Zincate mechanism on cast Al-Si alloy in non-cyanide multi-metal zincate solutions

HUANG Xiao-mei(黄晓梅), LI Ning(李宁), LI De-yu(黎德育), JIANG Li-min(蒋丽敏)
Department of Applied Chemistry, Harbin Institute of Technology, Harbin 150001, China

Received 19 August 2005; accepted 18 November 2005

Abstract: Arrhenius formula was applied to calculate the apparent activation energy of zincate reaction. The standard electrode potential of all the metal coordinating ions and the order of galvanic couple of different metals in zincate solution were also calculated. Electrochemical behavior of zincate process was studied by Tafel polarization curves, $E-t$ curves, and electrochemical impedance spectroscopy (EIS). The results show that the apparent activation energy of zincate reaction in non-cyanide multi-metal zincate solution is smaller than that in simple zincate solution, and precipitation sequence of all the metals in zincate solution is Cu, Ni, Fe and Zn. Relationship between the potential at 30 s before zincate and coverage was attained according to the change of potential of zincate. EIS shows that inductive reactance is produced during zincate.

Key words: cast Al-Si alloy; non-cyanide multi-metal zincate; zincate solution; zincate mechanism; apparent activation energy

1 Introduction

Si is an important element in Al-Si alloys, the content of which greatly affects the physical, mechanical and chemical properties of Al-Si alloys[1–3]. Due to special structure and cast techniques[4, 5], pre-treatment of cast Al-Si alloys before coating is difficult[6–8], and zincate is mainly applied at home and abroad at present[9–11]. LASHMORE[12] stated that zincate technique almost covered the whole market. STOYANOVA and STOYCHEV[13] researched zincate process on Al in alkaline media, and zincate mechanism was put forward by studying the kinetics of electrode reaction. PARAMASIVAM et al[15] and TANG et al[16] also studied the zincate mechanism successively, however, zincate mechanism on cast Al-Si alloys in non-cyanide multi-metal zincate solution was still not reported. Kinetics, thermodynamics and electrochemistry of zincate process in non-cyanide multi-metal zincate solution were systematically studied in this paper, and its zincate mechanism was also discussed.

2 Experimental

The A356 Al-Si alloy contained 7.5% Si, 0.3% Mg, 0.2% Cu, 0.2% Fe, 0.2% Ti, 0.1% Mn and 0.05% Ni (mass fraction). The main components of the non-cyanide multi-metal zincate solution were 110 g/L of NaOH, 18 g/L of Ni$^{2+}$, Cu$^{2+}$, Fe$^{2+}$ and Zn$^{2+}$, 90 g/L of composite complexing agents, 4–6 g/L of additives. The zincate process was the same as elsewhere[17, 18].

Electrochemical tests of zincate process were carried out on CHI 660 Electrochemical Station, with Hg-HgO electrode (NaOH 2 mol/L) as reference electrode, and Pt electrode as counter electrode.

3 Results and discussion

3.1 Kinetics of zincate process

1) Calculation of zincate velocity

Zero-gravity was applied to measure the mass of zincate film, which operated as the difference of the mass of Al-Si alloy test piece after and before zincate process, ignoring the minute dissolved part of the base. The dissolved part could not be calculated accurately right now, which was substituted by the mass loss in NaOH solution with the same concentration of zincate solution. The zincate process was the same as elsewhere[17, 18].

Mass variation curves during zincate process are shown in Fig.1.

It shows that mass increment of the test piece reaches its maximum at 30 s, and then a mass loss is detected. Zincate rate curves made from Fig.1 are shown
in Fig. 2, the macroscopic rate and corrosion rate of Al decrease with the increase of zincate time. From the beginning to 30 s, macroscopic rate and corrosion rate of Al decrease sharply, while after 30 s, they decrease gently, which may be due to the coverage of zinc layer on Al-Si alloy surface, thus the area for replacement reaction of Al and Zn is reduced. The growth rate of zinc reaches its maximum at 10 s, and decreases with the increase of zincate time. Macroscopic rate and deposit rate of Zn in simple zincate solution are larger than that in non-cyanide multi-metal zincate solution, indicating that the existence of multi-metal is beneficial for reducing the deposit rate of zinc, and attaining fine and even zinc crystals.

2) Calculation of apparent activation energy $E_a$ in zincate process

Assuming that the apparent activation energy $E_a$ in zincate process is independent of the temperature, from the following Arrhenius formula:

$$\frac{d \ln k}{dT} = \frac{-E_a}{RT^2}$$

One can obtain

$$\ln k = -\frac{E_a}{RT} + B$$

Thus a straight line can be drawn between $\ln k$ and $-1/T$. The apparent activation energy $E_a$ can be calculated from the slope of the line. Zincate rate, $k$ in this study is the mass of zinc layer per unit area and time. Relationship between the rate constant ($\ln k$) of zincate and temperature is shown in Fig. 3.

From linear fitting, the apparent activation energy $E_a$ of zincate process in non-cyanide multi-metal zincate solution is 34.207 kJ/mol, while that in simple zincate solution is 39.716 kJ/mol, indicating that the existence of Cu, Fe, Ni ions and complexing agents in zincate solution makes zincate reaction easier.

3.2 Thermodynamics in zincate process

1) Standard electrode potential of metal ions in zincate solution and electromotive force of replacement

Fig. 1: Mass variation curves during zincate process: (a) Mass variation curve of zincation layer; (b) Mass loss curve of substrate corrosion; (c) Measured mass of tested sample

Fig. 2: Zincate rate curves: (a) Macroscopic rate; (b) Corrosion rate of aluminum; (c) Growth rate of zinc deposition

Fig. 3: Relationship between zincate rate and temperature: (a) In multi-metal zincate solution; (b) In simple zincate solution
reaction

Ignoring the liberation of hydrogen, cathodic reaction in zincate process is the deposit of metal ions by obtaining electrons:

\[
[ML_x]^n \rightarrow M^{2+} + xL(n-2)/2 \quad (3)
\]

\[
M^{2+} + 2e^- \rightarrow M \quad (4)
\]

In formula (3), \( L \) represents the ligand, \( [ML_x]^n \) is the complexing ions, \( x \) is the coordination number and \( n \) is the charge number of the complexing ions.

Stable complexing ions are formed from metal ions and complexing agents in strong alkaline solutions, thus electrode potential of the metals is different from that under standard conditions. Assume that zinc complexes with NaOH, Fe with seignette salt, Ni and Cu with sodium citrate.

Firstly, oxide film on the surface of Al-Si alloy was dissolved in strong alkaline zincate solution:

\[
2Al_2O_3 + 2NaOH = 2NaAlO_2 + H_2O \quad (5)
\]

Then electrochemical reaction took place on the surface of Al-Si alloy:

\[
2Al + 3[Zn(OH)_4]^{2-} \rightarrow 2Al + 3Zn + 4H_2O + 4OH^- \quad (6)
\]

Electromotive force of the zincate reaction in zincate solution containing complexing agents can be described as

\[
e^0 = E^0_{\text{cathodic}} - E^0_{\text{anodic}} = E^0_{[Zn(OH)_4]^{2-}} - E^0_{Al} =\]

\[
-1.284 - (-0.3) = -1.284 - (-2.3) = 1.016 \quad (8)
\]

If \( e^0 = 0.3 \) V, formula (6) can take place spontaneously, Al on the surface of Al-Si alloy being replaced by Zn and a Zn layer formed. For multi-metal zincate solution, Fe, Ni and Cu can be deposited on the surface of Al-Si alloy together with Zn.

Table 1

<table>
<thead>
<tr>
<th>Metal ions</th>
<th>Complexing agents</th>
<th>( \lg K )</th>
<th>( E^c/V )</th>
<th>( E^s/V )</th>
<th>( e^0/V )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Zn^{2+}</td>
<td>NaOH</td>
<td>17.66</td>
<td>-0.76</td>
<td>-1.284</td>
<td>1.016</td>
</tr>
<tr>
<td>Fe^{2+}</td>
<td>Seignette salt</td>
<td>7.49</td>
<td>-0.44</td>
<td>-0.661</td>
<td>1.639</td>
</tr>
<tr>
<td>Ni^{2+}</td>
<td>Sodium citrate</td>
<td>6.90</td>
<td>-0.25</td>
<td>-0.454</td>
<td>1.846</td>
</tr>
<tr>
<td>Cu^{2+}</td>
<td>Sodium citrate</td>
<td>14.2</td>
<td>0.34</td>
<td>-0.079</td>
<td>2.221</td>
</tr>
</tbody>
</table>

and then Zn^{2+} will be replaced.

2) Galvanic couple order of Al-Si alloy, Zn, Fe, Ni and Cu in multi-metal zincate solution

Tafel polarization curves of Al-Si alloy, Zn, Fe, Ni and Cu in alkaline multi-metal zincate solution are shown in Fig.4. The self-corrosion potential is taken as stable potential and then galvanic couple order is attained, shown in Table 2.

Table 2

<table>
<thead>
<tr>
<th>Electrode</th>
<th>( E_{\text{cor}}(\text{vs Hg/HgO})/V )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cu</td>
<td>-0.238</td>
</tr>
<tr>
<td>Ni</td>
<td>-0.470</td>
</tr>
<tr>
<td>Fe</td>
<td>-0.760</td>
</tr>
<tr>
<td>Zn</td>
<td>-0.996</td>
</tr>
<tr>
<td>Al-Si alloy</td>
<td>-1.652</td>
</tr>
</tbody>
</table>

Galvanic couple order in zincate solution is Cu > Ni > Fe > Zn, in consistent with theoretical values, thus the deposit order of the metals is Cu, Ni, Fe and then Zn. Film dissolution took place in the late period of zincate process, the potential being close to that of Zn, and Cu, Ni and Fe would not be dissolved due to their high potential. EDS and EPMA analyses of the components of zinc layer show higher content of Fe than Ni, which is similar to ‘abnormal co-deposition’ in electroplating and the deposit of Fe restrains the deposit of nickel.
3.3 Electrochemical study of zincate process

1) Tafel polarization curves of zinc layers in 3.5% NaCl solution

Tafel polarization curves of zinc layers on Al-Si alloy attained from non-cyanide multi-metal zincate solution were tested in 3.5% NaCl solution, as shown in Fig.5, and the fitting results are listed in Table 3.

![Fig.5 Tafel polarization curves of zincate films in 3.5%NaCl at different zincate times](image)

<table>
<thead>
<tr>
<th>Zincate time/s</th>
<th>(E_{\text{corr}}) (vs Hg/HgO)/V</th>
<th>(J_{\text{corr}}/(10^{-4}\text{A} \cdot \text{cm}^{-2}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>10</td>
<td>-0.881</td>
<td>3.388</td>
</tr>
<tr>
<td>20</td>
<td>-1.017</td>
<td>2.483</td>
</tr>
<tr>
<td>30</td>
<td>-1.139</td>
<td>2.298</td>
</tr>
<tr>
<td>40</td>
<td>-1.196</td>
<td>2.093</td>
</tr>
<tr>
<td>50</td>
<td>-1.151</td>
<td>2.232</td>
</tr>
<tr>
<td>60</td>
<td>-1.075</td>
<td>1.819</td>
</tr>
</tbody>
</table>

From Fig.5 and Table 3, content of Zn is low at the beginning of zincate process, and large area of Al-Si alloy surface is bare, thus the corrosion current is high. With the increase of zincate time, Al-Si alloy is covered by zinc layer gradually, leading to the decrease in corrosion current. At 60 s, corrosion current reaches its minimum, and corrosion current varies slightly during 20–60 s. Corrosion potential shifts negatively with the increase of zincate time, which reaches its minimum at 40 s and then increases gradually, indicating that the surface is fully covered by zinc layer after 40 s.

2) Tafel polarization curves of zinc layers in zincate solution

Tafel polarization curves of zinc layers on Al-Si alloy attained from non-cyanide multi-metal zincate solution were tested in zincate solution, as shown in Fig.6, and the fitting results are listed in Table 4.

![Fig.6 Tafel polarization curves of zincate films in zincate solution at different zincate times](image)

<table>
<thead>
<tr>
<th>Zincate time/s</th>
<th>(E_{\text{corr}}) (vs Hg/HgO)/V</th>
<th>(J_{\text{corr}}/(10^{-3}\text{A} \cdot \text{cm}^{-2}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>-1.574</td>
<td>5.223</td>
</tr>
<tr>
<td>5</td>
<td>-1.597</td>
<td>5.414</td>
</tr>
<tr>
<td>10</td>
<td>-1.594</td>
<td>5.864</td>
</tr>
<tr>
<td>20</td>
<td>-1.612</td>
<td>6.141</td>
</tr>
<tr>
<td>30</td>
<td>-1.622</td>
<td>5.588</td>
</tr>
<tr>
<td>40</td>
<td>-1.585</td>
<td>5.307</td>
</tr>
<tr>
<td>50</td>
<td>-1.587</td>
<td>5.194</td>
</tr>
<tr>
<td>60</td>
<td>-1.580</td>
<td>5.158</td>
</tr>
</tbody>
</table>

It can be seen from Fig.6 and Table 4 that corrosion potential descends from the beginning to 30 s, and increases after 30 s, varies slightly after 40 s, indicating that the surface is fully covered by zinc layer after 40 s. \(E_{\text{Al/Zn}}^{2+} < E_{\text{Al/Zn}}^{2+} < E_{\text{Zn/Zn}}^{2+}\), which means that Al is dissolved at the beginning of zincate process. From the beginning to 20 s, corrosion current increases gradually, due to the corrosion of Al. After that, the corrosion current descends, mainly due to the deposit of zinc.

3) \(E-t\) curves of zinc layers at different zincate times

\(E-t\) curves of zinc layers at different zincate times are shown in Fig.7, and relationship between the lowest potential and zincate time from \(E-t\) curve is displaced in Fig.8. From the \(E-t\) curves of zinc layers in Fig.7, it
is found that electrode potential descends at the beginning, referring to the dissolution of Al. The electrode potential increases and finally reaches a stable value after its minimum. Fig. 8 shows that the lowest potential decreases in the first 30 s and reaches its minimum at 30 s, while an increase is seen from 30 s to 60 s, indicating that dissolution of Al is decreased due to the coverage of Zn on the surface of Al-Si alloy.

4) $E-t$ curves of Al-Si alloy, Zn, Fe, Ni, Cu, zincate layer in mixture of NaOH solution and composite complexing agents

$E-t$ curves of Al-Si alloy, Zn, Fe, Ni, Cu, zincate layer in a mixture of NaOH solution and composite complexing agents were tested to investigate the change of surface structure of the Al-Si alloy, shown in Fig. 9, and the stable potentials at 240 s are listed in Table 5.

![Fig.7](image7)

**Table 5** Stable potential of different samples after 240 s zincate

<table>
<thead>
<tr>
<th>Sample</th>
<th>$E$(vs Hg/HgO)/V</th>
</tr>
</thead>
<tbody>
<tr>
<td>Al-Si alloy</td>
<td>−1.149 0</td>
</tr>
<tr>
<td>Zincate layer</td>
<td>−1.093 0</td>
</tr>
<tr>
<td>Zn</td>
<td>−1.068 0</td>
</tr>
<tr>
<td>Ni</td>
<td>−0.636 1</td>
</tr>
<tr>
<td>Cu</td>
<td>−0.613 8</td>
</tr>
<tr>
<td>Fe</td>
<td>−0.523 7</td>
</tr>
</tbody>
</table>

Stable potential of Al-Si alloy is $-1.149 \ 0$ V, and that of Zn is $-1.068 \ 0$ V, indicating that the potential will shift positively when Al-Si alloy surface is covered by zincate layer. Practically, stable potential of Al-Si alloy covered by zincate layer is $-1.093 \ 0$ V, indicating that the zincate layer is very thin, however, it's the thin film that inhibits the growth of oxide film on Al-Si alloy, and provides a fine substrate for further electrochemical plating or chemical plating.

5) Relationship between zincate layer coverage and potential

At the beginning of the zincate process, discontinuous zinc alloy crystals grew on the Al-Si alloy surface. With the increase of zincate time, the zincate film became denser, and the surface would be fully covered by zincate film after 30 s. Theoretically, the substrate could not be fully covered by zincate film, for anodic reaction (dissolution of Al) always exists in replacement reaction. Therefore, some fine holes on the
zincate film were generated for the corrosion of Al. Compared with the area of the zincate layer, the holes could be ignored and the coverage \( \theta \) was assumed to be 1, thus the relationship between the zincate layer coverage \( \theta \) and potential \( E \) can be described as:

\[
\theta = aE + b \quad (a \text{ and } b \text{ are constants})
\]

when \( \theta = 0, E_0 = -1.574 \text{ V}; \theta = 1, E_1 = -1.622 \text{ V}. \)

Therefore, formula (9) becomes

\[
\theta = 20.833E - 32.792 \quad (10)
\]

6) EIS analysis of zincate process

EIS was firstly applied to study the zincate process. Due to the complicated and the fast reaction in zincate process, a mixture of NaOH and ZnO was used as the zincate solution. EIS of pure Zn and pure Al in zincate solution are shown in Figs. 10 and 11 respectively.

A large capacitance circle at high frequency is seen in the EIS of pure Zn, which is due to the electric double layer capacitance and electrochemical resistance, indicating that there is only one time constant, referring to the deposition of Zn. EIS of pure Al is more complicated, with a capacitance arc at high frequency, an inductive reactance arc at medium-low frequency.

Equivalent circuit models for pure zinc and pure aluminum in zincate solution are shown in Fig. 12, and the fitting results are listed in Tables 6 and 7.

It can be seen that electrochemical resistance \( R_1 \) of pure Al is much larger than that of pure Zn, indicating that the corrosion of Al in zincate process cannot be neglected. The inductive reactance arc at a medium frequency of pure Al may be caused by adsorption on the surface of the substrate covered by zincate film. With the increasing protection of zincate film, reaction resistance increases for further zincate, appeared as the increase in the radius of capacitance arc at high-intermediate frequency. With the growth of the zincate film, inductive reactance increases. The adsorption on the surface may be attributed by the formation of Al(OH)\(_3\) in alkaline zincate solution in the final period of zincate process.

4 Conclusions

1) The apparent activation energy of cast Al-Si alloy in non-cyanide multi-metal zincate solution was calculated from Arrhenius formula.

2) From the standard electrode potential of metal ions in zincate solution and electromotive force of replacement reaction replacement reaction in multi-metal zincate solution, it is shown that Cu\(^{2+}\), Ni\(^{2+}\), Fe\(^{2+}\) and then Zn\(^{2+}\) could be co-deposited in multi-metal zincate solution. Galvanic couple order of Al-Si alloy, Zn, Fe, Ni
and Cu in multi-metal zincate solution is Cu > Ni > Fe > Zn > Al-Si alloy.

3) Tafel polarization curves and $E-t$ curves show that corrosion potential descends with the increase of zincate time, while ascends in 30 s to 40 s. The coverage of zincate layer on the Al-Si alloy substrate is rather large after 30–40 s.

4) EIS shows inductive reactance arc at the mediate-low frequency in zincate process, due to the adsorption on the surface of the substrate covered by zincate film.

References


(Edited by LI Xiang-qun)