Oxidation electroquímica de plata y cobre en medio acuoso básico y en hidróxidos fundidos

E. M. TEJADA-ROSALES, N. CASAÑ-PASTOR, P. GÓMEZ-ROMERO*
Institut de Ciència de Materials de Barcelona (CSIC). Campus de la UAB, 08193 Bellaterra, Barcelona, Spain.

1. INTRODUCTION

The synthesis of novel metal oxides has relied traditionally upon conventional solid-state reactions at high temperatures. More recently, the advantages of soft-chemistry and low-temperature techniques to yield new metastable compounds have been recognized[1]. But when it comes to the preparation of oxides with metals in unusual oxidation states further control is necessary. Certain soft chemistry routes have been shown to be well suited for the preparation of oxidized compounds (by in-situ oxidation)[2], but the most obvious choice for a better control of oxidation state is the use of electrochemical techniques.

As part of our continuing effort to probe novel synthetic techniques for the preparation of electroactive oxides we have explored the possibilities of electrochemical reactions and crystallization processes in basic (molten hydroxide) fluxes. This combination allows for the simultaneous control of basicity and electrochemical potential at a relatively low reaction temperature and has been used before for the isolation and crystallization of silver oxides from silver anodes [3-7] or from silver solutions [8-10] and Cu(III) oxides for instance [11]. Furthermore, molten hydroxide media offer an electrochemical-potential window large enough to stabilize not only metals but also oxygen in various oxidation states: $O_2^+$, $O_2$, $O_2^-$. [12-13].

We report here an electrochemical study of the oxidation of silver and copper anodes in aqueous basic media and in molten hydroxide fluxes with different electrolytes added [14]. In addition to the characterization of the oxidation products obtained, this study also contributes to establish the electrochemical stability of the anodes in the strongly basic media used.

2. EXPERIMENTAL

The hydroxide used as electrolyte, NaOH, Panreac for analysis (97%) and KOH, Merck, ultrapure (85.90%) were used as purchased. In the molten hydroxide fluxes 20 g of the equimolar eutectic NaOH (8.32g) / KOH (11.68g) were fused at 185°C (the theoretical melting point of the mixture is 170°C) in a Teflon crucible exposed to the atmosphere in a sand bath. The hydroxide temperature was controlled with a Teflon-coated thermocouple. The hydroxides were heated for 12 hours before conducting each experiment, in order to assure a reproducible low level of water in the system.

The working electrodes used were silver foil (Goodfellow, 99.99%, 0.5cm$^2$, 0.125mm thick) or copper foil (Goodfellow, 99.9%, 1cm$^2$, 0.125mm thick). A platinum wire (Goodfellow, 99.99%, 0.5mm diameter) was used as counter-electrode, and silver (Goodfellow, 99.9%, 0.5cm$^2$, 0.125mm thick) or gold foil (Goodfellow, 99.9%, 0.5cm$^2$, 0.125mm thick) were used as pseudo-reference electrodes. These were used both in molten hydroxide and in aqueous solutions for comparison (instead of more conventional HgO electrodes, which could only be used in the experiments carried out in aqueous media). Some experiments included the addition of 0.02g of CuO (maximum amount soluble in molten hydroxides under the conditions of our experiments) or 0.2g of AgNO$_3$(Panreac, for analysis, 99.8%).

Electrochemical studies were performed in a three-electrode, one-compartment-cell geometry. An EG&G PAR 273A and 263A potentiostat were used for these studies.

X-Ray Diffraction patterns of the deposits were measured with a Rigaku “Rotaflex” Ru-200B diffractometer (CuKα wavelength 1.5418Å)

3. RESULTS AND DISCUSSION

Electrochemical studies of silver and copper anodes were carried out both in aqueous basic solutions or molten hydroxides. A comparison of the behavior of both metals in those media is presented in this paper. In an attempt to get more easily comparable results, the same reference electrodes used in molten hydroxides (gold and silver) were also used in basic aqueous media instead of the more conventional HgO reference electrode. Yet, these gold or silver electrodes cannot be...
strictly considered reference electrodes. They have been used in earlier studies as pseudo-reference electrodes [15] and they are certainly useful in cases like the present one, with corrosive molten hydroxide electrolytes, but their lack of a reliable redox couple which could be applied both to aqueous and molten media makes very difficult a direct meaningful comparison of the results in both solvents. On the other hand, the use of these pseudo-reference electrodes has been very valuable for controlling potentials in a particular medium and carrying out electrolysis experiments which could be correlated with the corresponding cyclic voltammogram curves in the same medium.

3.1. Silver anodes

Figure 1 shows a cyclic voltammogram of a silver electrode in 3M NaOH aqueous solution. Two oxidation and the corresponding two reduction waves can be clearly distinguished. Chronoamperometric oxidations were carried out at each of the oxidation potentials (600mV and 900mV vs. Au) for a period of 1-3 hours leading to the formation of smooth dark deposits on the electrode. The resulting compounds were characterized by X-Ray diffraction confirming the presence of cubic Ag$_2$O in the first oxidation step and a mixture of Ag$_2$O and AgO in the last one (see figure 2). It should be noted that AgO is not a Ag(II) phase but has been shown to contain Ag(I) and Ag(III) by means of neutron diffraction [16] and EXAFS data [17-18].

Similar experiments were carried out in a mixture of fused NaOH/KOH. The cyclic voltammogram resulting from this medium is remarkably different from that obtained in aqueous solutions (Figure 3). First of all, in fused hydroxides the redox waves are shifted towards less positive potentials. This must be due to the more basic medium affecting the formation of oxide species. Secondly, in the molten hydroxide CV only a couple of waves are observed, although the oxidation wave shows in some experiments a shoulder, better resolved than in aqueous electrolytes which could correspond to the formation of silver suboxide films. Finally, whereas the CV in aqueous media shows a self-limiting oxidation process, corresponding to the formation of a passivating layer, the oxidation of silver anodes in molten hydroxides is not self-limited (a complete oxidation of the electrode is possible). This different behavior could be explained in terms of the higher temperature in molten hydroxides (which would facilitate oxygen diffusion through the oxide layer) but also with the different nature of the oxide at the electrode surface.

In order to assign the anodic wave to the corresponding process chronoamperometric oxidations of the silver electrodes at 150 mV vs. Au were carried out and the deposits formed were analyzed by X-Ray diffraction (Figure 4). The microcrystalline deposit obtained corresponded to a fairly pure new phase that crystallizes in the orthorhombic system (a=3.12, b=9.96, c=12.79 Å). Although these lattice parameters are metrically very close to the unit cell of the oxide Ag$_2$O$_3$ (JCPDS 400909), it should be noted that the diffraction patterns are radically different both concerning the number, position and intensity of the peaks. Indeed, the diffraction pattern of the new phase cannot be assigned to any known compound, and its composition and crystal structure are currently under study. Microcrystals suitable for single-crystal X-Ray diffraction measurements have been obtained although the crystal structure solution seems to be hindered by twinning or pseudosymmetry problems.

3.2. Copper anodes

Electrochemical oxidations were carried out on copper electrodes. As for silver electrodes, those studies were made both in basic aqueous solution and molten NaOH/KOH.
A typical cyclic voltammogram of a copper electrode in 3M NaOH (aq) showing basically two redox waves is presented in figure 5. The first wave at -0.1 Volts vs. Au corresponds to the two-electron oxidation of Cu directly to CuO, avoiding the formation of Cu(II) as previously reported [19]. At a potential near 600mV there is a second smaller wave which we assign to the oxidation to Cu(III) by comparison with the results obtained in molten hydroxides (see below). So far we have not been able to isolate this oxidized compound. In the same cyclic voltammogram the corresponding reduction waves can also be observed, indicating a reversible nature for the oxidation reactions studied. First the Cu(III) to Cu(II) process is detected as a weak wave at 0.6 volts vs. Au. Then, instead of a single two-electron process, two reduction waves only partially resolved are observed, which indicate the possibility of stabilizing Cu(I) species under these conditions.

Similarly to what happened with silver anodes, the cyclic voltammograms of copper in molten hydroxides are remarkably different from the ones obtained in basic aqueous solutions. In this case, the molten hydroxides medium is very useful for separating the different processes as can be seen in figure 6, where three pair of clear redox waves are resolved. The first oxidation step must correspond to the formation of cuprite (CuO). Further oxidation of this system in molten hydroxide gets complicated by dissolution processes not present in aqueous basic media. In this respect, the solubility of CuO in molten hydroxides, is well known and we have repeatedly ascertained it in our work. Furthermore, a chronoamperometric experiment with a copper anode in molten hydroxides (17 hours at -0.050 vs. Ag) resulted in the complete dissolution of the anode. Thus, the anodic wave centered at ca. 0.0 Volt shown in figure 6 must correspond to a process of oxidation taking place with simultaneous dissolution of the Cu(II) oxide formed. The shoulder at lower potentials could correspond to the initial oxidation of CuO to CuO, which would get dissolved and then a two-electron oxidation could take place (Cu(II) to Cu(II) dissolved). The dissolution process is also confirmed by the observation of a blue coloration of the molten flux at this point, which corresponds to well known Cu(OH)\textsuperscript{2-} and Cu(OH)\textsuperscript{4+} species. Further oxidation of the system at 0.5 Volts vs. Ag is enough to lead to the formation of NaCuO\textsubscript{2}, a Cu(III) compound that has been previously reported.[11, 20-22] In this high oxidation state copper does not form a binary oxide, and Cu(III) only gets stabilized through the formation of ternary compounds with alkaline metals.[23] The nature of this oxide, which we have also obtained on Pt electrodes from Cu(II)-containing hydroxide melts has been confirmed by powder x-ray diffraction.

Table I summarizes the electrochemical processes studied, including the values of redox potentials and products isolated in each case.

<table>
<thead>
<tr>
<th>ANODE</th>
<th>MEDIA</th>
<th>OXIDATION POTENTIAL</th>
<th>REDUCTION POTENTIAL</th>
<th>PRODUCT</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ag</td>
<td>NaOH+aq.3M</td>
<td>0.56 V vs. Au</td>
<td>0.25 V vs. Au</td>
<td>Ag/Ag2O</td>
</tr>
<tr>
<td>Ag</td>
<td>NaOH+aq.3M</td>
<td>0.62 V vs. Au</td>
<td>0.56 V vs. Au</td>
<td>Ag2O / AgO</td>
</tr>
<tr>
<td>Ag</td>
<td>Fused NaOH/KOH</td>
<td>0.15 V vs. Au</td>
<td>0.027 V vs. Au</td>
<td>Ag/Ag2O</td>
</tr>
<tr>
<td>Cu</td>
<td>NaOH+aq.3M</td>
<td>-0.8 V vs. Au</td>
<td>-0.89 V vs. Au</td>
<td>Cu/CuO</td>
</tr>
<tr>
<td>Cu</td>
<td>NaOH+aq.3M</td>
<td>0.65 V vs. Au</td>
<td>0.299 V vs. Au</td>
<td>CuO / CuO(III) (not isolated)</td>
</tr>
<tr>
<td>Cu</td>
<td>Fused NaOH/KOH</td>
<td>-0.56 V vs. Au</td>
<td>-0.48 V vs. Ag</td>
<td>Cu/CuO</td>
</tr>
<tr>
<td>Cu</td>
<td>Fused NaOH/KOH</td>
<td>-0.015 V vs. Ag</td>
<td>-0.22 V vs. Ag</td>
<td>CuO/CuO (CuII in solution)</td>
</tr>
<tr>
<td>Cu</td>
<td>Fused NaOH/KOH</td>
<td>0.59 V vs. Ag</td>
<td>0.26 V vs. Ag</td>
<td>CuII in solution / NaCuO</td>
</tr>
</tbody>
</table>

A second round of experiments included the addition of copper or silver salts for the oxidation of silver and copper anodes respectively. The purpose of these additions was to test the possibility of obtaining mixed silver copper oxides as the one we recently reported[24]. Nevertheless, the presence of these additional ions in the electrolyte had no significant effect on the oxides obtained.

4. CONCLUSIONS

Electrochemical oxidation of silver or copper anodes in strongly basic media is a straightforward way to prepare films of their oxides. In comparison with more widely studied experiments in basic aqueous media, the use of an eutectic NaOH/KOH flux strongly affects the redox behavior of these metals and provides an opportunity to extend the range of phases that can be obtained. This is emphasized by the isolation of an hitherto unknown silver oxide electrocrystallized from that eutectic, the crystal structure of which is presently under investigation.

ACKNOWLEDGMENTS

This work was funded by the MCyT(Spain), Plan Nacional de Materiales (MAT2001-1709-C04-01 and MAT2002-04529-C03, MAT96-1057-c02-01and PGC PB98-0491). We also thank the Ministry of Education and Culture (Spain) for a predoctoral fellowship awarded to EMT-R.
REFERENCES


Recibido: 1.2.03
Aceptado: 30.11.03