Reduction of Solid Al₂O₃ with Electrolysis of CaCl₂-Based Melt

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Cyclic and square-wave voltammetry methods are used to investigate the mechanism of the cathode process on molybdenum in a CaCl₂-CaF₂ melt at 750 °C. It is shown that calcium reduces on an inert cathode in the form of its solution in catholyte at potentials more positive than are required for metallic calcium formation. The presence of Al₂O₃ in the catholyte increases currents of calcium reduction on the forward cathodic wave at potentials more positive than are required for metallic calcium formation. A decrease in the ohmic resistance between the anode and cathode is observed to have been caused by the appearance of electronic conduction during the calcium cathodic reduction in the studied melt. A mechanism for the reduction of Al₂O₃ in the catholyte during electrolysis of the CaCl₂-CaF₂ melt, including Ca⁺ subions and calcium cathode formation and the secondary reduction of Al₂O₃, is proposed. In order to demonstrate the supposed mechanism for the Al₂O₃ reduction, the electrolysis tests were performed in two laboratory-scale electrolyzers with different separation type of anolyte and catholyte. The necessity of separating the anolyte and catholyte for providing stable continual electrolysis in the CaCl₂-based melt is demonstrated.

The aim of this work was to obtain additional experimental data for the electrolysis tests in two laboratory electrolyzers supplied with different types of anolyte-catholyte separation.

Experimental

Melt preparation.— The subject of testing was a (wt%) 80CaCl₂-20CaF₂ melt with decreased liquidus temperature and lowered hydrolysis capability in comparison with pure CaCl₂. The following individual reagents were used for its preparation: anhydrous CaCl₂ and CaF₂—both chemically pure grade (99.9 wt%, CJSC “Vektón”). The salt mixture was placed in an experimental reactor (alumina crucible) in a closed quartz tube under dry argon atmosphere and gradually heated over a period of 24 hours to an operating temperature of 750 °C. The prepared melt was electrolyzed in potentiostatic mode using a molybdenum cathode (99.95 wt%, CJSC “OZTM&TS”) and graphite anode (spectral pure grade) with the cathode potential of −2.5 V relative to the CO/CO₂ gas electrode, which was placed in a porous alumina container (open porosity—1.2 vol. %) together with the (wt%) 80CaCl₂-20CaF₂ melt. No oxide was specifically added into the CO/CO₂ gas electrode semi-element electrolyte due to the presence of residual oxygen-containing forms in CaCl₂-CaF₂ being taken into account. The duration of the pre-electrolysis was 12–18 hours, with the termination being determined experimentally (with the relative achievement of a current minimum). The melt temperature was regulated by a Varta TP-703 thermoreactor and Chromel-Alumel thermocouple.

Aluminium oxide was chosen as a reducible reagent due to its minor solubility (less than 0.005 wt%) in the melts under study. This data supported by the elemental composition of the metals and alloys sintered by electrolysis of CaCl₂-melts in alumina crucibles.

Electrochemical measurements.— The electrochemical measurements were carried out in a closed three-electrode alumina cell in a dry argon atmosphere using a PGSTAT AutoLab 302N and NOVA 1.11 software (Metrohm, Netherlands) (with a cell scheme similar to that of the electrolyser presented in Fig. 1). Cyclic and square-wave voltammetry methods were used to determine the features of the cathode processes occurring on the molybdenum substrate. Voltammograms were recorded at potential sweep rates from 0.01 to 1 V s⁻¹. The square-wave voltammograms were recorded at a frequency of 15 Hz and an amplitude of 50 mV. A molybdenum rod (surface area—1.1 cm²) served as the working electrode; the counter electrode was...
graphite (surface area = 14 cm²). The gas CO/CO₂ electrode was used as a quasi-reference one. The gas CO/CO₂ electrode was realized by placing the graphite rod (forming the electrode substrate) in the porous sintered alumina case (OJSC “Ogneupory”, open porosity = 1.2 vol. %) containing the CaCl₂-CaF₂ melt. Due to the Boudouard reaction between the graphite and residual oxygen species in the melt (CO₂ + C = 2CO), an equilibrium atmosphere was established and maintained in the alumina container for the duration of the experiment. There were no holes in the alumina container since this function (electron transfer) was performed by the pores. The optimal open porosity value was chosen experimentally. Details of the behavior of the CO/CO₂ gas electrode in oxide-halide melts in terms of potential stability, reproducibility and irreversibility was described in our previous work. In order to measure and compensate the ohmic resistance (R) of the electric circuit, the current interrupt (I-Interrupt) method was used. The ohmic resistance value was derived from the ohmic drop (ΔU = IR) value, which was typically achieved 2–5 mks after the current (I) interruption. The accuracy of the measured R-values was also checked from the Nyquist plots obtained from the impedance measurements (FRA method).

**Electrolysis tests.**—Electrolysis of the CaCl₂-CaF₂ melt was performed at 750 °C in two lab electrolyzers (amperage = 20 A) with different electrode arrangements. Both electrolyzers were placed in a closed quartz tube filled with dried argon. A molybdenum rod (Ø = 1 mm, immersion = 15–40 mm) in an alumina container served as the cathode, and a dense graphite rod (Ø = 10 mm, immersion = 50 mm) served as the anode. The cathode potential was set relative to the potential of the gas CO/CO₂ electrode. An alumina tube (for the both electrolyzers – OJSC “Ogneupory”; inner diameter = 18 mm; thickness = 3 mm; open porosity = 0.5 vol. %) was used to separate the anolyte and catholyte. In the first construction (Fig. 1, left) the cathode and Al₂O₃ powder (8 g) were placed into the alumina tube; two holes (Ø = 0.5 mm) were made to provide a passage for the electrical current. In a second construction the electric current was passed through the layer of Al₂O₃ powder at the bottom of alumina container. The alumina tube was immersed in the Al₂O₃ layer, which provided a good separation of the anolyte and catholyte in this construction. The molybdenum cathode and Al₂O₃ powder (8 g) were placed in an alumina crucible (Fig. 1).

**Pre-electrolysis.**—The voltammograms obtained on molybdenum with a potential scan rate of 0.1 V s⁻¹ after melt preparation (curve 1) and following pre-electrolysis (curve 2) for 2 hours at cathode potential E₉₀ ≈ 0.9 V (recalculated against the potential of saturated calcium electrode) are presented in Fig. 2. The peak at the cathode potential E₉₀ ≈ 1.8 . . . 2.4 V is apparently associated with the discharge of the residual hydrogen, which presented itself in the form of hydrolysis products (Ca(OH)₂, Ca(OH)Cl and others). The discharge in the measuring cell with the CO/CO₂ electrode can be described as follows:

\[
\text{Ca(OH)₂} + \frac{1}{1 + x} \text{CO} + \frac{1}{1 + x} \text{CO₂} + \text{H₂} + \text{CaO}
\]

where x is the volume fraction of CO₂ in the semi-element of the reference electrode, which corresponds to the equilibrium of the Boudouard reaction at certain temperature, (1 - x) is the volume fraction of CO. Evaluation of the hydrogen discharge potential according this Reaction 1 is complicated because of the unknown activities of Ca(OH)₂ and CaO and partial pressure of H₂ in the semi-element of the working electrode.

In our previous experience with the CO/CO₂-electrodes, we have been able to achieve stable, reproducible potential even while utilizing simple manufacturing techniques. We use electrodes of this type in a study for these reasons. It is not possible to carry out thermo-dynamic calculations in accordance with reactions of type (1) using the CO/CO₂ electrode. The potential of the CO/CO₂ electrode was estimated experimentally against the potential of the saturated calcium electrode. The saturated calcium electrode consists of calcium supplied with the molybdenum rod as a sensor of its electrical potential. In our experiments it’s a (dynamic) working electrode, on which the calcium mole-fraction activity is closed to (1) due to the high cathode current density. Further, all potentials were recalculated against the potential of the saturated calcium electrode (marked on the graphs as a dashed line). An analogous approach was used by Bard. Following pre-electrolysis, the peak value decreased from 0.33 to 0.16 A cm⁻².

**Analysis of melt and synthesis products.**—In order to determine the peculiarities of the Al₂O₃ reduction mechanism, the composition of the products in the catholyte and pores of the alumina tube were analyzed by means of Raman spectroscopy (RAMAN), X-ray diffraction (XRD) and scanning electron microscopy (SEM + EDX) methods. The following equipment was used: Renishaw U-1000 Raman microscope-spectrometer (Renishaw, UK); Rigaku D/MAX-2200VL/PC automatic X-ray diffractometer (Rigaku, Japan); JEOL X-act ADD + JSM-5900LV scanning electron microscope with
nitrogen-free energy dispersive detector (JEOL, Japan) supplied with a wave dispersive micro analyzer, sluice chamber and INCA Energy 250 and INCA Wave500 devices for suppression of electromagnetic interference (Oxford Instruments, UK).

Results and Discussion

**Kinetics of the cathode process.**—A gradual increase in the cathodic current density from the potential of about 0.9 V and its sharp increase at the potential range of 0.1 . . . 0.0 V is observed on the voltammogram obtained on molybdenum in the CaCl₂-CaF₂ melt without Al₂O₃ in the catholyte (Fig. 2, curve 2). Such a course of the non-stationary i-E dependence confirms the results of stationary polarization studies on inert cathodes in the CaCl₂-KCl melt. Analysis of stationary polarization curves revealed that a number of electrons of the cathode process occurred at more positive potentials than calcium metal reduction is equal to one. This process can be explained by the formation of the calcium subions solution in calcium chloride:

\[
\text{Ca}^{2+} + e^- \rightarrow \text{Ca}^+ \tag{2}
\]

A similar mechanism of the process is observed during cathodic reduction of alkali and alkaline earth metals.

The addition of non-conductive insoluble oxide powder (Al₂O₃) into the catholyte leads to an additional rising of the cathode currents at potentials more negative than 0.9 V. In our opinion, this happens due to the fact that cathodically formed Ca⁺ (in the form of calcium sub-chloride CaCl₉) can be consumed in the catholyte by oxide, for example, according to reaction:

\[
\text{Al}_2\text{O}_3 + 6\text{CaCl} \rightarrow 2\text{Al} + 3\text{CaO} + 3\text{CaCl}_2 \tag{3}
\]

This leads to a kinetic increase in cathodic Reaction 2 currents during the non-steady experiment at potentials more positive than the thermodynamic potential of the calcium metal formation. Such an interpretation of the reduction mechanism of oxides to metals in the CaCl₂-based melts at cathode potentials more positive than the potential of the metal calcium formation is consistent with the results of electrolysis tests described in previous works.

An increase in the potential sweep rate to 5 V s⁻¹ results in an increase in the cathodic current in the potential range of 0.9 . . . 0.1 V (Fig. 3) even while the features of the solid oxide reduction peaks remain the same (increased cathode currents on the right semi-waves of voltammograms). In accordance with these observations, we think it likely that the subsequent chemical Reaction 3 limits the summary process in the melt under study even at high solubility of CaO in the CaCl₂-based melts. This can explained by limitations in CaO (or O²⁻ ions) transfer from the reducible oxide into the melt though a sponge layer of reduced metal, pores and capillaries which are, most probably, filled with double oxides and melt saturated with CaO. Moreover the presence of a double oxide in pores can complicates both the removal of CaO and admission of reductant (electrons from the Ca⁺ cations) to the reaction front of oxide reduction. This assumption has been confirmed by the electrolysis tests performed using lab scale electrolyzers having different catholyte arrangements.

In order to clarify features of the cathode process square-wave voltammetry was also used. The typical square-wave voltammograms obtained on molybdenum with and without Al₂O₃ powder in the catholyte are demonstrated in Fig. 4. One can see two precise peaks at the cathode potentials of about 0.75 and 0.30 V. Wherein peaks

![Figure 2. Voltammograms, obtained on molybdenum in the CaCl₂-CaF₂ melt at 750°C and a potential sweep rate of 0.1 V s⁻¹ without (curves 1, 2) and with Al₂O₃ powder (curve 3) in the catholyte. Gas CO/CO₂ quasi-reference electrode.](image)

![Figure 3. Voltammograms, obtained on molybdenum in the CaCl₂-CaF₂ melt at 750°C and potential sweep rates of 0.1–5.0 V s⁻¹ with the Al₂O₃ powder in the catholyte. Gas CO/CO₂ quasi-reference electrode.](image)
occur at voltammogram obtained in a melt without the Al₂O₃ powder. This suggests that process (2) takes place at the potential of 0.75 V, and the electroreduction of Ca²⁺ subions to calcium thorough the reaction

\[
\text{Ca}^+ + e^- \rightarrow \text{Ca}
\]  

[4]
takes place at the potential of 0.30 V. One can see the wave of metal calcium formation at cathode potentials more negative than minus 0.1 V, which are accompanied by fluctuations of the measured current. We explain this feature in terms of the formation of an ion-electron liquid in a catholyte saturated with Ca²⁺ subions.⁷–¹⁰ The geometry (geometric shape and surface area of the region where an ion-electron liquid distributed) of the latter is variable because of the parallel (geometric shape and surface area of the region where an ion-electron liquid distributed) of the latter is variable because of the parallel

Electrolysis tests.—Formation of Ca⁺ subions by Reaction 2 or by reaction:⁵,⁶

\[
\text{Ca} + \text{CaCl}_2 \rightarrow 2\text{CaCl}
\]  

[5]
was observed during the preliminary electrolysis tests in the electrolyser as shown in Fig. 1 (left). The latter leads to the appearance of the electronic component of the common conductivity of the investigated melt.⁵,⁷

Even small amounts of passed current (0.1–0.4 A · h) decrease the melt electroresistance from 1.5–2.3 to 0.15–0.22 Ohms in our experiments. Therefore, a correction (especially at the initial stage) of the compensated electroresistance should be carried out for stable organi-
part of the melt after its washing in distilled water) are given in Fig. 6. One can see the phases of Ca$_{12}$Al$_{14}$O$_{33}$, CaF$_2$, and Al$_2$O$_3$ in the melt as well as in the alumina tube. The phases of Al, Al$_2$Ca, Al$_4$Ca were obtained in the metal droplets. The absence of intermediate CaCl is associated with its instability, as well as with the fact that the analyses were not carried out in situ in the molten state. The presence of metallic calcium (in the intermetallic compounds) may be caused both by the Reaction 5 and by the direct calcium discharge at the cathode with the depolarization. In the case of electrolysis, the depolarization can be caused by the formation of alloys (Al$_2$Ca, Al$_4$Ca).

The obtained results confirm the presence of the subsequent chemical reaction, which can limit the summary process in the case of the electrolysis products, the synthesis mechanism of aluminum and catholyte. However, a systematic study of the synthesis parameters is required in order to develop the necessary technology for their production.

Conclusions

The kinetics of the cathodic formation of calcium in the CaCl$_2$-CaF$_2$ melt at 750°C was studied by means of cyclic and square-wave voltammetry. It was revealed that the formation of calcium takes place at potentials more positive than the thermodynamic potential of the metallic calcium formation. The presence of the Al$_2$O$_3$ powder in the catholyte leads to the subsequent chemical reaction of the consumption of the reduced forms of calcium (Ca$^+$, Ca). The increase in the cathode current confirms this statement.

In order to demonstrate the features of the suggested Al$_2$O$_3$ reduction mechanism in the CaCl$_2$-CaF$_2$ melt at 750°C, the electrolysis tests were performed in two electrolysers with different approaches to separating the anolyte and the catholyte. Based on the analysis of the electrolysis products, the synthesis mechanism of aluminum and its alloys during the electrolysis of CaCl$_2$-based melts was described. The possibility in principle of the production of aluminum-calcium alloys using an electrolyser with graphite anode was demonstrated and the main features of this process described.

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Figure 6. XRD (a) and RAMAN (b) analysis of the compositions of the different phases after the potentiostatic electrolysis (at $E_{Mo} = 0.6...0.4$ V during 24 hours) of the CaCl$_2$-CaF$_2$ melt at 750°C with the screened anolyte and Al$_2$O$_3$ powder in the catholyte.

Figure 7. Photography of the oxidized metal droplets (a), microphotography (SEM) of the shear of the alumina tube following their holding in air (b) and carbon distribution in alumina tube (c).
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