Anodic process on Cu-Al Alloy in KF-AlF₃-Al₂O₃ melts and suspensions

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Abstract

Inert anode development has been a topic of interest for decades. Cu-Al, Fe-Ni based alloys have shown promising results in low-temperature melts. Metallic anodes tend to contaminate the produced aluminium with their corrosion products and introducing alumina suspension particles in the electrolyte could resolve this problem by suppressing the convective transfer of corrosion products toward aluminium produced. In this paper, anodic processes on Cu-10Al electrode in molten KF-AlF₃-Al₂O₃ (5 wt. %) and suspensions were characterized. Effects of cryolite ratio CR (Mole % KF/Mole % AlF₃), temperature and particle volume fraction (φ = 0 – 0.15) on the electrochemical behaviour of the anode were demonstrated. The results suggest that CR = 1.4 at 800 °C with the particle volume fraction of not more than 0.09 are better parameters for the usage of Cu-10Al anode. At melts with φ = 0.15 at all temperatures and CR values, a low current density was observed.

Keywords: aluminium, suspension, oxidation, inert anode, cryolite melt, low-temperature electrolysis

Introduction

Production of aluminium is carried out by aluminium oxide decomposition in the molten sodium cryolite at an industrial scale by using Hall-Heroult cell. The process is been in use for more than 100 years with periodic upgrades. Carbon is used as an anode where CO₂ evolution takes place, leading to the consumption of anodes on a regular basis enforcing the periodic replacement of anodes. Other than CO₂, greenhouse gases like CO, COF₂, CF₄, and C₂F₆ are also emitted. The following reactions occur in the process of aluminium reduction:

\[ \text{Al}_2\text{O}_3(\text{dis}) + \frac{3}{2}\text{C}_l(\text{s}) \rightarrow 2\text{Al}_l(\text{i}) + \frac{3}{2}\text{CO}_2(\text{g}) \] (1)

with (EMF) \(E^0 = 1.168\)V at 1000 °C

\[ \text{Al}_2\text{O}_3(\text{dis}) + 3\text{C}_l(\text{s}) \rightarrow 2\text{Al}_l(\text{i}) + 3\text{CO}_2(\text{g}) \] (2)

with \(E^0 = 1.033\)V,

\[ 4\text{AlF}_3(\text{dis}) + 3\text{C}_l(\text{s}) \rightarrow 4\text{Al}_l(\text{i}) + 3\text{CF}_4(\text{g}) \] (3)

with \(E^0 = 2.155\)V and,

\[ 2\text{AlF}_3(\text{dis}) + 2\text{C}_l(\text{s}) \rightarrow 2\text{Al}_l(\text{i}) + \text{C}_2\text{F}_6(\text{g}) \] (4)

with \(E^0 = 2.378\)V.

This scenario can be avoided by introducing inert anode to the process, which not only eliminates the emission of greenhouse gases, but also the cost associated with the fabrication of the carbon anodes. Inert anodes have been the centre of interest for researchers for a while and attempts are made to find suitable materials to replace existing carbon anode. Oxygen is evolved at the inert anodes and the reaction (5) takes place.

\[ \text{Al}_2\text{O}_3(\text{dis}) \rightarrow 2\text{Al}_l(\text{i}) + \frac{3}{2}\text{O}_2(\text{g}) \] (5)

with \(E^0 = 2.337\)V at 750 °C.

A material should meet the following requirements to qualify as an inert anode: (1) good electrical conductivity, (2) low corrosion rate, (3) high stability at high-temperature electrolysis, (4) easy fabrication and low maintenance [1, 2]. The energy balance and environmental challenges using inert anodes were discussed extensively earlier [3] [4]. Metallic, ceramic and cermet materials were...
tested to date and each has its advantages and disadvantages [5-12]. Metallic anodes have been showing promising results, as they possess high electrical conductivity, good thermal shock resistance. Moreover, the scales formed on the surface of the anode protects it from the oxidation.

During the electrolysis process, the oxide layers are formed on the surface of the metallic anode. If the oxide layer shows good conductivity and stable microstructure while adhering well with the anodic material then it can be considered as a good candidate for anodic material. However, in most cases, the oxide layers react with the cryolite, which results in corrosion. Reaction (6) would occur when the oxide scale reacts with the molten cryolite.

\[
M_xO_y + \frac{2y}{3} Na_3AlF_6 = M_xF_{2y} + \frac{y}{3}Al_2O_3 + 2yNaF
\] (6)

The performance of an anode depends upon the electrolyte used and it is advantageous to use melts with low liquidus temperature while using metallic inert anodes to minimize the corrosion. Extensive studies have been carried out on sodium and potassium cryolites. The aluminium electrolysis process is carried out at around 960 °C using NaF-AlF₃ melt but while using the KF-AlF₃ melts, the process can be performed at 700-800 °C. Low-temperature melts are highly desirable as inert anodes possess low corrosion rate and thermal shocks at these conditions[13-21]. Nevertheless, decreasing the electrolyte temperature leads to an increase in the decomposition voltage by 0.10-0.15V. Selection of Cu-Al alloy as an anode for the present study in the melts and suspensions was based on the following properties possessed by the alloy [22, 23]:

- melting point of the anode is around 1050 °C which allows electrolyte overheating;
- relatively high electrical conductivity of the alloy and its oxide scale (copper oxide and copper aluminates) formed during the electrochemical process;
- stability of the copper aluminates in fluoride melts saturated by the oxygen ions;
- high corrosion resistance at 750 °C in KF-AlF₃ melts with cryolite ratio CR \(\frac{\text{Mole} \% \text{KF}}{\text{Mole} \% \text{AlF}_3}\) = 1.3-1.8.

Suspension melts have also been suggested for the aluminium electrolysis process. The particles should be evenly dispersed with no sedimentation. Zero sedimentation of alumina was obtained at a volume fraction of 0.32 [24]. This type of melt increases the purity of the produced liquid aluminium by stopping the corrosion products from contacting it and resolve the problem of maintaining the electrolyte saturated with alumina [25, 26]. Although, an increase in the volume fraction of alumina results in the increased electrical resistance of the suspension. Although, this issue can be minimized by decreasing the anode-cathode distance.

In the previous work [27, 28], three anodic materials with the compositions Cu – 9Al – 5Fe, Cu – 10Al and Cu – 10Al – 1.7Be were investigated at 750 °C with KF-AlF₃-Al₂O₃ at CR=1.3. Results suggested that Cu – 10Al was performing better out of the three at given conditions and further research must be conducted to explore its performance at different temperatures in different melts and suspensions. Reactions shown in table 1 occur on Cu- 10Al electrode during the oxidation process and the sum of half-reactions of the oxidation-reduction which correspond to the Gibbs energies and are associated with the potential difference (EMF) as shown:

\[
E^0 = -\frac{\Delta G^0}{zF}
\] (7)

where z is the number of electrons, F = 96585 [C/mol] is the Faraday constant.

<table>
<thead>
<tr>
<th>Reaction</th>
<th>(\Delta G^0_{1023K}) kJ/mol</th>
<th>(\Delta H^0_{1023K}) kJ/mol</th>
<th>(\Delta S^0_{1023K}) kJ/mol K</th>
<th>(E^0), V</th>
</tr>
</thead>
<tbody>
<tr>
<td>(Al_2O_3(s) + 3Cu(s) \Rightarrow 2Al(l) + 3CuO(s))</td>
<td>580.21</td>
<td>621.63</td>
<td>40.48</td>
<td>2.00</td>
</tr>
<tr>
<td>(Al_2O_3(s) + 6Cu(s) \Rightarrow 2Al(l) + 3Cu_2O(s))</td>
<td>535.79</td>
<td>595.54</td>
<td>58.40</td>
<td>1.85</td>
</tr>
<tr>
<td>(2Al_2O_3(s) + 3Cu(s) \Rightarrow 1.5Cu_2Al_2O_4(s) + Al(l))</td>
<td>513.90</td>
<td>58.44</td>
<td>44.19</td>
<td>1.78</td>
</tr>
<tr>
<td>(2Al_2O_3(s) + 1.5Cu(s) \Rightarrow 1.5CuAl_2O_4(s) + Al(l))</td>
<td>393.77</td>
<td>453.57</td>
<td>58.44</td>
<td>1.36</td>
</tr>
</tbody>
</table>

| Table1. Possible reactions occurring on Cu – 10 Al anode at 750 °C per 1 mole of Al. |
The present study aims to determine the electrochemical behaviour of Cu – 10Al anode in KF-AlF₃-Al₂O₃ melts and suspensions. The anode was tested with melt (CR = 1.3) at 750 °C with varying particle volume fraction (φ = 0 to 0.15). Ultra-fine alumina particles were used to minimize the sedimentation. CR values and temperatures were varied in the range of 1.2-1.5 and 700–850 °C respectively. This article presents the results of the galvanostatic polarization and cyclic potentiometry studies performed.

**Experimental**

**Material preparation and characterisation**

Alloy with a composition of Cu- 10 Al (in wt. %) was prepared in a vacuum melting furnace at 1050 °C. The materials used for the preparation had a purity of Cu (99.95 %) and Al (99.999%). The equilibrium phase of the Cu-Al system was fcc (α Cu-Al) solid solution [29]. The specimens were cut in a cylindrical shape with dimension φ 15mm × 50 mm. The specimens were treated with degreasing agents (acetone and ethanol) and then dried in the air before the usage. A hole with φ 2.5mm was drilled and threading was made to assemble current lead (steel rod) with the anode. The unused anodic part and the steel current lead were protected with a BN tube insulator.

Electrolytes with CR between 1.2 and 1.5 were synthesized. Firstly, a crystalline form of KF was heated in the vertical furnace to remove the water present in the salt at 400 °C for 4 hours. While synthesizing the electrolyte, the AlF₃ salt was placed at the bottom of the crucible and the KF salt was set at the top as AlF₃ tends to sublimate quickly at 800 °C and higher. The synthesis process was carried out for 3 hours. The melt was mixed well and a portion of alumina was added corresponding to its saturation point in the melt (5 wt.%) with continuous stirring. The temperature during the preparation of melt was monitored throughout the process with the help of a k-thermocouple protected with BN sleeve, which was immersed in the electrolyte. The melt was transferred to a container where it was left to solidification. The solidified melt was then crushed and used accordingly. All the chemicals used were analytical grade and alumina used has an average particle size less than 5µm as shown in Fig. 1.

![Fig. 1. The particle size distribution of Al₂O₃](image)

**Cell design and experimental method**

The three-electrode cell as shown in Fig. 2 was used to perform the experiments. Graphite crucible with high purity was used to contain KF-AlF₃-Al₂O₃ (5wt. %) melt. The crucible also acted as a counter electrode. The reference electrode Al/AlF₃ was used which connected to the measuring device using tungsten rod. The reference electrode setup had a porous BN tube containing the liquid aluminium and KF-AlF₃ electrolyte.
Fig. 2. Experimental setup: 1 - Furnace; 2 - graphite crucible h = 110 mm, d\text{int} = 76 mm, d\text{ext} = 92 mm; 3 - electrolyte/suspension KF-AlF\textsubscript{3}-Al\textsubscript{2}O\textsubscript{3}; 4 - thermocouple; 5 - anode sampled = 15 mm, h = 10 mm; 6 - reference electrode (Al/AlF\textsubscript{3}); 7 - steel current leads, 8 – potentiostat, 9 – BN shields

Autolab PGSTAT302n potentiostat equipped with a 20A booster and controlled by NOVA 2.1.2 software was used to carry out the electrolysis. Electrolysis was conducted using chrono potentiometric method according to which the stable potential was obtained and a relatively stable oxide layer was formed on the anodic surface. Stationary polarization was performed for 0.005 to 1.5 A/cm\textsuperscript{2} current densities where the recording was made with 30 µs of current interruption after a 120s current passage to determine the ohmic voltage drop (IR). Cyclic voltammetry was performed at 0.05 V/s scan rate to examine the possible anodic reactions. The anodic overvoltage was calculated according to the equation:

\[ \eta_a + \eta_c = E_i - E_r - IR \]  

where \( \eta_a \) is the activation overvoltage, \( \eta_c \) is the concentration overvoltage, \( E_i \) is the potential difference between the anode under current and the reference electrode, \( E_r \) is the anodic reversible potential relative to the reference electrode, \( i \) is the current, \( R \) is the resistance. The activation and concentration overvoltages were not divided, and the sum was treated as total anodic overvoltage in this paper.

Results and Discussion

Oxide layer characterisation

Initially, anodes were polarized in melts (CR = 1.3) and suspensions (\( \phi = 0.12 \)) for 1 to 1.5 hours at 750 °C with the current density \( i = 0.4 \) A/cm\textsuperscript{2} to record the change of total voltage being supposedly associated with the oxide layer formation vs. time as shown in Fig. 3

Fig. 3. Galvanostatic polarization CR=1.3 750 at 0.4 A/cm\textsuperscript{2}
Stationary anodic polarization in melt changed between 2.5 V and 3.1 V. The anodic potential was increased from 2V (open circuit potential) up to 3.1 V. Consistent change in the anodic potential reflects the formation of different types of oxide layers (copper oxides and copper aluminates) and changes in its structure. High oscillations of the anodic potential might be associated with the formation of new oxide layer on a consistent basis or due to the bubble evolution on the anodic surface leading to the reduction of the active surface area. No stable oxide layer was formed even after an hour of polarization. The stable oxide layer was formed after 5000 seconds of polarization. In case of anodic polarization in suspension ($\varphi = 0.12$), a stable oxide layer was formed from the beginning of the polarization and only a minor shift of anodic potential was observed. No fluctuations in the voltage were seen unlike anode tested in melt. The sedimentation was not observed.

The XRD results from Fig. 4 and Table 2 show the composition of oxide layers formed on the anode during the polarization.

![XRD-graphs recorded on oxide layers after anodic polarization in melts and in suspensions with $\varphi = 0.15$ at 750 °C: A) electrolyte phases, B) Cu$_2$O, C) CuAlO$_2$, D) Cu, E) CuAl$_2$O$_4$, F) CuO](image)

**Table 2.** Oxide layer composition after polarization in melt and suspension (wt.%)  

<table>
<thead>
<tr>
<th>$\varphi$</th>
<th>CuO</th>
<th>Cu$_2$O</th>
<th>CuAlO$_2$</th>
<th>CuAl$_2$O$_4$</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.00</td>
<td>0</td>
<td>30.34</td>
<td>53.63</td>
<td>16.03</td>
</tr>
<tr>
<td>0.15</td>
<td>31.16</td>
<td>42.84</td>
<td>11.77</td>
<td>14.22</td>
</tr>
</tbody>
</table>

No CuO oxide layer was observed from XRD results and the dominant oxide layer in terms of proportion was CuAlO$_2$ for melts ($\varphi=0$), while in suspensions ($\varphi= 0.15$) the most dominant oxide layer on the surface of the anode was Cu$_2$O, while CuAlO$_2$ content was low unlike in melts ($\varphi=0$). No CuF or CuF$_2$ was found on XRD examinations meaning CuF and CuF$_2$ produced from the process are dissolved in the electrolyte. The anodes were tested with stationary and non-stationary methods in melts ($\varphi=0$) and suspensions ($0.03 \leq \varphi \leq 0.15$) at different temperatures and CR to find the best conditions at which the anode can perform.

**Effect of temperature and CR**

To characterize the effect of temperature and CR on the anodic behaviour of Cu – 10Al alloy both stationary and non-stationary polarization techniques were applied. Cyclic voltammetry was performed on anodes at 700 and 800 °C in melts saturated with alumina at CR = 1.2 – 1.5. Voltammograms recorded to observe the response of anode with the change in potential at 0.05 V/s scan rate are illustrated in the Fig. 5.
Several regions can be distinguished on the curves. The $ab$ region between 2.1 and 2.3 V is where the metal oxidation takes place. After that, a region $bc$ with a plateau or a slight decrease in current with the potential shift toward more positive values can be observed. In [30] similar regions were indicated as the passivation of the anode. The transpassivation process of the anode takes place in the region $cd$ where the oxygen starts to evolve. The current density fluctuation at potentials after 2.5 V is due to the bubbles evolution on the anode surface leading to the constant change in the active anode surface and the interpolar resistance. No passivation of the anode was observed at 800 °C for $CR = 1.4$ and $CR = 1.5$. With the decrease in the CR, reduction of anodic current density at the particular anodic potentials (in oxygen region $cd$) can be seen.

For melts with $CR = 1.2$ at 700 °C, current density remains low, indicating the passivation of the anode from the beginning of the process. After increase in the temperature to 800 °C the shape of the curve in the oxygen region had changed. The region $cc''$ appeared where the $\partial i / \partial E$ value was higher than that of $cc'$ and $c'd$ regions. The reason might be associated with change in resistance during the sweep due to passivation and transpassivation phenomena. High current densities were achieved while using melts with $CR = 1.5$. This can be related to higher alumina solubility in melt at higher CR.

The reversible potentials and the stationary potentials at $i=0.4$ A cm$^{-2}$ obtained through the galvanostatic process for $1.2 \leq CR \leq 1.5$ are illustrated in Fig. 6. An increase in the CR results in the decrease of the anodic potential and evolution of oxygen at low current densities. The reason is that an increase in AlF$_3$ concentration in the melts leads to the formation of oxyfluoroaluminates like $\text{AlOF}_x^1-x \text{Al}_2\text{OF}_x^4-x \text{Al}_2\text{O}_2\text{F}_x^2-x$, $\text{Al}_4\text{OF}_6$, $\text{Al}_6\text{O}_2\text{F}_{10}$ at higher rates leading to the difficulties in the ions transport between the anode and the cathode [32,33].

![Fig. 5. Cyclic voltammograms for the anode in melts with different $CR$ (1.2-1.5) at 700 and 800 °C](image)

![Fig. 6. Dependence between reversible potential (left) and stationary potential at 0.4 A/cm$^2$ (right) in melts at 700, 750 and 800 °C vs. CR](image)
In case of melts with CR = 1.5, the effect of temperature on the anodic potential was not observed until the oxygen evolution potential (2.34V). The limiting current densities were observed at high potentials (3.2-4.2 V) for CR = 1.2 at both the temperatures. In case of CR= 1.4, limiting currents were in the region from the potential 2.9V. At CR = 1.5, the limiting current region is between the potentials (3.4 -4.2V) in melts at 700 ℃ and no limiting currents were observed at 800 ℃. In melts with CR= 1.4, drastic variation in current density w.r.t anodic potential was seen with the change in temperature.

According to the obtained results, the anode performed better at 800 ℃. A further increase in temperature is undesirable due to probable decrease in corrosion resistance. The CR should be maintained at CR≈1.4. The reasons for the chosen parameters are:

- the onset potential of oxygen evolution is more negative at higher temperatures and higher CR;
- high overvoltage is observed at temperatures below 800 ℃;
- the stationary anodic potentials at current density 0.4 A cm⁻² were similar in melts with CR=1.4 and CR=1.5;
- in terms of the cathodic process the lower CR is preferable due to higher limiting currents of aluminium reduction [31], so the optimal CR value can be between 1.3 and 1.4.

**Effect of particles volume fraction**

Stationary galvanostatic polarization curves were recorded for melts (φ=0) and suspension (φ=0.03…0.15) under chosen conditions (T=800 ℃, CR=1.4) to estimate the limiting current density and the effect of particle volume fraction on the polarization curve. The data was compared to the previously obtained results for initially chosen conditions (T=750 ℃, CR=1.3). The plots are shown in the Fig. 7 [27].

![Stationary galvanostatic polarization curves of the anodic process in melts and suspensions (φ=0-0.15) with CR 1.3 and 1.4 at 750 ℃ and 800 ℃](image)

The metal oxidation takes place in the region ab. The potentials in melt (φ=0) with CR=1.3 indicate the formation of copper aluminates rather than oxides (see Table 1). At point b, the potential was equal to that of the Cu/CuO electrode, which shows the formation of CuO layer on the electrode in the region bc. Point c is where the oxygen evolution starts and it can be seen that the oxygen evolution takes place at high i in the melts (φ=0) and starts at low i while using suspensions. A sharp increase in the current density indicating the oxygen evolution onset can be seen in the regions cd and c’d in suspension. The electrode in suspensions at φ=0.15 had no metal oxidation region ab.

The limiting current densities were observed in the de region. It is observed that the electrode used in melts (φ=0) possess high limiting current density while an increase in φ drops the limiting
currents, which might be due to the decrease in mass transfer coefficient $K_s$ of the oxyfluoride ions. States that with increasing suspension volume fraction in melts, limiting current decreases.

The comparison of the described results with those obtained in suspensions with CR 1.4 at 800 °C allows us to conclude that the limiting currents can be observed at 2.5-3 V in the suspension with $\varphi = 0.12-0.15$. In case of suspensions with lower $\varphi$ (0.09) and melts, limiting current densities were not achieved up to 1 A cm$^{-2}$. With an increase in the volume fraction $\varphi$ from 0.12 to 0.15, the passivation of the anode was observed. From the galvanostatic curves, it can be concluded that the volume fraction with no more than 0.09 can be used. Further increase in the $\varphi$ leads to drastic increase in overvoltage.

The cyclic voltammetry recordings were made at a scan rate of 0.05 V/s in suspensions ($\varphi=0.03-0.15$) at 800 °C and compared to those obtained previously [27] as shown in the Fig. 8.

![Fig. 8. Cyclic voltammograms recorded in melts suspension with CR 1.3 and 1.4 at 750 °C and 800 °C](image)

In case of pure melts, low current densities were observed, reflecting metal oxidation stage in the regions $ab$ and $bc$. The potential at point $b$ was 2.07 V, which is closer to the potential corresponding to Cu/CuO electrode (2.00 V). CuO might have been a dominating oxide layer on the surface of the electrode in the $bc$ region while before that rather copper aluminates or Cu$_2$O were formed. Oxygen evolution happens from point $c$ where the potential of the anode was above 2.5 V in the melt with CR=1.3. It has shifted toward more negative value after increase in CR. Point $e$ separates the pre-oxygen and oxygen evolution regions. At the region $dc$, an increase in the current density can be seen and reason might be due to the reaction (13). In suspensions with high $\varphi$ (0.12 and 0.15) at point $e$, the potential is more than 2.6 V, which is close to the EMF of the reaction (14). The active CuO layer oxidation may take place leading to the formation of CuF$_2$ and O$_2$ evolution and it states the catastrophic corrosion of an oxide layer.

$$\text{AlF}_3(l) + \text{Cu}_3(s) \rightarrow \frac{3}{2} \text{AlF}_2(s) + 3\text{Cu}^{2+}(l)$$ \hspace{1cm} (13)

with $E^0 = 2.31$ V

$$\frac{4}{3}\text{AlF}_3(l) + 2\text{CuO}_3(s) \rightarrow 2\text{CuF}_2(s) + \frac{4}{3}\text{Al}^{3+}(l) + \text{O}_2(g)$$ \hspace{1cm} (14)

with $E^0 = 2.65$ V

In suspensions with $\varphi = 0.12$, oxygen evolution takes place from the beginning at 2.45V and reaction (15) is expected which occurs at potential 2.29 V, readily formed Cu$_2$Al$_2$O$_4$ oxide on the anodic surface at the beginning of the voltammetry process is transformed to CuO and oxygen evolution takes place at the same time. In the regions $cd$ and $de$, the dissolution of the CuO oxide layer takes place with continuous O$_2$ evolution at potentials close to those required for the reaction (14) where CuO layer reacts with the AlF$_3$ present in the electrolyte and CuF is expected to be the product along with Al$_3$ and evolution of O$_2$. 
\[1.5\text{Cu}_2\text{Al}_2\text{O}_4 + \text{Al}_2\text{O}_3 \xrightarrow{4\text{F}} 3\text{CuO} + 5\text{Al}_\ell + 3\text{O}_2(\ell)\]  \tag{15}

with \(E^0 = 2.29\) V

In the \(cd\) region, passivation of the anode may take place where the reason might be the formation of the specific oxyfluoroaluminates. In suspensions with \(\varphi = 0.15\), since the beginning of the sweep, high resistance was observed and no reactions occurred with considerable rate. The anode surface was completely passivated.

The dependence between the stationary current density at 2.7 V and the \(\varphi\) for suspension with CR=1.4 at 800 °C is shown in Fig. 9.

![Fig. 9. Dependence between stationary current density \(i\) at anodic potential \(E=2.7\) V (relative to Al reference electrode) and alumina particles volume fraction in suspension with CR=1.4 at 800 °C](image)

It can be observed that the stationary current slightly decreases in the range of \(\varphi\) between 0 and 0.09. After the introduction of additional alumina particles, the limiting current of oxygen evolution seems to be decreased. As a result the observed stationary current decreases rapidly with increase in \(\varphi\).

\[k_s = \frac{i_l}{zF\mathcal{C}}\]  \tag{16}

where \(k_s\) [cm/s] is the mass transfer coefficient, \(z\) is the number of electrons, \(i_l\) [A/cm\(^2\)] is the limiting current density, \(F= 96500\) [C/mol] is the Faraday’s constant, \(\mathcal{C}\) [mol/cc] is the solubility. The mass transfer coefficients are calculated for the set of conditions and presented in Table 3.

<table>
<thead>
<tr>
<th>(T/\degree C)</th>
<th>CR</th>
<th>(\varphi)</th>
<th>(i_l/A\cdot cm^{-2})</th>
<th>(\mathcal{C}/mol\cdot cc^{-1})</th>
<th>(k_s\cdot 10^4/cm\cdot s^{-1})</th>
</tr>
</thead>
<tbody>
<tr>
<td>750</td>
<td>1.3</td>
<td>0</td>
<td>0.90</td>
<td>0.00901</td>
<td>5.175</td>
</tr>
<tr>
<td>750</td>
<td>1.3</td>
<td>0.12</td>
<td>0.60</td>
<td>0.00901</td>
<td>3.450</td>
</tr>
<tr>
<td>750</td>
<td>1.3</td>
<td>0.15</td>
<td>0.15</td>
<td>0.00901</td>
<td>0.862</td>
</tr>
<tr>
<td>800</td>
<td>1.4</td>
<td>0.03</td>
<td>1.10</td>
<td>0.00738</td>
<td>7.722</td>
</tr>
<tr>
<td>800</td>
<td>1.4</td>
<td>0.12</td>
<td>0.20</td>
<td>0.00738</td>
<td>1.404</td>
</tr>
<tr>
<td>800</td>
<td>1.4</td>
<td>0.15</td>
<td>0.02</td>
<td>0.00738</td>
<td>0.140</td>
</tr>
</tbody>
</table>

Clearly, \(k_s\) value decreases with the increasing volume fraction in both cases; although the \(k_s\) values decrease with the increase in CR. The limiting currents for CR 1.4 at low \(\varphi\) were not observed resulting in the difficulties to estimate \(k_s\). It is worth noting that \(k_s\) slightly decreases with an increase in \(\varphi\) if \(\varphi\leq0.12\). However, any further increase in \(\varphi\) leads to the drastic drop of \(k_s\). The obtained \(k_s\) values are relatively less compared to the values obtained for gas evolving electrodes [37,38].

The industrial process of aluminium reduction in alumina suspensions seems to be operable with \(\varphi=0.09\) if the highly dispersed alumina is used.

**Conclusion**

The influence of dispersed alumina particles content in melts on the current-potential characteristics is discussed. The main conclusions:

- The dominant oxide layer on the anode surface in melts (\(\varphi=0\)) was CuAlO\(_2\) and while using suspension (\(\varphi=0.15\)) Cu\(_2\)O was the dominant one;
□ with the increase in the alumina volume fraction, the limiting current densities of metal oxidation and oxygen evolution decrease;
□ passivation of the anode at the beginning of the cyclic voltammetry process was seen at 700 °C at all CR values (1.2-1.5). The reasons might be the oxide formation or the reduction in the anodic active surface area;
□ particle volume fraction (φ) of 0.09 is suggested for the further process development;
□ anode possesses high oxygen evolution limiting currents at 800 °C.

Further work will be conducted to analyze the current efficiency with Cu-10Al alloy at 800 °C in suspension with φ = 0.09 and CR=1.4.

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References
