Pentavalent Vanadium Species as Potential Corrosion Inhibitors of Al$_2$Cu Intermetallic Phase in the Sulfuric(VI) Acid Solutions

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Abstract: The objective of this work was to test vanadium isopolyoxoanions as potential corrosion inhibitors of the intermetallic phase Al$_2$Cu in sulfuric acid solutions at pH = 1.3 and 2.5. The intermetallic was melted in an electric arc furnace. Its phase composition was confirmed using X-ray diffraction, light microscopy, and differential scanning calorimetry. Then Al$_2$Cu corrosion kinetics was studied. Chemical composition of the solution after corrosion was determined using inductively coupled plasma-optical emission spectroscopy. The surface of corroded specimens was analyzed using scanning electron microscopy and X-ray photoelectron spectroscopy. Subsequent electrochemical studies involved determination of open-circuit potential, electrochemical impedance spectra, and polarization curves. It was found that the Al$_2$Cu phase corrodes selectively and vanadium isopolyoxoanions increase this process both at pH = 1.3 and 2.5 with two exceptions. Corrosion inhibition was observed for 100 and 200 mM of Na$_3$VO$_4$ at pH 1.3, with inhibition efficiency 78% and 62% respectively, due to precipitation of V$_2$O$_5$.

Keywords: intermetallics; selective corrosion; sodium orthovanadate; isopolyoxoanions

1. Introduction

Aluminum is the most widely applied non-ferrous metal. Components made of aluminum alloys are mainly used in transportation, engineering, construction, and packaging, where their good strength, corrosion, and wear resistance is often required. Phase composition of aluminum alloys can be very complex because this metal forms numerous intermetallics with alloying elements and impurities. These intermetallics significantly affect alloy properties e.g., strength and corrosion resistance. One of the most important intermetallic in aluminum alloys is Al$_2$Cu. It ensures good strength of Al-Cu-Mg-Mn and Al-Zn-Mg-Cu wrought alloys as well as Al-Cu cast alloys [1]. Unfortunately, the precipitation of intermetallic in the matrix diminishes corrosion resistance of these alloys. This occurs due to strong galvanic coupling between Al$_2$Cu and the matrix and local dissolution of the latter [2–10]. At the same time, Al$_2$Cu itself is susceptible to the selective corrosion in acidic and alkaline solutions [2,3,11,12].

Corrosion studies of intermetallic phases are important for understanding the corrosion mechanism of copper-rich aluminum alloys. Components made of these alloys are usually applied in neutral environments. Their surface treatment, however, often requires acid etching. This occurs e.g., when anodic coatings are stripped in hot orthophosphoric acid solution. Presently, industry commonly uses chromium trioxide to prevent corrosion of the metallic substrate during etching/stripping [13]. Unfortunately, hexavalent chromium species are highly toxic [14]. Thus, new, less toxic inhibitors must be developed. Sodium molybdate was successfully applied instead of chromium trioxide in the
stripping bath. Unfortunately, sufficient corrosion inhibition was achieved only for aluminum 1050, but not for commonly applied 2024 alloy [15–17]. It was also found that sodium molybdate decreases the dissolution rate of anodic coating [18].

Recently, sodium orthovanadate, tungstate, and molybdate, were tested as the potential inhibitors of corrosion of Al2Cu intermetallic phase in orthophosphoric acid [12]. When adding increasing amounts of a mineral acid to aqueous solutions of orthometalates, addition and condensation processes occur and isopolyoxoanions are formed. They are composed of a transition metal, oxygen, and hydrogen. Condensation in a solution containing e.g., orthophosphoric or silicic acid gives heteropolyoxoanions. They incorporate one or more elements e.g., phosphorus or silicon, in addition to the transition metal, oxygen and hydrogen [19]. It was shown that neither molybdenum nor tungsten heteropolyoxoanions inhibit corrosion of intermetallic Al2Cu [12]. Corrosion was inhibited however in 0.5 and 1 M H3PO4 due to reduction of vanadium heteropolyoxoanions and precipitation of an insoluble salt [12,20].

In this work, isopolyoxovanadates were tested as corrosion inhibitors of intermetallic Al2Cu. Literature survey indicated that tetrahedrally coordinated species, existing in mildly acidic solutions, such as [H2VO4]− and [V4O12]4−, inhibit the cathodic process on aluminum [21]. They also decrease the open-circuit potential and corrosion current density and increase the pitting potential of intermetallic Al2Cu in 0.5 M NaCl solutions at pH = 9.17 [22]. Decavanadate ions, such as [HV10O28]4− and [H2V10O28]4+, which are stable in mildly and strongly acidic solutions, do not exhibit an inhibiting effect on the 2024 alloy [21,23]. Vanadium isopolyoxoanions, however, were not tested as corrosion inhibitors of intermetallic Al2Cu in acidic solutions. The idea of such studies is related to development of new corrosion inhibitors for aluminum alloys in acidic media. Although Al2Cu intermetallic in the alloy does not need corrosion protection because it is more noble than the matrix, the situation may change in the presence of the inhibitor. When it is adsorbed onto anodic sites e.g., on the matrix it may change polarity in the corrosion cell and Al2Cu phase will become exposed to corrosion. Alternatively, when Al2Cu surface is covered with an insoluble film, the cathodic reaction will be hampered, and corrosion rate of the alloy reduced. Anyway, successful replacement of toxic hexavalent chromium species requires systematic studies of the influence of potential corrosion inhibitors on the corrosion behavior of both the alloy and its microstructural constituents, both cathodic and anodic with respect to the matrix.

2. Materials and Methods

Al2Cu alloy, ca. 4.5 g, was obtained by melting of Al and Cu (99.999 wt.% purity, Alfa Aesar, Lancashire, UK under Ar atmosphere in an electric arc furnace using tungsten electrode. Weight loss of the specimen due to vaporization during melting was negligible (<0.1%). The alloy was annealed in air, T = 823 K, t = 30 h, cut using an electrical discharge machine, mounted in an epoxy resin, grinded with SiC abrasive paper (320 and 500 grit), rinsed with water and isopropyl alcohol, and air-dried. Geometric surface area of the Al2Cu specimen was 0.79 cm².

Microstructure of obtained specimen was revealed using Keller’s reagent (HF 1% v/v, HCl 1.5% v/v, HNO3 2.5% v/v, t = 45 s at the room temperature) and examined using light microscope Leica DMI 3000M, Leica Microsystems CMS GmbH, Wetzlar, Germany. Phase composition was confirmed using ARL X’Tra X-ray diffractometer, Thermo Fisher Scientific Inc., Waltham, U.S., with Cu Kα radiation source, and STA 449 F3 Jupiter thermal analyzer, Netzsch-Gerätebau GmbH, Selb, Germany. Differential Scanning Calorimetry (DSC) curves were recorded for 42 mg of the alloy, in PtRh20 crucible with Al2O3 liner obtained from Netzsch-Gerätebau GmbH, under He atmosphere, with heating rate 5 K·min⁻¹.

Corrosion rate of Al2Cu phase was determined as a function of pH and vanadium initial concentration cv. Na3VO4 was dissolved in 0.04 M sulfuric(VI) acid and solutions with cv = 0, 10, 50, 100, and 200 mM were obtained. Their pH was adjusted using concentrated H2SO4 at 1.3 and 2.5. UV-Vis absorption spectra of these solutions were measured using Cary 60 spectrophotometer, Agilent
Technologies Inc., Santa Clara, CA, US from 1100 to 190 nm in the quartz cuvettes with 10 mm optical path length. Samples were diluted 300 times using H₂SO₄ solutions with pH 1.3 or 2.5.

The volume of solutions used for corrosion tests was 100 cm³, temperature was 303 K and immersion time was equal to 576 min. All the solutions were in equilibrium with air. Aluminum and copper concentrations in solutions after corrosion tests were measured using inductively coupled plasma-optical emission spectroscopy ICP-OES (Ultima 2 Horiba Jobin Yvon SAS, Longjumeau, France) and corrosion rates were calculated.

Microscopic examination of the surface of selected corroded Al₂Cu specimens was performed using Phenom XL scanning electron microscope equipped with energy-dispersive X-ray spectrometer (EDX), Thermo Fisher Scientific Inc., Waltham, U.S. Further examination of corrosion product precipitated on the specimens was performed using X-ray photoelectron spectroscopy. Scanning XPS Microprobe PHI 5000 Versa Probe II (ULVAC-PHI, Chigasaki, Japan) system with microfocused (100 µm, 25 W) Al Kα X-ray beam was applied. The photoelectron spectra were acquired from 400 × 400 µm areas with analyzer pass energy of 46.95 eV and a photoelectron take-off angle of 45°. The charging effect was compensated using a dual-beam charge neutralizer. The operating pressure in the analytical chamber was <5 × 10⁻⁷ Pa. All XPS peaks were referenced to the oxide O1s peak at 531.0 eV . Pre-sputtering of the samples was performed by 10 min. with an argon gas cluster ion beam (Ar-GCIB) and the average Ar cluster size was 4000 atoms. The beam energy was set to 10 keV, beam current was 5 nA, and the sputtered area was 3 × 3 mm². The samples were rotated during pre-sputtering (Zalar rotation) to minimize surface roughening due to cluster bombardment. Application of the pre-sputtering by Ar-GCIB allowed to remove potential contamination on sample surfaces. Spectrum background was subtracted using the Shirley method. PHI MultiPak™ data analysis software (version 9.9.0.8) was used to calculate elemental compositions from the peak areas.

Electrochemical tests were conducted in a 3-electrode electrochemical cell open to air. Platinum wire with the surface area 20 cm² was applied as a counter electrode and Ag|AgCl (3 M KCl) as a reference electrode. The latter was placed in a Luggin probe filled with 1 M KNO₃ solution. The solution volume was 100 cm³ and temperature 303 K. The electrochemical cell was placed in a Faraday cage and connected to a SP-300 potentiostat, Bio-Logic SAS, Seyssinet-Pariset, France.

The open-circuit potential (OCP) of Al₂Cu was recorded for 576 min as a function of vanadium initial concentration, \( c_V = 0, 10, 50, 100, \) and 200 mM, and pH = 1.3 and 2.5. Further electrochemical analysis was limited to the following cases. Impedance spectra of the intermetallic together with polarization curves were measured at pH = 1.3, \( t = 576 \) min, as a function of vanadium concentration. An additional impedance spectrum was measured at pH = 2.5, \( c_V = 200 \) mM, \( t = 159 \) min to confirm the possibility of temporary corrosion inhibition at these conditions. The kinetics of corrosion of Al₂Cu was studied for \( c_V = 0 \) mM at pH = 1.3 and 2.5. In this particular case, the impedance spectra were recorded after \( t = 159, 298, 437, \) and 576 min. The frequency domain of all the measured impedance spectra was between 200 kHz and 10 mHz. A sinusoidal perturbation of potential equal to 5 mV of a root mean square was applied. The impedance spectra were validated using a Kramers–Kronig transformation with a KK Test software, version 1.01 [24,25]. Examples of obtained results are presented for \( c_V = 50 \) and 100 mM in Figure 1a and b, respectively. Random spread of residuals \( Z' \) and \( Z'' \) indicates sufficient stability of the studied system. These residuals were obtained using Equations (1) and (2) [24,25]:

\[
\Delta Z' = \frac{Z'_{\text{meas}} - Z'_{\text{calc}}}{|Z_{\text{calc}}|}
\]

\[
\Delta Z'' = \frac{Z''_{\text{meas}} - Z''_{\text{calc}}}{|Z_{\text{calc}}|}
\]

where \( Z'_{\text{meas}} \) and \( Z''_{\text{meas}} \) stand for measured values of a real and imaginary parts of the impedance, for a given frequency, whereas \( Z'_{\text{calc}} \) and \( Z''_{\text{calc}} \) were calculated using KK Test software. \( |Z_{\text{calc}}| \) indicates modulus of the calculated impedance.
was obtained after prolonged annealing (30 h) at 550 °C (XRD method). All the diffraction lines were ascribed to Al\(_2\)Cu tetragonal phase, International Centre for Diffraction Data (ICDD) card 04-001-0923 (data not shown). Because XRD is not able to detect possible small amount of impurities, additional characterization, using light microscopy (LM) and DSC, was conducted (Figure 2).

In our previous work it was shown that the Al\(_2\)Cu intermetallic was contaminated with a small amount of eutectic mixture of Al\(_2\)Cu and Al(Cu) solid solution [27]. This phase composition, however, is non-equilibrium one due to rapid crystallization. Thus, the single-phase alloy in the present work was obtained after prolonged annealing (30 h) at 550 °C. This was confirmed using X-ray diffraction (XRD) method. All the diffraction lines were ascribed to Al\(_2\)Cu tetragonal phase, International Centre for Diffraction Data (ICDD) card 04-001-0923 (data not shown). Because XRD is not able to detect possible small amount of impurities, additional characterization, using light microscopy (LM) and DSC, was conducted (Figure 2).

3. Results and Discussion

3.1. Characterization of Al\(_2\)Cu Intermetallic Phase

In our previous work, it was shown that the Al\(_2\)Cu intermetallic was contaminated with a small amount of eutectic mixture of Al\(_2\)Cu and Al(Cu) solid solution [27]. This phase composition, however, is non-equilibrium one due to rapid crystallization. Thus, the single-phase alloy in the present work was obtained after prolonged annealing (30 h) at 550 °C. This was confirmed using X-ray diffraction (XRD) method. All the diffraction lines were ascribed to Al\(_2\)Cu tetragonal phase, International Centre for Diffraction Data (ICDD) card 04-001-0923 (data not shown). Because XRD is not able to detect possible small amount of impurities, additional characterization, using light microscopy (LM) and DSC, was conducted (Figure 2).

Eutectic mixture would appear on the photomicrograph at the grain boundaries of Al\(_2\)Cu hypereutectic crystals. This however was not observed (Figure 2a). LM examination of the metallographic specimen revealed only large grains of Al\(_2\)Cu phase and cracks. The latter arose during crystallization because Al\(_2\)Cu phase is very hard and brittle.

DSC analysis definitely confirmed that the single-phase alloy was obtained. Only one endothermic peak appeared during heating with the onset at 862 K (Figure 2b). This temperature is very close to the peritectic temperature 863.5 K and well above the eutectic temperature (823 K) [28].
3.2. Corrosion Rate of Al$_2$Cu Intermetallic Phase

Isopolyoxovanadates were tested as potential corrosion inhibitors of intermetallic Al$_2$Cu in H$_2$SO$_4$ solutions at pH = 1.3 and 2.5. It was observed that at both pH, Al concentration in the solutions after $t = 576$ min Al$_2$Cu exposition was higher than that of Cu (Figure 3a,b). This suggests that the intermetallic corroded selectively. The higher the difference between concentration of both metals, the more severe possible dealloying of the intermetallic. Thus, at pH = 2.5, it occurred to much lesser extent when compared to pH = 1.3.

![Graph](image)

**Figure 3.** Concentration of Cu and Al in H$_2$SO$_4$ solutions after $t = 576$ min immersion of intermetallic Al$_2$Cu, $T = 303$ K as a function of the initial concentration of Na$_3$VO$_4$: (a) pH = 1.3; (b) pH = 2.5.

SEM examination of a specimen after corrosion confirmed its selective character. The surface of intermetallic Al$_2$Cu became porous and enriched with copper (Figure 4). The porous outer layer can be easily peeled off from the substrate e.g., during specimen rinsing or drying after the corrosion test. It should be noted here that color intensity differences on EDX maps came not only from the spatial distribution of elements but also from differences in height of the specimen.
The lowest values of corrosion rate were obtained at \( cV \geq 50 \text{ mM} \). The corrosion rate of the intermetallic was higher when compared to pH = 2.5, it decreased when \( cV \geq 50 \) mM. The lowest values of corrosion rate were obtained at \( cV = 0 \text{ mM} \). At pH = 2.5, its lowest value was obtained for \( cV = 0 \text{ mM} \). It then increased with increasing initial concentration of Na\(_3\)VO\(_4\), thus, no corrosion inhibition was observed. Although at pH = 1.3, \( cV = 0 \text{ mM} \) the corrosion rate of the intermetallic was higher when compared to pH = 2.5, it decreased when \( cV \geq 50 \text{ mM} \).

Data presented in Figure 3 were used for calculation of Al\(_2\)Cu corrosion rate (Figure 5). At pH = 2.5 its lowest value was obtained for \( cV = 0 \text{ mM} \). It then increased with increasing initial concentration of Na\(_3\)VO\(_4\), thus, no corrosion inhibition was observed. Although at pH = 1.3, \( cV = 0 \text{ mM} \) the corrosion rate of the intermetallic was higher when compared to pH = 2.5, it decreased when \( cV \geq 50 \text{ mM} \). The lowest values of corrosion rate were obtained at \( cV = 100 \) and 200 mM. They correspond to inhibition efficiency equal to 78% and 62%, respectively.

**Figure 4.** Micrograph of intermetallic Al\(_2\)Cu after corrosion in H\(_2\)SO\(_4\) solution without Na\(_3\)VO\(_4\), pH = 1.3, immersion time \( t = 576 \text{ min} \), \( T = 303 \text{ K} \) together with spatial distribution of elements, color intensity differences on EDX maps occur due to different concentration of elements and are also related to height of the specimen.

**Figure 5.** Corrosion rate of intermetallic Al\(_2\)Cu in H\(_2\)SO\(_4\) solutions after \( t = 576 \text{ min} \) at \( T = 303 \text{ K} \) as a function of pH and initial concentration of Na\(_3\)VO\(_4\).
Vanadium speciation in the studied solutions explains differences in the corrosion rate. UV-Vis spectroscopy revealed that different vanadium forms exist at pH = 1.3 and 2.5 (Figure 6a). The shape of the spectra and peak intensities for pH = 2.5 are different when compared to pH = 1.3 at corresponding vanadium initial concentrations. Solutions of isopolyoxovanadates at the higher pH absorb UV and visible light much stronger than at the lower pH. Color difference between vanadium solutions at these two pH values was visible with the unaided eye. It was also observed that change in vanadium initial concentration at a given pH changed only absorption peak intensities, but not their positions. This means that vanadium speciation does not change with increasing V(IV) provided that pH is constant. According to thermodynamic considerations, pentavalent vanadium species in strongly acidic solutions exist as pervanadyl VO$_2^+$ or decavanadate H$_2$V$_{10}$O$_{28}^{4-}$ ions (Figure 6b). Thus, the former is probably responsible for certain extent of corrosion inhibition at pH = 1.3 whereas the latter only accelerated intermetallic corrosion at pH = 2.5.

In fact, corrosion inhibition when $c_V \geq 100$ mM was related to precipitation of vanadium-containing film on the surface of the electrode. EDX analysis indicated that Al, Cu, V and O were uniformly distributed on the surface and no severe corrosion loss was observed (Figure 7). Precipitation occurred also within bulk of the solution. According to Figure 6b, V$_2$O$_5$ should precipitate from the solution at pH = 1.3, when $c_V \geq 130$ mM. This however is only an approximate distribution diagram. Positions of demarcation lines in the diagram depend on the ionic strength of the solution, the counterion as well as temperature. Therefore, precipitation of V$_2$O$_5$ from solution containing 100 mM of vanadium cannot be excluded.

Further examination of precipitate in the form of powder and thin film on the surface of the electrode was performed using X-ray photoelectron spectroscopy (XPS). Major components of the powder are oxygen and vanadium. Oxygen was mainly bound to vanadium with small admixture of sulfate (data not shown). Much more interesting is V 2p$^{3/2}$ spectrum (Figure 8a). It shows two forms of vanadium at 516.8 eV and 518.0 eV, corresponding to V(IV) and V(V) states, respectively [29]. Therefore, the powder is composed of V$_2$O$_5$ with a small admixture of adsorbed VO$^{2+}$. Data presented in Table 1 confirms this assumption i.e., the sum of concentrations of oxygen in vanadium pentoxide and vanadyl cation, calculated from the vanadium concentrations, equals to total oxide oxygen concentration.
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Figure 7. Micrograph of intermetallic Al$_2$Cu after corrosion in H$_2$SO$_4$ solution, $c_V = 100$ mM, $pH = 1.3$, immersion time $t = 576$ min, $T = 303$ K together with spatial distribution of elements, color intensity differences on EDX maps occur due to different concentration of elements and are also related to height of the specimen.

Further examination of precipitate in the form of powder and thin film on the surface of the electrode was performed using X-ray photoelectron spectroscopy (XPS). Major components of the powder are oxygen and vanadium. Oxygen was mainly bound to vanadium with small admixture of sulfate (data not shown). Much more interesting is V 2p 3/2 spectrum (Figure 8a). It shows two forms of vanadium at 516.8 eV and 518.0 eV, corresponding to V(IV) and V(V) states, respectively [29]. Therefore, the powder is composed of $V_2O_5$ with a small admixture of adsorbed VO$_2^+$. Data presented in Table 1 confirms this assumption i.e., the sum of concentrations of oxygen in vanadium pentoxide and vanadyl cation, calculated from the vanadium concentrations, equals to total oxide oxygen concentration.

Figure 8. V 2p 3/2 fitted peaks: (a) for powder precipitated from the solution; (b) for vanadium-containing film deposited onto electrode.

| Measurement | O 1s | V 2p | Al 2s | Cu 2p |
|-------------|------|------|-------|-------|
| O$_2^-$     | 70.0 | 18.0 | 9.0   | 2.0   |
| V$^{5+}$    | 70.0 | 18.0 | 9.0   | 2.0   |
| V$^{4+}$    | 70.0 | 18.0 | 9.0   | 2.0   |

Average 70.0 18.0 9.0 2.0

Figure 8. V 2p 3/2 fitted peaks: (a) for powder precipitated from the solution; (b) for vanadium-containing film deposited onto electrode.
Table 1. The contribution of fitted peaks for powder sample (at.%).

| Measurement | O 1s | V 2p |
|-------------|------|------|
|              | O2-  | V5+  | V4+  |
| 1            | 69.8 | 26.3 | 3.9  |
| 2            | 68.5 | 26.3 | 5.2  |
| Average      | 69.2 | 26.3 | 4.6  |

The spectrum obtained for vanadium-containing film on the electrode surface, besides aforementioned O 1s and V 2p\(^{3/2}\) peaks consists also of Al 2s and Cu 2p peaks at 120 and 933 eV, respectively. Al 2s peak overlaps with Cu 3s which occurs around 123 eV. Taking it into consideration Cu 3s peak was eliminated from the Al 2s spectra. Oxygen again is mainly bound to vanadium, with small admixture of sulfate. Vanadium in the film is mainly in the form of V\(^{2+}\)O\(_5\), but the contribution of VO\(^{2+}\) is higher when compared to the powder (Figure 8a, Table 2). It is reasonable because reduction of VO\(^{2+}\) to VO\(^{2+}\) occurs at the electrode surface. Al 2s peak is mainly related to Al\(^{3+}\) formed during Al\(_2\)Cu corrosion and adsorbed onto the vanadium-containing film. In the case of copper, it is difficult to determine whether it is in ionic or metallic form. Both possibilities are plausible. Cu is present in the solution due to corrosion process. Its concentration in the solution is ca. 4 times lower when compared to aluminum (Figure 3a). Small contribution from the substrate i.e., Al\(_2\)Cu also cannot be excluded. Big discrepancy between Cu content obtained in two different areas on the specimen seems to justify this speculation. Additional general conclusion is that the chemical composition of the vanadium-containing film is less homogeneous when compared to the powder.

Table 2. The contribution of fitted peaks for vanadium-containing film on the Al\(_2\)Cu (at.%).

| Measurement | O 1s | V 2p | Al 2s | Cu 2p |
|-------------|------|------|-------|-------|
|              | O2-  | V5+  | V4+  | Al\(^{3+}\)| Cu\(^{2+}/Cu\) |
| 1            | 54.8 | 16.0 | 9.1   | 19.0  | 1.2   |
| 2            | 61.9 | 15.7 | 6.9   | 15.3  | 0.3   |
| Average      | 58.3 | 15.8 | 8.0   | 17.1  | 0.7   |

3.3. Electrochemical Analysis—Influence of Vanadium Species

OCP of the intermetallic Al\(_2\)Cu was studied as a function of time and initial concentration of Na\(_3\)VO\(_4\) in the solutions. When their pH was 1.3 and c\(_V\) = 0 and 10 mM OCP values initially dropped and then slowly increased with time (Figure 9a). When vanadium initial concentration was equal to 50 mM OCP decreased slowly with time. OCP values after \(t = 576\) min, for solutions with c\(_V\) ≤ 50 mM at pH = 1.3, were between −99 and −185 mV vs. Ag/AgCl (3 M KCl) electrode. On the contrary, when c\(_V\) = 100 and 200 mM, OCP increased as the time increased and reached 340–360 mV vs. Ag/AgCl (3 M KCl). It should be noted here that equilibrium potential of hydrogen electrode is −287 mV vs. Ag/AgCl (3 M KCl). Thus, one of the possible cathodic process is oxygen reduction. Equilibrium potential of oxygen electrode is 940 mV vs. Ag/AgCl (3 M KCl). Another is reduction of V\(^V\) to V\(^{IV}\) with the equilibrium potential 790 mV vs. Ag/AgCl (3 M KCl) [30]. The latter was confirmed using XPS. Further reduction of tetravalent to trivalent vanadium species also is possible when c\(_V\) ≤ 100 mM, because the equilibrium potential is equal to 127 mV vs. Ag/AgCl (3M KCl) [30]. V\(^{III}\) species however were not detected neither in the powder, nor within the vanadium-containing film. When pH = 2.5 OCP values decreased with time (Figure 9b). Interestingly, when c\(_V\) = 50 and 200 mM, plateaus on the OCP vs. time curves were observed. This may indicate temporary effect of corrosion inhibition. OCP values after 576 min for c\(_V\) = 0, 10, 50, and 200 mM were between −150 and −300 vs. Ag/AgCl (3 M KCl). This is slightly lower than pH = 1.3. Selective corrosion in the solutions
with pH = 2.5 was less severe when compared to pH = 1.3. Thus, electrodes contained less copper on the surface. This caused lower values of OCP. Relatively high value of OCP was achieved when \( c_V = 100 \) mM. This, however, does not mean lower corrosion rate (Figure 5). At pH = 2.5 the same cathodic processes can occur as mentioned above. Reduction of \( V^{IV} \) to \( V^{III} \) is possible when \( c_V = 10, 50, \) and 200 mM.

![Impedance spectra of intermetallic Al2Cu for pH = 1.3, t = 576 min were analyzed as a function of sodium orthovanadate initial concentration. This pH was selected because certain degree of corrosion inhibition was achieved on the contrary to pH = 2.5 (Figure 5). The characteristic features of the spectra obtained when \( c_V \leq 50 \) mM are linear parts at high frequency, \( f \geq 23.8 \) Hz (0 mM), 12.7 Hz (10 mM) and 8.3 Hz (50 mM, Figure 10a,b). They have an angle between 20 and 42° with respect to the real axis. This is related to the porous layer formed on the surface due to selective corrosion (Figures 3 and 4). The impedance response of the porous electrode was initially introduced by de Levie [31]. According to his model, the Nyquist plot calculated for a single, cylindrical, pore, in the absence of a faradaic current consisted of a straight line with 45° angle at high frequencies and a vertical line at low frequencies. In the real systems, however, it is necessary to include different shape of the pores, size of their openings, possible branching, contribution from the flat part of the electrode on its top and ohmic drop within the solution and electrode material. These factors cause deviation from the theoretical angle or even from the linear shape of the high-frequency part. Due to the charge-transfer processes, semicircle or depressed semicircle appears at lower frequencies instead of the horizontal line [32–34]. The high-frequency parts of the spectra are related to geometry of the porous electrode and do not contain valuable information regarding corrosion kinetics. Thus, they were not fitted. Distorted semi-circles observed at medium frequencies (23.8–0.4 Hz for 0 mM, 12.7–0.4 Hz for 10 mM and 8.3–0.4 Hz for 50 mM) correspond to the double-layer capacitance \( C_{dl} \) in parallel to the charge-transfer resistance \( R_{ct} \). Due to frequency dispersion, constant phase element CPE was used for fitting instead of the capacitor [32]. Additional one or more time constants can be observed at lower frequencies. They include inductive and capacitive loops and are related to adsorption processes. Scattered impedance values observed at the lowest frequencies were excluded from fitting.

Impedance spectra obtained for \( c_V = 100 \) and 200 mM are different when compared to the other ones. First, linear, high-frequency parts indicating electrode porosity do not exist. Secondly, due to precipitation of vanadium-containing film, much higher impedances were observed (Figure 10c).

When pH = 2.5, \( t = 159 \) min, \( c_V = 0 \) and 200 mM also high impedances were obtained (Figure 10d). This confirms temporary corrosion inhibition deduced from OCP vs. time curve (Figure 9b).
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Impedance spectra obtained for $c_V = 100$ and 200 mM are different when compared to the other ones. First, linear, high-frequency parts indicating electrode porosity do not exist. Secondly, due to precipitation of vanadium-containing film, much higher impedances were obtained (Figure 10c). When $pH = 2.5$, $t = 159$ min, $c_V = 0$ and 200 mM also high impedances were obtained (Figure 10d). This confirms temporary corrosion inhibition deduced from OCP vs. time curve (Figure 9b).

Figure 10. Impedance spectra of intermetallic Al$_2$Cu as a function of initial concentration of Na$_3$VO$_4$ in the solutions: (a) 0 mM, pH = 1.3, $t = 576$ min; (b) 10 and 50 mM, pH = 1.3, $t = 576$ min; (c) 100 and 200 mM, pH = 1.3, $t = 576$ min; (d) 0 and 200 mM, pH = 2.5, $t = 159$ min; $T = 303$ K, continuous lines indicate fitted parts of the spectra.

Impedance spectra were fitted using EECs presented in Figure 11. Faradaic impedance of the first one (Figure 11a) consists of $R_{ct}$, $R_1$, and $L_1$. This model was initially introduced by Epelboin et al. for passivation of metals. The resistance $R_1$ was related to changes of the surface coverage or film resistivity with time. The inductance $L_1$ was inversely proportional to the time constant of this process \cite{35}. In this work, connection of $R_1$ and $L_1$ in series is related to adsorption of species formed during the anodic dissolution of the intermetallic Al$_2$Cu. These are probably aluminum ions. It should be noted here that detection of different intermediates adsorbed during aluminum corrosion in acids is impossible using EIS \cite{36}. However, in other environments, such as aqueous alkaline and non-aqueous solutions, three step mechanisms of Al dissolution were successfully investigated using EIS \cite{36,37}.

The alternative EEC, where passive behavior is modelled using capacitor in parallel to resistor, was also proposed \cite{35}. This model was used in this work for fitting the spectra obtained for $c_V = 100$ and 200 mM at pH = 1.3 and $c_V = 200$ mM at pH = 2.5, where inductive behavior was not observed (Figure 11b). However, constant phase element CPE$_1$ was used instead of the capacitor. $R_1$ and CPE$_1$ depend both on the charge-transfer resistance and faradaic current change caused by increase of the surface coverage, presumably with vanadium-containing film \cite{35}.

When $c_V = 10$ and 50 mM more complicated equivalent circuit, with additional time constant, was used (Figure 11c). Cao derived general equation of faradaic impedance for electrode processes whose rate depend on electrode potential and two other, potential-dependent, variables such as surface
coverage [38]. He also proposed several equivalent circuits applicable for such a case. In this work, these two potential-dependent variables are related to adsorption of two different species, presumably aluminum ions and vanadium isopolyoxoanions. Physical interpretation of inductor \( L_1 \), capacitor \( C_2 \), and resistors \( R_1 \) and \( R_2 \), applied in this model, is not straightforward. In general, these parameters are mutually related; e.g., \( C_2 \) depends on \( R_1 \), \( R_2 \), \( R_{ct} \), and \( L_1 \). They depend on changes in faradaic current caused by changes in surface coverage and changes in adsorption kinetics caused by electrode potential. In addition, all these parameters depend on both adsorbing species [38].

\[ R_{ct} = \text{charge transfer resistance}, \ C_{PE} \text{ is the constant phase element corresponding to double layer capacitance}, \ R_s \text{ is the solution resistance,} \ L_1, L_2 \text{ are resistors and capacitors related to adsorption and precipitation of insoluble film: (a) case of selective corrosion, when } c_V = 0 \text{ mM, pH} = 1.3, \ t \geq 437 \text{ min, } c_V = 0 \text{ mM, pH} = 2.5; \text{(b) case of corrosion inhibition, when when } c_V = 100 \text{ and } 200 \text{ mM, pH} = 1.3, c_V = 200 \text{ mM, pH} = 2.5, t = 159 \text{ min; (c) case of selective corrosion, when } c_V = 0 \text{ mM, pH} = 1.3, t < 437 \text{ min, } c_V = 10 \text{ and } 50 \text{ mM.}

The results of the impedance spectra fitting are given in Tables 3–5.

In general, to estimate the corrosion rate from the impedance spectra one should determine polarization resistance \( R_p \) value i.e., the resistance when \( f \rightarrow 0 \). Unfortunately, resistances obtained at low frequencies are scattered and reliable values of \( R_p \) cannot be obtained. Approximation of the impedance spectra allowed, however, determination of the charge-transfer resistance. It decreases as \( c_V \) increases from 0 to 50 mM (Figure 12). This indicates that at pH = 1.3 vanadium species, presumably peryanadyl ions, act as additional depolarizers in the corrosion cell. In fact, reduction of peryanadyl to vanadyl was even confirmed for \( c_V = 100 \) mM using XPS. From Figure 5 it can be concluded that certain corrosion inhibition is expected at \( c_V = 50 \) mM. It should be noted here, however, that corrosion rates presented in Figure 5 are averaged over \( t = 576 \) min whereas impedance spectrum gives information at the moment when it is recorded. In addition, \( R_{ct} \) values are normalized to geometric surface area of the specimen but not the real one. The latter remains unknown but possibly very high due to intensive selective corrosion [27]. The increase of the real surface area also causes the decrease of the resistance.

When \( c_V \geq 100 \) mM, \( R_{ct} \) achieves much higher values. Vanadium-containing film precipitating on the surface of the intermetallic, hampers the charge transfer through the solid/liquid interphase.

\[ R_{ct} (\Omega \cdot \text{cm}^2) \]

\[ c_V (\text{mM}) \]

Figure 12. Charge-transfer resistance as a function of initial concentration of sodium orthovanadate in H\(_2\)SO\(_4\) solution, pH = 1.3, t = 576 min, T = 303 K.
Table 3. The results of fitting of the impedance spectra obtained at pH = 1.3 using EEC from Figure 11a,c: $\chi^2$ and $S$ indicate the quality of the fit (chi-square and residual sum of squares, respectively); $R_s$ is the solution resistance; $T_{dl}$ is the parameter of the constant phase element CPE$_{dl}$ representing the electrical double layer; $\alpha_{dl}$ is the exponent of the CPE; $R_{ct}$ is the charge-transfer resistance; $L_1$, $R_1$, $R_2$, and $C_2$ are the inductance, resistances, and capacitance related to the adsorption processes, respectively, uncertainties of the fitted parameters are given in brackets.

| Experimental Conditions | $\chi^2$ | $S$ | $R_s$ (Ω cm$^2$) | $T_{dl}$ (mF s$^{0.5}$ cm$^{-2}$) | $\alpha_{dl}$ | $R_{ct}$ (Ω cm$^2$) | $R_1$ (Ω cm$^2$) | $L_1$ (H cm$^2$) | $C_2$ (mF cm$^{-2}$) | $R_2$ (Ω cm$^2$) |
|-------------------------|---------|-----|-----------------|-----------------|------------|-----------------|-----------------|----------------|-----------------|----------------|
| $cV = 0$ mM, $t = 159$ min | $1.05 \times 10^{-4}$ | 0.11 | 13 (1) | 2.055 (0.022) | 0.95 (0.01) | 97 (1) | 0 | 211 (16) | 17.805 (1.368) | 91 (2) |
| $cV = 0$ mM, $t = 298$ min | $5.46 \times 10^{-4}$ | 0.04 | 15 (1) | 4.201 (0.067) | 0.96 (0.01) | 56 (1) | 0 | 95 (6) | 28.958 (2.534) | 44 (1) |
| $cV = 0$ mM, $t = 437$ min | $2.23 \times 10^{-4}$ | 0.01 | 17 (1) | 7.804 (0.883) | 0.91 (0.01) | 46 (1) | 61 (1) | 121 (4) |
| $cV = 0$ mM, $t = 576$ min | $1.59 \times 10^{-4}$ | 0.01 | 18 (1) | 12.856 (0.168) | 0.86 (0.01) | 57 (1) | 36 (1) | 65 (2) |
| $cV = 10$ mM, $t = 576$ min | $3.76 \times 10^{-4}$ | 0.02 | 11 (1) | 35.294 (1.202) | 0.85 (0.02) | 18 (1) | 0 | 17 (1) | 99.884 (6.842) | 16 (1) |
| $cV = 50$ mM, $t = 576$ min | $2.59 \times 10^{-4}$ | 0.01 | 11 (1) | 66.191 (2.919) | 0.60 (0.03) | 15 (1) | 0 | 23 (1) | 201.722 (17.043) | 17 (1) |

Table 4. The results of fitting of the impedance spectra obtained when corrosion was inhibited, using EEC from Figure 11b: $\chi^2$ and $S$ indicate the quality of the fit (chi-square and residual sum of squares, respectively); $R_s$ is the solution resistance; $T_{dl}$ is the parameter of the constant phase element CPE$_{dl}$ representing the electrical double layer; $\alpha_{dl}$ is the exponent of the CPE; $R_{ct}$ is the charge-transfer resistance; $R_1$, $T_1$, and $\alpha_1$ are the resistance and constant phase element CPE$_1$ parameters related to the adsorption/formation of a protective layer processes, respectively, uncertainties of the fitted parameters are given in brackets.

| Experimental Conditions | $\chi^2$ | $S$ | $R_s$ (Ω cm$^2$) | $T_{dl}$ (mF s$^{0.5}$ cm$^{-2}$) | $\alpha_{dl}$ | $R_{ct}$ (Ω cm$^2$) | $R_1$ (Ω cm$^2$) | $T_1$ (mF s$^{0.5}$ cm$^{-2}$) | $\alpha_1$ |
|-------------------------|---------|-----|-----------------|-----------------|------------|-----------------|-----------------|-----------------|---------|
| pH = 1.3 | $cV = 100$ mM, $t = 576$ min | $2.68 \times 10^{-4}$ | 0.03 | 7 (1) | 0.043 (0.001) | 0.88 (0.01) | 1108 (29) | 379 (33) | 0.660 (0.083) | 0.64 (0.04) |
| pH = 2.5 | $cV = 200$ mM, $t = 576$ min | $2.42 \times 10^{-4}$ | 0.03 | 5 (1) | 0.044 (0.001) | 0.88 (0.01) | 919 (47) | 521 (52) | 0.437 (0.066) | 0.54 (0.03) |
| pH = 2.5 | $cV = 200$ mM, $t = 159$ min | $2.91 \times 10^{-4}$ | 0.04 | 11 (1) | 0.048 (0.001) | 0.87 (0.01) | 1450 (11) | 581 (21) | 2.205 (0.117) | 0.68 (0.02) |
Table 5. The results of fitting of the impedance spectra obtained at pH = 2.5 using EEC from Figure 11a: $\chi^2$ and $S$ indicate the quality of the fit (chi-square and residual sum of squares, respectively); $R_s$ is the solution resistance; $T_{dl}$ is the constant phase element CPE$_{dl}$ parameter representing the electrical double layer; $\alpha_{dl}$ is the exponent of the CPE; $R_{ct}$ is the charge-transfer resistance; $L_1$ and $R_1$ are the inductance and resistance related to the adsorption processes, respectively, uncertainties of the fitted parameters are given in brackets.

| Experimental Conditions | $\chi^2$ | $S$ | $R_s$ (Ω·cm$^2$) | $T_{dl}$ (mF·s$^{\alpha_{dl}}$·cm$^{-2}$) | $\alpha_{dl}$ | $R_{ct}$ (Ω·cm$^2$) | $R_1$ (Ω·cm$^2$) | $L_1$ (H·cm$^2$) |
|-------------------------|---------|-----|-------------------|---------------------------------|-----------|-------------------|-----------------|--------------|
| $cV = 0$ mM, $t = 159$ min | $7.13 \times 10^{-4}$ | 0.07 | 133 (1) | 0.557 (0.009) | 0.64 (0.01) | 1325 (33) | 249 (6) | 772 (11) |
| $cV = 0$ mM, $t = 298$ min | $4.49 \times 10^{-4}$ | 0.04 | 80 (1) | 0.587 (0.008) | 0.68 (0.01) | 766 (11) | 150 (3) | 398 (4) |
| $cV = 0$ mM, $t = 437$ min | $8.39 \times 10^{-4}$ | 0.08 | 65 (1) | 0.722 (0.011) | 0.70 (0.01) | 657 (12) | 133 (2) | 368 (4) |
| $cV = 0$ mM, $t = 576$ min | $1.97 \times 10^{-4}$ | 0.01 | 61 (1) | 0.966 (0.012) | 0.71 (0.01) | 658 (12) | 138 (2) | 356 (3) |
The increase of the surface area of the electrode caused by selective corrosion can be easily monitored using the double-layer capacitance. The capacitive behavior of the electrical double layer is usually modelled using CPE instead of capacitor due to frequency dispersion. The impedance of the CPE is given by Equation (4) [32]:

$$Z_{CPE} = \frac{1}{T_{dl}(j\omega)^{\alpha}},$$

where $T_{dl}$ is equal to capacitance when $\alpha = 1$, $\omega$ is the angular frequency and $j$ is imaginary number.

Provided that time constants for the faradaic process are distributed along the surface of the electrode, the double-layer capacitance can be obtained as follows (Equation (5)) [32]:

$$C_{dl} = T_{dl}^{\alpha} \left( \frac{1}{R_s} + \frac{1}{R_{ct}} \right)^{1-\frac{\alpha}{2}}$$

This equation works well for relatively high $\alpha$ values, but when $\alpha < 0.85$, calculation of $C_{dl}$ can lead to significant error [39]. Thus, double-layer capacitance was estimated only for $c_V = 0$ mM at $pH = 1.3$. After $t = 576$ min it is equal to 9.632 (0.482) mF·cm$^{-2}$. Again, it should be noted that capacitance was normalized to geometric surface area of the specimen. The value obtained is 482 times higher when compared to theoretical value 20 $\mu$F·cm$^{-2}$ [34]. This is characteristic for highly porous electrodes and is related to their enormous surface area [12,27,33,34].

It should be noted here that resistances $R_1$ obtained from impedance spectra approximation were very low. Thus, in Table 3 0 $\Omega$ values are reported. This is probably caused by a very small diameter of the closed loops e.g., in Figure 10b. Inductances gradually decrease as the surface area of the electrode increases.

Polarization resistances $R_p$ were estimated using polarization curves recorded in the vicinity of the corrosion potential at $pH = 1.3$ (data not shown). They were approximated using Equation (3). It can be concluded that $R_p = 32, 16, 15, 912$ and 1146 $\Omega$·cm$^2$ for $c_V = 0, 10, 50, 100, $ and 200 mM. Again, the discrepancy between the corrosion rate and polarization resistance exists for $c_V = 50$ mM. The possible explanation was given during discussion of $R_{ct}$ values. These $R_p$ values are normalized to geometric surface area of the electrode i.e., they decrease as the surface area increases. The values of Tafel constant were in most cases obtained with very high uncertainties. Furthermore, their practical importance is in this case insignificant, because they characterize all cathodic and anodic processes occurring at the electrode. These two obstacles make calculation of the corrosion current density from polarization resistance impossible.

3.4. Electrochemical Analysis—The Kinetics of Selective Corrosion

The kinetics of corrosion was studied in the solutions without vanadium species at $pH = 1.3$ and 2.5. The impedance of the intermetallic decreases as the immersion time increases. It is also much lower at $pH = 1.3$ than at $pH = 2.5$ (Figure 13). The results of fitting of these spectra are given in Table 5. Intermetallic Al$_2$Cu corrodes selectively in acidic solutions. This process is more severe at $pH = 1.3$ than 2.5 (Figure 3). In the former case double-layer capacitance was calculated using Equation (5). Obtained values are high and increase with time due to increase of the surface area of the electrode (Figure 14a). In the latter case $C_{dl}$ was not calculated because $\alpha_{dl} < 0.85$. Charge-transfer resistance, in turn, was determined for both cases (Figure 14b). It decreases with time as already explained. Approximately 10-fold lower values were obtained for $pH = 1.3$.

It can be observed that the product $C_{dl} R_{ct}$ should not depend on the immersion time of the intermetallic. Interestingly, this is not valid at $pH = 1.3$ (Figure 15). It means that $C_{dl}$ and $R_{ct}$ changes with time are caused not only by the increases of electrode surface area but also by the change in mechanism. In fact, it is quite reasonable because the chemical composition of the electrode changes with time i.e., aluminum content on the surface decreases as the immersion time increases. The chemical composition of the solution also changes. Its pH raises in the vicinity of the electrode due to oxygen
reduction and aluminum concentration increases. The surface of the electrode becomes highly porous and enriched with copper. Copper corrodes in aerated acidic solution when is not electrically coupled with aluminum. At the same time, porous copper structure can locally detach from the surface and dissolve in the solution. Thus, copper ions may easily act as the additional depolarizers in the corrosion cell. These two factors i.e., changes of chemical composition of the electrode and solution probably affect the double-layer capacitance and charge-transfer kinetics.

![Figure 13](image1)

**Figure 13.** Impedance spectra of intermetallic Al<sub>2</sub>Cu as a function of immersion time: (a) pH = 1.3; (b) pH = 2.5; T = 303 K, c<sub>V</sub> = 0 mM.

![Figure 14](image2)

**Figure 14.** Kinetics of selective corrosion of intermetallic Al<sub>2</sub>Cu: (a) double-layer capacitance at pH = 1.3; (b) charge-transfer resistance at pH = 1.3 and = 2.5; c<sub>V</sub> = 0 mM, T = 303 K.

![Figure 15](image3)

**Figure 15.** C<sub>dl</sub>·R<sub>ct</sub> product as a function of immersion time, pH = 1.3, c<sub>V</sub> = 0 mM, T = 303 K.
4. Conclusions

Pentavalent vanadium species were tested as corrosion inhibitors of intermetallic Al\textsubscript{2}Cu in H\textsubscript{2}SO\textsubscript{4} at pH = 1.3 and 2.5, respectively. When pH = 1.3 and \(c_V\) was < 100 mM severe dealloying of the Al\textsubscript{2}Cu phase occurred. The inhibition effect, observed when \(c_V\) was ≥ 100 mM, was mainly related to precipitation of vanadium-containing film on the surface of the electrode and in the bulk of the solution. XPS spectra analysis showed that V\textsubscript{2}O\textsubscript{5} is the main component of the film which agrees with thermodynamic considerations. Precipitation of vanadium pentoxide within the bulk of the solution makes application of vanadium species, possibly pervanadyl ions, as corrosion inhibitors of Al\textsubscript{2}Cu at pH = 1.3 impractical. However, their influence on the corrosion behavior of an Al-Cu solid solution and other intermetallics, existing in aluminum alloys, in H\textsubscript{2}SO\textsubscript{4} should also be studied. When pH = 2.5 and vanadium was probably in the form of decavanadate ions, corrosion of the intermetallic was not inhibited. On the contrary, it was accelerated due to introduction of additional depolarizer to the corrosion cell. At the same time, corrosion rate was lower when compared to pH = 1.3. The corrosion mechanism of Al\textsubscript{2}Cu is complex. Several cathodic processes are possible in the studied system. In addition, surface area of the electrode changes with time which complicates the comparison between the impedence spectra and impede calculation of corrosion current density.

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References

1. Belov, N.A.; Eskin, D.G.; Aksenov, A.A. Multicomponent Phase Diagrams: Applications for Commercial Aluminum Alloys; Elsevier: Oxford, UK, 2005; pp. 152–192.
2. Scully, J.R.; Knight, T.O.; Buchheit, R.G.; Peebles, D.E. Electrochemical characteristics of the Al\textsubscript{2}Cu, Al\textsubscript{3}Ta and Al\textsubscript{3}Zr intermetallic phases and their relevancy to the localized corrosion of Al alloys. Corros. Sci. 1993, 35, 185–195. [CrossRef]
3. Buchheit, R.G. A compilation of corrosion potentials reported for intermetallic phases in aluminum alloys. J. Electrochem. Soc. 1995, 142, 3994–3996. [CrossRef]
4. Buchheit, R.G.; Martinez, M.A.; Montes, L.P. Evidence for Cu Ion Formation by Dissolution and Dealloying the Al\textsubscript{2}CuMg Intermetallic Compound in Rotating Ring-Disk Collection Experiments. J. Electrochem. Soc. 2000, 147, 119–124. [CrossRef]
5. Vukmirovic, M.B.; Dimitrov, N.; Sieradzki, K. Dealloying and Corrosion of Al Alloy 2024-T3. J. Electrochem. Soc. 2002, 149, B428. [CrossRef]
6. Zhang, X.; Hashimoto, T.; Lindsay, J.; Zhou, X. Investigation of the de-alloying behaviour of \(\theta\)-phase (Al\textsubscript{2}Cu) in AA2024-T351 aluminum alloy. Corros. Sci. 2016, 108, 85–93. [CrossRef]
7. Osório, W.R.; Spinelli, J.E.; Ferreira, I.L.; Garcia, A. The roles of macrosegregation and of dendritic array spacings on the electrochemical behavior of an Al–4.5 wt% Cu alloy. Electrochim. Acta 2007, 52, 3265–3273. [CrossRef]
8. Osório, W.R.; Spinelli, J.E.; Freire, C.M.A.; Cardona, M.B.; Garcia, A. The roles of Al\textsubscript{2}Cu and of dendritic refinement on surface corrosion resistance of hypoeutectic Al-Cu alloys immersed in H\textsubscript{2}SO\textsubscript{4}. J. Alloys Compd. 2007, 443, 87–93. [CrossRef]
9. Osório, W.R.; Siqueira, C.A.; Santos, C.A.; Garcia, A. The Correlation between Electrochemical Corrosion Resistance and Mechanical Strength of As-Cast Al-Cu and Al-Si Alloys. Int. J. Electrochem. Sci. 2011, 6, 6273–6289.
10. Obispo, H.M.; Murr, L.E.; Arrowood, R.M.; Trillo, E.A. Copper deposition during the corrosion of aluminum alloy 2024 in sodium chloride solutions. *J. Mater. Sci.* 2001, 36, 3479–3495.

11. Lebouli, S.; Tardelli, I.; Rocca, E.; Volovitch, P.; Ogle, K. Dealloying of Al$_2$Cu, Al$_2$Cu$_2$Fe, and Al$_2$CuMg intermetallic phases to form nanoparticulate copper films. *Mater. Corros.* 2014, 65, 416–424. [CrossRef]

12. Kwolek, P.; Dychton, K.; Pytel, M. Orthophosphoric acid solutions of sodium orthovanadate, sodium tungstate, and sodium molybdate as potential corrosion inhibitors of the Al$_2$Cu intermetallic phase. *J. Solid State Electrochem.* 2019, 23, 3019–3029. [CrossRef]

13. ASTM B137-95. *Standard Test Method for Measurement of Coating Mass Per Unit Area on Anodically Coated Aluminum*; ASTM International: West Conshohocken, PA, USA, 2009.

14. Katz, S.A.; Salem, H. The toxicology of chromium with respect to its chemical speciation: A review. *J. Appl. Toxicol.* 1993, 13, 217–224. [CrossRef] [PubMed]

15. Li, X.; Deng, S.; Fu, H. Sodium molybdate as a corrosion inhibitor for aluminium in H$_3$PO$_4$ solution. *Corros. Sci.* 2011, 53, 2748–2753. [CrossRef]

16. Kwolek, P.; Kaminski, A.; Dychton, K.; Drajewicz, M.; Sieniawski, J. The corrosion rate of aluminium in the orthophosphoric acid solutions in the presence of sodium molybdate. *Corros. Sci.* 2016, 106, 208–216. [CrossRef]

17. Dychton, K.; Kwolek, P. The replacement of chromate by molybdate in phosphoric acid-based etch solutions for aluminium alloys. *Corros. Eng. Sci. Technol.* 2018, 53, 234–240. [CrossRef]

18. Kwolek, P.; Pustula, A.; Nowak, W.J. Influence of molybdocophosphoric acid on the kinetics of the anodic coating dissolution. *Surf. Coat. Technol.* 2019, 357, 535–542. [CrossRef]

19. Pope, M.T. *Heteropoly and Isopoly Oxometalates*; Springer: Berlin, Germany, 1983; pp. 1–3, 34–40.

20. Kwolek, P. Effect of Na$_3$VO$_4$ inhibitor on the corrosion resistance of Al$_2$Cu intermetallic phase in H$_3$PO$_4$ aqueous solution. *Arch. Metall. Mater.* 2020, 65, 175–183.

21. Ralston, K.D.; Chrisianti, S.; Young, T.L.; Buchheit, R.G. Corrosion Inhibition of Aluminum Alloy 2024-T3 by Aqueous Vanadium Species. *J. Electrochem. Soc.* 2008, 155, C350–C359. [CrossRef]

22. Mansfeld, F. The Polarization Resistance Technique for Measuring Corrosion Currents. In *Advances in Corrosion Science and Technology*; Fontana, M.G., Staehle, R., Eds.; Plenum Press: Boston, MA, USA, 1970; pp. 163–262.

23. Iannuzzi, M.; Frankel, G.S. Mechanisms of corrosion inhibition of AA2024-T3 by vanadates. *Corros. Sci.* 2007, 49, 2371–2391. [CrossRef]

24. Boukamp, B. A linear Kronig-Kramers transform test for immittance data validation. *J. Electrochem. Soc.* 1995, 142, 1885–1894. [CrossRef]

25. Boukamp, B. Electrochemical impedance spectroscopy in solid state ionics: Recent advances. *Solid State Ionics* 2004, 169, 65–73. [CrossRef]

26. Mansfeld, F. Influence of the electrochemical properties of vanadium oxides on specific capacitance by molybdenum doping. *Bull. Mater. Sci.* 2019, 42, 37. [CrossRef]

27. Zobac, O.; Kroupa, A.; Zemanova, A.; Richter, K.W. Experimental Description of the Al-Cu Binary Phase Diagram. *Mater. Trans. A Phys. Metall. Mater. Sci.* 2019, 50, 3805–3815. [CrossRef]

28. Huang, Y.; Zhang, Y. Influence of the electrochemical properties of vanadium oxides on specific capacitance by molybdenum doping. *Bull. Mater. Sci.* 2019, 42, 37. [CrossRef]

29. De Levie, R. Electrochemical response of porous and rough electrodes. In *Advances in Electrochemistry and Electrochemical Engineering*; Delahay, P., Ed.; Interscience: New York, NY, USA, 1967; pp. 329–397.

30. Lasia, A. *Electrochemical Impedance Spectroscopy and Its Applications*; Springer: New York, NY, USA, 2014; pp. 177–196, 210–214.

31. Chen, L.; Lasia, A. Ni-Al Powder Electro catalyst for Hydrogen Evolution. *J. Electrochem. Soc.* 1993, 140, 2464–2473. [CrossRef]

32. Chen, L.; Lasia, A. Study of the kinetics of hydrogen evolution reaction on Nickel-Zinc powder electrodes. *J. Electrochem. Soc.* 1992, 139, 3214–3219. [CrossRef]
35. Epelboin, I.; Gabrielli, C.; Keddam, M.; Takenouti, H. The Study of the Passivation Process by the Electrode Impedance Analysis. In Electrochemical Materials Science. Comprehensive Treatise of Electrochemistry; Bockris, J.O., Conway, B.E., Yeager, E., White, R.E., Eds.; Springer: Boston, MA, USA, 1981; pp. 151–192.

36. Péter, L.; Arai, J.; Akahoshi, H. Impedance of a reaction involving two adsorbed intermediates: Aluminum dissolution in non-aqueous lithium imide solutions. J. Electroanal. Chem. 2000, 482, 125–138. [CrossRef]

37. Macdonald, D.D.; Lee, K.H.; Mocari, A.; Harrington, D. Evaluation of Alloy Anodes for Aluminum-Air Batteries: Corrosion Studies. Corrosion 1988, 44, 652–657. [CrossRef]

38. Cao, C. On the impedance plane displays for irreversible electrode reactions based on the stability conditions of the steady state II. Two state variables besides electrode potential. Electrochim. Acta 1990, 35, 837–844. [CrossRef]

39. Łosiewicz, B.; Jurczakowski, R.; Lasia, A. Kinetics of hydrogen underpotential deposition at iridium in sulfuric and perchloric acids. Electrochim. Acta 2017, 225, 160–167. [CrossRef]

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