Effect of Electric Pulse Treatment on the Corrosion Resistance of HAl52-2 Aluminum Brass

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In this work, the effects of electric pulse treatment (EPT) on the corrosion resistance of HAl52-2 aluminum brass has been investigated using electrochemical workstation, scanning electron microscope and X-ray diffraction. The results have shown that after EPT, the corrosion potential increased and the corrosion current density decreased. At the same time, the diameter of the capacitive reactance arc of the HAl52-2 aluminum brass after EPT was 5 ~ 6 times of the original sample, and the thickness of dezincification layer was reduced from 59.2 µm to 50.2 µm by 84.8%. The aluminum brass alloy with EPT has demonstrated better corrosion resistance of than the original sample.

Keywords: corrosion behavior, electric pulse treatment, γ phase, HAl52-2 aluminum brass.

1. Experiment Materials and Methods

HAl52-2 aluminum brass (Cu: 52wt%, Al: 2wt%, Zn: bar.) block (dimensions: diameter 10mm, height 10mm) was used for the present study. Firstly, the specimen was preheated to 400 ℃, and the temperature of the furnace with silicon carbide rods was heated to 1150 ℃. When Cu and Al were melted, Zn was pressed to the melt by a bell jar for 5 minutes. The melt was then poured into a permanent mold, and after solidification, the original samples were obtained. The graphite electrode was inserted vertically into the alloy melt with 3 cm in depth. The electric pulse frequency was 15 Hz, and the pulse voltages were 500 V and 700 V, respectively, and the treatment time was 30 s. The samples were sealed by epoxy resin in a size of 10 mm × 10 mm. The samples for corrosion test were prepared, after polished, degreased by ethanol and ultrasonic cleaned SEM. IVIUM electrochemical workstation and D/max-2500/PC X-ray diffractometer were used to examine the samples.

The polarization curve and EIS AC impedance spectroscopy of the samples with and without EPT were tested in 3.5% NaCl solution by an IVIUM electrochemical workstation corrosion resistance test system. The platinum was used as the auxiliary electrode with 1 cm×1 cm in size. The reference electrode was standard calomel electrode (SCE). The scan speed was 0.5 mV/s and the open circuit potential was in the range from -250 mV to 1000 mV in the polarization curve tests. The sine-wave amplitude modulation was 10 mV and the frequency was 100 KHz-0.01 Hz in EIS AC impedance spectroscopy testing.

2. Results and Discussion

2.1 The effect of microstructure and property of HAl52-2 aluminum brass by EPT

The metallographic microstructure of the samples was observed by a scanning electron microscope and a ZEISS.
Axiovert2000MAT metallurgical microscope, and the results are presented in Figure 1. It can be found that coarse grains are observed in sample (a) without EPT, where the γ phase is star-shaped and distributed aggregatedly on the grain boundary; the grain size of sample (b) with 500 V EPT and the γ phase has decreased and dispersed; the grain size of sample (c) with 700 V EPT has decreased more obviously, and the γ phase has changed into chunk-shaped on the β phase boundary and grains. Based on above, the problem of segregation was solved efficiently.

The microhardness of the samples was tested using an HVS-1000 microhardness tester. The volume fraction of the γ phase was measured and calculated, as shown in Figure 2. It can be seen that the γ phase of the samples with EPT spreads into the matrix in solution treatment. The volume fraction decreased and the microhardness was obviously improved.

2.2 The effect of corrosion resistance of HA152-2 aluminum brass by EPT

The electrochemical corrosion of the samples was tested using an IVIUM electrochemical workstation and the steady-state polarization curves of samples were given in Figure 3. It can be seen that the polarization curve of cathode is smooth, while the polarization curve of anode is flat, suggesting smaller electrode reaction and resistance of the polarizability. Two characteristic regions, the active dissolution region and the transition region of active deactivation, are presented in the polarization curve of cathode of all the three samples. The free corrosion potential of samples with EPT is higher than that without EPT, and the corrosion current density of samples with EPT is lower, implying that the ability of passivation of the samples with EPT was improved obviously.

The polarization curve tested in the experiment was simulated with Tafel curve extrapolation by CorrView Version 3.0 simulation software, and the parameters for the corrosion test are listed in the Table 1.

Based on Figure 3 and Table 1, the free corrosion potential of aluminum brass samples increased by 1% from -0.26602 V without EPT to -0.23969 V with EPT, and the corrosion current density decreased from 9.5110×10⁻⁶ A·cm⁻² to 1.3892×10⁻⁷ A·cm⁻² by two magnitude orders.

It can then be concluded that the corrosion resistance of aluminum brass alloy has been significantly increased.
2.3 The effect of AC impedance spectroscopy of HAl52-2 aluminum brass by EPT

The samples were tested using an IVIUM electrochemical workstation with a sine-wave amplitude modulation of 10 mV and a frequency of 100 KHz-0.01 Hz. The EIS AC impedance spectroscopy tested in the experiment is shown in Figure 4. It can be found that different features were formed in the Nyquist graph of the samples with and without EPT. After EPT, the solid solubility of Al was increased, and Al$_2$O$_3$ oxide film was formed on the surface of the samples. The diffusion of Cu and Zn atom was then blocked and the mass transfer rate was smaller than the formation rate of corrosion products. The mass transfer process was then obstructed, resulting from that the corrosion product layer was formed due to the gradual deposition of corrosion products on the surface of samples. A bigger diameter of the capacitive reactance arc is usually related to a better corrosion resistance of the samples. After EPT, the diameter of capacitive reactance arc increased by 4-5 times from 365.0 to 2020.0 µm, suggesting the enhancement of the corrosion resistance of the aluminum brass with EPT.

![Figure 4. Nyquist pots of aluminum brass alloy in 3.5% NaCl](image)

2.4 The effect of morphology and thickness of dezincification of aluminum brass by EPT

The samples were scanned using an S-3000N scanning electron microscope. The distribution of elements on the electrochemical corrosion surface of HAl52-2 aluminum brass samples was detected, and the results are given in Figure 5. It has shown that Zn distributed heterogeneity on the corrosion surface of the original samples, while the relative Zn content increased and the relative Cu content decreased after EPT.

Figure 6 gives the corrosion morphology of electrochemical corrosion section of the aluminium brass samples. The thicknesses of dezincification layer of the aluminium brass samples were compared, as shown in Figure 7. From Figure 6 and Figure 7, it can be found that the dezincification layer of the original aluminium brass samples was porous (Fig. 6a) and the depth was up to 59.2 µm. The thickness of the dezincification layer with EPT decreased obviously by 90 % at 53.3 µm with 500V EPT. For the samples with 700V EPT (Fig. 6c), it decreased by 84.8 % at 50.2 µm, and the surface of matrix was covered tightly and uniformly. It can then be concluded that the re-corrosion of matrix was prevented, and the corrosion resistance of aluminium brass was efficiently improved.

2.5 The mechanism analysis of the corrosion resistance of HAl52-2 aluminum brass alloys by EPT

The preferential dissolution and resolution deposit mechanism, the double-space mechanism and the seepage mechanism were studied as the main mechanisms of dezincification corrosion. The double-space mechanism and the seepage mechanism were combined by Jihui Wang, and the mechanism of seepage channel was explained by the crystal models of alloys. The double-space was formed during the corrosion process of brass and the resultant selective dissolution of Zn has also been proved.
The XRD analysis of the phase structure of samples with and without EPT is given in Figure 8 and Figure 9, respectively, to further explore the internal relations between the changes of the corrosion resistance and phase structures. The findings show that dezincification occurred on the phase boundary between β and γ phases in the corrosion of aluminium brass alloys, and the β phase was then corroded as anode. The corrosion of the γ phase did not occur until the porous shape was formed on the corrosion layer.

The nucleation rate of the aluminium brass alloy melt was obviously improved and the problem of segregation was solved efficiently by EPT. Therefore, the Zn atom was uniformly distributed in the grains and the quantity of galvanic couple was decreased. The dezincification progress was then inhibited efficiently by EPT. The undercooling of aluminium brass alloy melt was improved by EPT, thus the size of atomic cluster decreased and the quantity increased. Additionally, the critical dimension of nucleation decreased during the nucleation process. The morphology, size and distribution of γ phase can be ameliorated by EPT, and the rate of dezincification was decreased obviously.

3. Conclusions

1. The free corrosion potential of samples with EPT was higher than that in the samples without EPT, and the corrosion current density decreased from $9.5110 \times 10^{-6}$ $\text{A} \cdot \text{cm}^{-2}$ to $1.3892 \times 10^{-7}$ $\text{A} \cdot \text{cm}^{-2}$ by two magnitude orders.
2. After EPT, the thickness of dezincification layer with EPT decreased from 59.2 µm to 50.2 µm. The continuous and pelleted corrosion products were uniformly distributed on the surfaces of the samples. The diffusion of Zn atom was blocked, and the corrosion resistance of aluminium brass was significantly improved.

3. The volume fraction of $\gamma$ phase in the solidification structure of aluminium brass decreased by 0.12%. The microhardness increased by 40% from 198 HV without EPT to 274 HV with 700 V EPT. The average grain size decreased by 50% at 221.3 µm. The diameter of capacitive reactance arc increased by 4~5 times from 365.0 µm without EPT to 2020.0 µm with EPT. The optimum parameters for EPT were determined as 700 V, 15 Hz, and 30 s.

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