studies of localized corrosion in Al alloys, ²⁴ suggesting that overall alloy corrosion, at least initially, proceeds almost selfcatalytically.

In contrast, $MgZn_2$ did not reveal any stability or passive response over the range of pH tested, indicating that such a compound will ultimately undergo appreciable dissolution in the environments

Table I. Tabulated electrochemical data as a function	of pH for intermetallic	compounds tested in this study.
---	-------------------------	---------------------------------

Stoichiometry	Phase	pН	$E_{ m corr}$ m ${ m V}_{ m SCE}$ $^{ m a}$	$E_{ m pit}~{ m mV}_{ m SCE}^{~~a}$	Current (A/cm^2) at $-900 \text{ mV}_{SCE}^{b}$
Al ₃ Fe β	β	2.5	-510	-50	-4.2×10^{-5}
	•	6	-539	106	-1.6×10^{-4}
		10	-408	40	-1.2×10^{-4}
		12.5	-230	550	-1.1×10^{-4}
Al_2Cu θ	θ	2.5	-546	-458	-3.8×10^{-5}
		6	-665	-544	-4.1×10^{-5}
		10	-739	-408	-1.1×10^{-4}
		12.5	-743	-407	-3.8×10^{-5}
$MgZn_2$ M, η	Μ, η	2.5	-1007	-	1.2×10^{-4}
	•	6	-1003	-	1.0×10^{-3}
		10	-999	-	5.0×10^{-4}
		12.5	-1012	-	4.8×10^{-4}
Mg_2Si	β	2.5	-1408	-	3.4×10^{-3}
	•	6	-1538	-	3.4×10^{-4}
		10	-1430	-	9.6×10^{-4}
		12.5	-1553	-951°	5.0×10^{-5}
Al ₇ Cu ₂ Fe	-	2.5	-535	-451	-3.2×10^{-4}
		6	-551	-448	-4.2×10^{-4}
		10	-604	-420	-2.4×10^{-3}
		12.5	-594	-410	-2.6×10^{-3}
Al ₂ CuMg	S	2.5	-750	-	-2.0×10^{-5}
		6	-883	80^{d}	$\pm 1 \times 10^{-5}$
		10	-850	-390^{d}	$\pm 1 \times 10^{-6}$
		12.5	-670	-350	-3.6×10^{-5}

^a Dispersion of these values seen in Fig. 7-12.

tested and as a component of Al alloys, with microstructural evidence of this existing. 13 This is significant because the MgZn₂ is present in a large number density for 7xxx series Al alloys, 13,25,26 and more specifically populates the grain-boundary regions.

The electrochemical response of Mg_2Si reveals some interesting characteristics. Under acidic conditions, high dissolution rates are observed (viz., in the mA/cm² range). No evidence of passivity is observed at pH 6 or 10; however, at pH 12.5, the dissolution characteristics of Mg_2Si are markedly different. There is a well-defined window of passivity that may be expected to be observed for pure Mg at elevated (>11) pH. Consequently, for environments where the bulk solution is not heavily basic, significant self-dissolution of Mg_2Si ought to be expected. Other recent work in our laboratory revealed that, during dissolution of Mg_2Si , there is an apparent incongruent dissolution, whereby the loss of Mg reveals remnant Si on the alloy surface, suggesting that Mg may dominate the electrochemical response of Mg_2Si .

As previously suggested, the electrochemical behavior of Al $_2$ CuMg is complex and unique owing to the de-alloying processes. Unambiguous characterization of the precise kinetics and associated phenomenology of the electrochemical response ultimately requires an associated surface analysis and detailed analytical microscopy. 21,28 However, for the purpose herein, the potentiodynamic results presented in Fig. 6 indicate that, as pH tends toward basic conditions, the attendant passive current density decreases remarkably to about 10 $\,\mu\text{A/cm}^2$. This may indicate a reduction in the de-alloying processes. Again, this could be associated with the thermodynamic stability of Mg at an elevated pH, and may also be influenced by the low electrochemical activity of Cu at an elevated pH. $^{4.5}$

The data presented herein may be incorporated into a discussion of the corrosion damage evolution in the context of a 7xxx series Al alloy. The $E_{\rm corr}$ of most Al-Zn-Mg based alloys, is nominally in the

vicinity of about -900~mV in a quiescent neutral NaCl solution. 8,12,29 Information pertaining to the average currents measured upon compounds tested as a function of the pH at such a potential is included in Table I (along with mean E_{corr} and E_{pit} values). This information provides a deeper insight into the overall behavior of the alloy, further indicating the weaknesses of E_{corr} values or potential gradients (i.e., the difference between the E_{corr} of the particle and the matrix) alone as the basis for mechanistic interpretations of alloy corrosion. Similarly, this indicates the weakness of grouping together all particles as simply "anodic" or "cathodic", or only characterizing particles in a qualitative manner, which is often the case in prior works. Processes such as de-alloying, incongruent dissolution, a heavily pH-dependent electrochemical response, and current sustaining ability (which has been shown to be unrelated to the value of $E_{\rm corr}$ or even $E_{\rm pit}$, which is commonly used as a measure of the relative nobility of intermetallics) need to be uniquely considered in understanding the extent, morphology, and rate of the corrosion damage. Such information may also be useful in the formulation of customized corrosion protection regimes, because certain aspects of the corrosion cycle, i.e., Fe-containing particles leading to peripheral pitting (Fig. 16), can be identified as key contributors to the corrosion cycle and be targeted by the appropriate inhibitors.

Behavior and corrosion morphology of AA7075-T651 as a function of bulk solution pH.— The above discussion related largely to the localized corrosion in a near-neutral pH solution, where pH gradients can exist locally upon the alloy surface. This section, however, deals with varying the pH of the bulk solution. Thus specific tests were conducted on AA7075-T651 in order to reconcile the fundamental (microcell) electrochemical information collected and presented with damage evolution upon a bulk alloy, combining some results with the discussion.

^b Average values reported. Results varied typically within 20% of the average.

^c Breakdown was only revealed for this compound for the highest pH tested (pH 12.5).

^d E_{corr} of this compound is in the vicinity of -900 mV_{SCE}, making it difficult to yield an approximate average current because in cases it may be net anodic or net cathodic; however, in all cases the current is low with an absolute value $<1 \times 10^{-5}$ A/cm².

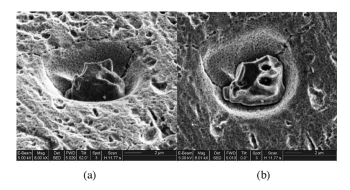


Figure 16. High-resolution SEM images revealing the corrosion morphology associated with an Al_7Cu_2Fe particle in AA7075-T651 immersed for 10 h in 0.1 M NaCl at pH 6. Images taken at (a) 52° tilt, (b) 0° tilt (secondary electron image).

Although an acid or alkaline bulk environment nominally represents unlikely extremes in the prescription of Al alloys, time or service conditions may conspire to produce such undesirable conditions. The examples herein will highlight that, in cases where the local electrolyte may tend to acid or alkaline conditions, the attendant damage morphology then will be different, while such exposures can complement the individual microcell results collected for specific intermetallics.

Figure 17 reveals the evolution of open-circuit potential (OCP) of AA7075-T651 for acid (pH 2.5), near-neutral (pH 6), and alkaline (pH 12.5) conditions. We see that in acid conditions the alloy initially displays a comparatively noble OCP which decreases with time toward $\sim\!-900~{\rm mV_{SCE}}.$ For neutral conditions we see the well-documented response that, following an initial transient period commencing at OCP $<-1~{\rm V}$, the potential varies within a narrow window close to $\sim\!-750~{\rm mV_{SCE}}.$ In the case of an alkaline solution, the OCP reveals a marked decrease, commencing at a comparatively very negative value of $\sim\!-1650~{\rm mV_{SCE}},$ followed by an ennoblement with time.

The OCP testing suggests a markedly different electrochemical response for the bulk alloy with pH, and this is seen clearly in Fig. 18 for potentiodynamic testing upon AA7075-T651. We see, at a near-neutral pH, a very small window of passivity (i.e., $<100~\rm{mV}).^{26}$ This window is however time dependent and diminishes with continued immersion. 26 However, at pH 2.5, dissolution occurs at high rates for $E>E_{\rm corr}.$ Conversely, the self-dissolution rate for testing at pH 12.5 is nominally nearly two orders of magnitude greater, and again no passivity is seen.

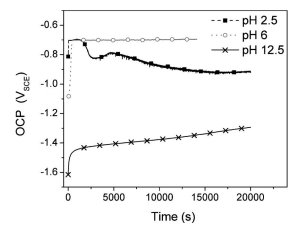


Figure 17. OCP vs time for AA7075-T651 collected in 0.1 M NaCl at pH 2.5, 6, and 12.5.

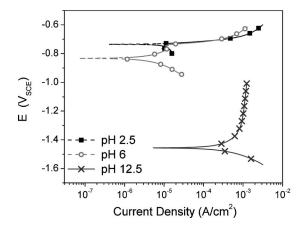


Figure 18. Potentiodynamic polarization diagrams for AA7075-T651 collected in 0.1 M NaCl at pH 2.5, 6, and 12.5.

Figure 19 reveals the typical damage morphology, as determined by SEM, for AA7075-T651 in 0.1 M NaCl at pH 2.5 following 24 h immersion. A large-scale pitting is observed. These pits are in no way associated with cathodic particles (which are seen as light stringers in the SEM image of Fig. 19). What is seen is loss (i.e., dissolution) of "anodic" particles (viz., Mg2Si and MgZn2) as gauged by their lack of presence from navigating the sample. However, pitting is also randomly occurring, to a large extent, in the matrix at local sites, likely related to stochastic defects in the matrix oxide and in accordance with the thermodynamic instability of Al and its solid solutions in acid conditions. The determination of the corrosion morphology in Fig. 19 is important, as it highlights physically the thermodynamic stability of so-called "cathodic" particles, because they remain largely unaffected as the corrosion proceeds. Furthermore, particles such as Mg2Si and MgZn2, expected to be dissolved, show no evidence of existence, in accordance with the microcell test prediction.

A higher resolution image of the damage seen in an apparently unaffected area in Fig. 19 is seen in Fig. 20. In this case, a strong secondary electron contrast was used to reveal localized pitting patterns, such that they appear to reveal a "mesh" or network of damage (see the darkest regions), possibly associated with individual grain boundaries. Such a detailed investigation is the focus of future work. Figure 20 again highlights that corrosion damage appears unrelated to cathodic particles, which can be summarily defined as stochastic in its nature.

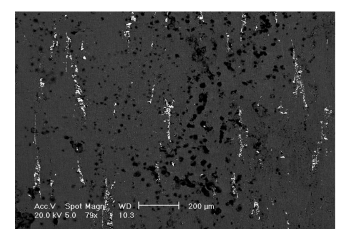


Figure 19. SEM image of AA7075-T651 exposed in 0.1 M NaCl at pH 2.5 for 24 h. Electron detector used was an equal mix of secondary and back-scattered electrons.

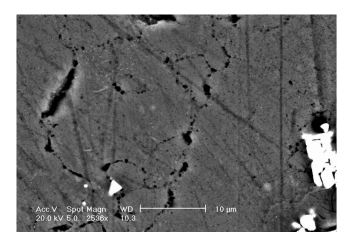


Figure 20. Higher magnification SEM image of AA7075-T651 exposed in 0.1 M NaCl at pH 2.5 for 24 h (secondary electron image).

For the case of exposure to an alkaline solution (0.1 M NaCl at pH 12.5), the associated damage is seen in Fig. 21. The exposure in this case was limited to 5 h because large-scale corrosion and sample discoloration was evident. It is clear that the damage morphology is markedly different from acid exposure, and indeed neutral pH exposure. ¹³ Again, corrosion is not associated directly with cathodic intermetallics, although such intermetallics can likely function as cathodes nonetheless. Instead, the damage is best described as a large-scale matrix dissolution in a general sense, concomitant with the large currents revealed by the potentiodynamic testing (Fig. 18). This matrix dissolution is clearly evident from Fig. 21, because secondary electrons were used to generate the image such that the light particles represent material that sits "proud" of the (receded) surface. Furthermore, the "mud cracking" on the matrix materials is related to oxide cracking, posited to be associated with a corrosion product several µm in thickness. In fact, this damage phenomenon is so marked that it is visible with the naked eye given the correct illumination.

The somewhat higher resolution images in Fig. 22 emphasize the reconciliation between the microcell results and the observed damage. Figure 22a shows a secondary electron image such that intermetallic particles appear proud of the surface; however, the associated backscattered electron image reveals an interesting result. Particle A, which was seen to be unaffected in Fig. 22a, appears dark on the basis of the compositional contrast offered by the backscattered electron imaging. Subsequent EDXS reveals the presence of

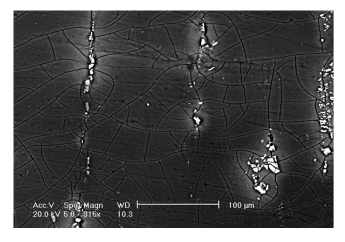


Figure 21. SEM image of AA7075-T651 exposed in 0.1 M NaCl at pH 12.5 for 5 h (secondary electron image).

abundant Mg and Si as seen in Fig. 22c, whereas the stringer particle reveals the presence of Cu and Fe, Fig. 22d. Based on prior alloy stereology work, we believe these particles to be Mg_2Si and Al_7Cu_2Fe . This therefore confirms that Al_7Cu_2Fe -type particles and indeed the Mg_2Si intermetallic are unattacked at an elevated pH, as predicted from the microcell testing. The physical response of both Al_2CuMg and Al_2Cu was not determined from the exposure of AA7075-T651 because these compounds appear in a low number density in such an alloy, and are often difficult to resolve when corrosion has occurred. 13

General discussion.— Overall, the work herein extends beyond the somewhat first-order notion that the corrosion of such alloys is simply related to the microstructure, indicating that the relationship is especially complex and heavily dependent on the environment and the level of corrosion being experienced, because the latter can also influence the microenvironment in the vicinity of anodic or cathodic activity upon the metal surface. The incorporation of such a complexity is critical in the prediction of corrosion, such that any predictive models do not under- or overpredict corrosion damage based upon an incorrect assessment or lack of sensitivity of corrosion upon the environment. In the case of aircraft, the corrosive environment will alter significantly, both as electrolytes become concentrated or diluted, and as corrosion proceeds. We explored the physical aspects of the fundamental microcell data via the extreme conditions of bulk alloy exposure in highly acidic and highly alkaline bulk environments as two model examples. Although such conditions may not be encountered in general service, the testing highlights the pH-dependent response of intermetallic particles, along with the bulk alloy, and conditions that could be seen locally in a corroding specimen. It is seen rather clearly that the mode of corrosion upon AA7075-T651 is stochastic at acid pH, general at alkaline pH, and largely deterministic, based heavily on the stereology (i.e., microstructure) and electrochemistry of intermetallics, at a nearneutral pH. This has vast consequences in the development of predictive models for localized damage accumulation, highlighting the notion that the environment is as critical as the microstructure in terms of corrosion, and that the ultimate damage represents the intersection of the two. This also indicates heavily that the correct interpretation of the corrosion of such alloys must be sensitive to the environment; hence, qualitative indices such as the Volta potential³⁰ are of limited use for such complex systems because the Volta potential is a qualitative value measured in air; which has no relevance to what may occur in say, as an example, alkaline pH. Another critical factor that represents work in progress is the electrochemical characteristics as a function of temperature.

Conclusions

The information provided herein is a useful resource for clarification pertaining to the electrochemical behavior of Al alloys, drawing together a large amount of characteristic data in standard test solutions to allow for a consolidated presentation. It is seen that the electrochemical response of intermetallics that populate high strength Al alloys, tested herein, is heavily dependent on the pH of the environment, and significantly more complex than the relative activity or nobility that may be guessed from the composition or corrosion potential, while thermodynamics may not be a satisfactory index of kinetics. A detailed treatise of this was not presented on theoretical grounds; however, empirical data reported clearly reveal this to be the case. A simple example of this is that, in cases where Mg_2Si reveals its noblest potential (in acid pH), it also displays its highest dissolution rates.

Some highlights include the finding that the intermetallics containing Fe increase their cathodic efficiency as the pH increased to more alkaline values. This can have attendant ramifications in the damage evolution upon bulk alloys in near-neutral conditions, and indicates both that large amounts of corrosion ought to be synonymous with such particles, while any corrosion protection mechanism should also be likely aimed at incorporating a mechanism for halting

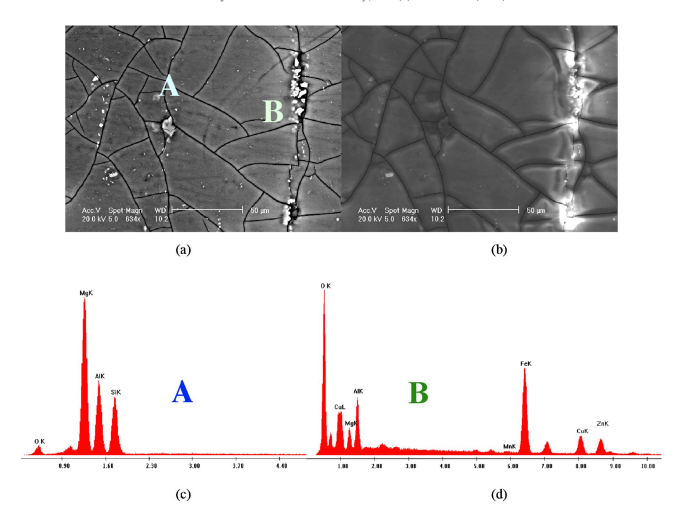


Figure 22. (Color online) Higher magnification SEM image of AA7075-T651 exposed in 0.1 M NaCl at pH 12.5 for 5 h. (a) Secondary electron image. (b) Backscattered electron image. (c) EDXS spectrum of particle A, revealing Mg and Si along with matrix background. (d) EDXS spectrum of particle B, revealing Fe and Cu along with matrix background.

such cathodic activity. It was also seen that Mg₂Si, nominally spontaneously active, was capable of displaying active-passive behavior at pH 12.5, suggesting that its electrochemical response is dominated by Mg, allowing the formation of a stable oxide film at an alkaline pH.

Investigation of the morphology of corrosion following exposure to acid and alkaline environments for AA7075-T651 reconciles the fundamental microcell data and highlights the highly pH-specific damage evolution modes, and attendant morphology, that occur in Al alloys. This allows us to now interpret corrosion damage in a far more detailed, quantitative, and meaningful manner.

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

References

- N. Birbilis and R. G. Buchheit, J. Electrochem. Soc., 152, B140 (2005).
 G. O. Ilevbare, O. Schneider, R. G. Kelly, and J. R. Scully, J. Electrochem. Soc.,
- 3. M. Büchler, T. Watari, and W. H. Smyrl, Corros. Sci., 42, 1661 (2000).
- 4. M. Pourbaix, Atlas of Electrochemical Equilibria in Aqueous Solutions, NACE, TX (1974).
- 5. O. Schneider, G. O. Ilevbare, R. G. Kelly, and J. R. Scully, J. Electrochem. Soc., 151, B465 (2004).
- G. O. Ilevbare and J. R. Scully, J. Electrochem. Soc., 148, B196 (2001).
- J. R. Scully, T. O. Knight, R. G. Buchheit, and D. E. Peebles, Corros. Sci., 35, 185

- 8. R. G. Buchheit, J. Electrochem. Soc., 142, 3994 (1995).
- R. P. Wei, C.-M. Liao, and M. Gao, Metall. Mater. Trans. A, 29, 1153 (1998).
 P. Leblanc and G. S. Frankel, J. Electrochem. Soc., 149, B239 (2002).
 I. J. Polmear, Light Alloys, 3rd ed., Arnold, London (1995).

- 12. Q. Meng and G. S. Frankel, J. Electrochem. Soc., 151, B271 (2004).
- 13. N. Birbilis, M. K. Cavanaugh, R. G. Buchheit, D. G. Harlow, and R. P. Wei, in Proceedings of Symposium: Application of Materials Science to Military Systems, Materials Science and Technology '05, Pittsburgh, PA, TMS (2005).
- 14. L. F. Mondolfo, The Aluminum-Magnesium-Zinc Alloys, Research and Development Center, Revere Copper and Brass Inc., Rome, NY (1967).

 15. J. O. Park, C. H. Paik, Y. H. Huang, and R. C. Alkire, *J. Electrochem. Soc.*, **146**,
- T. Suter and H. Böhni, Electrochim. Acta, 43, 2843 (1998).
- M. Lohrengel, Corros. Eng., Sci. Tech., 39, 53 (2004).
 T. Suter and R. C. Alkire, J. Electrochem. Soc., 148, B36 (2001).
- 19. N. Birbilis, B. N. Padgett, and R. G. Buchheit, Electrochim. Acta, 50, 3536 (2005).
- 20. R. M. Leard, M.Sc. Thesis, The Ohio State University, Columbus, OH (2001).
- 21. R. G. Buchheit, M. A. Martinez, and L. P. Montes, J. Electrochem. Soc., 147, 119 (2000).
- T. J. R. Leclère and R. C. Newman, J. Electrochem. Soc., 149, B52 (2002).
- 23. Z. Szklarska-Smialowska, Corros. Sci., 41, 1743 (1999).
- 24. C. M. Liao, J. M. Olive, M. Gao, and R. P. Wei, Corrosion (Houston), 54, 451 (1998).
- 25. P. K. Poulose, J. E. Morral, and J. McEvily, Metall. Trans., 5, 1393 (1974).
- 26. J. A. Wert, Scr. Metall., 15, 445 (1981).
- S. Jain, Ph.D. Thesis, The Ohio State University, Columbus, OH (2006).
- 28. R. G. Buchheit, R. P. Grant, P. F. Hlava, B. McKenzie, and G. L. Zender, J. Electrochem. Soc., 144, 2621 (1997).
- 29. N. Birbilis, M.K. Cavanaugh, and R.G. Buchheit, ECS Trans., 1(4), 115 (2006).
- 30. J. H. W. de Wit, Electrochim. Acta, 49, 2841 (2004)