Parameters of Oxide Films Anodically Formed on Ag-Zn Alloys with Different Concentrations of Vacancy Defects in the Surface Layer

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Abstract
The nature and the properties of oxide films anodically formed on metals and alloys depend on the chemical composition and energetic state of the electrode surface. This study was aimed at revealing the composition and the parameters of oxide films formed in 0.1 M KOH on silver-zinc alloys (up to 30 at.% of Zn) with different concentrations of vacancy defects in the surface layer. The enhanced concentration of vacancy defects in the alloys surface layer was created by preliminary selective dissolution of zinc at different potentials in 0.01 M HNO₃ + 0.09 M KNO₃. These alloys were moved into 0.1 M KOH for the formation of oxide films. The composition of films was monitored by cathodic voltammetry. The film thickness was calculated from the anodic coulometry taking into account the current efficiency. The latter was determined by cathodic coulometry. The morphology of the electrode surface was monitored by scanning electron microscopy (SEM).

It was revealed that the concentration of superequilibrium vacancies formed in the surface layer depended on the potential of the selective dissolution of zinc from the alloy. The peak of Ag(I) oxide reduction was registered on the cathodic voltammograms of the alloys after their anodic oxidation. The current efficiency of oxide formation on Ag-Zn alloys was less than 100%. It decreased with zinc concentration in the alloys. The thickness of Ag(I) oxide on alloys did not exceed 25 nm. SEM images showed the uniform distribution of oxide particles along the electrode surface. The shape of the particles was close to spherical.

The Ag(I) oxide was the main product of oxidation of silver-zinc alloys (up to 30 at% of Zn) with different vacancy defects of the surface layer. The current efficiency and the thickness of Ag(I) oxide films formed on alloys with an increased concentration of vacancy defects were less as compared with the current efficiency and the thickness of Ag(I) oxide films formed on pure silver. However, these parameters were higher as compared with the parameters of Ag(I) oxide films formed on alloys with an equilibrium concentration of vacancy defects. It was found that the diameter of the Ag(I) oxide particles decreased, and their quantity on the unit of the electrode surface increased with an increase of the concentration of vacancy defects in the alloys surface layer.

Keywords: silver-zinc alloys, Ag(I) oxide, selective dissolution, anodic oxide formation.


1. Introduction
Silver and zinc oxides, in particular nanosized ones, are considered to be functional materials with the desired properties [1–4]. Structures of the oxide/oxide [5, 6] or oxide/metal type [7–10] play a decisive role in anode and corrosion processes. They are also used in modern semiconductor technologies and in the manufacture of sensors. The scope of application for oxide structures is determined by a set of their main parameters, depending on the method and conditions of their preparation.

One of the possible ways to obtain the oxide phase and control its parameters is the

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anodic oxidation of homogeneous alloys with different contents of their components. During the transition from individual metals to alloys, the composition of oxidation products and the structure of the oxide film become more complicated [11–13]. In addition, a number of side electrochemical reactions can occur at the potentials of oxide growth; therefore, data on the current efficiency of the anodic oxide formation process are required for the correct evaluation of its thickness. Finally, during the anodic oxidation of alloys, the possibility of their selective dissolution (SD) should be taken into account, as a result of which the electronegative component predominantly passes into the solution, and the surface zone of the alloy is enriched with the electropositive component and super-equilibrium vacancies [14–17]. Due to a significant increase in the concentration of vacancies in comparison with the equilibrium concentration, the energy saturation of the system as a whole also increases, which can be reflected both in the kinetics of further oxide formation and in the properties of anodically formed oxides. The concentration of vacancy defects in the surface layer of the alloy can be controlled by changing the conditions of the SD.

The goal of this study was the calculation of the current efficiency of the anodic oxide formation and determination of the composition and thickness of oxide films anodically formed in 0.1 M KOH on silver-zinc alloys with different concentrations of vacancy defects in the surface layer.

2. Experimental

Homogeneous polycrystalline alloys were made of silver and zinc (metal purity of 99.99 at%) in vacuum quartz ampoules with exposure at a temperature of 1273 K for at least two hours and their subsequent slow cooling. Predicted zinc concentration $N_{\text{Zn}}$ was from 5 to 30 at% (Table 1). The phase (α-phase) and chemical compositions of the alloys were confirmed by X-ray diffractometry and energy-dispersive microanalysis [18]. Samples were mounted in epoxy resin in such a way that the working surface with the geometric area $S_g$ from 0.4 to 0.5 cm$^2$ remained open (Table 1). The preparation of the surface of the electrodes for the experiment included mechanical processing on abrasive paper, polishing on suede and degreasing with isopropyl alcohol.

Electrochemical studies were carried out using an IPC-Compact potentiostat (Institute of Physical chemistry and Electrochemistry, RAS, Moscow). Anodic oxide formation and determination of current efficiency were performed in 0.1 M KOH (pH 12.89), prepared on double distilled water from a chemically pure reagent and deaerated by sparging with chemically pure argon. The sequence of stages was as follows:

1. Cathodic polarization in an alkaline solution for 300 s at a potential $E_c = -1.1$ V for reduction of oxide traces and standardization of the surface.

2. Potential-controlled and charge-controlled anodic synthesis of Ag(I) oxide of a specified thickness (34 nm under the assumption of 100% current efficiency of the anodic formation of $\text{Ag}_2\text{O}$) at potential $E_a = 0.56$ V. This value was chosen based on the results of anodic voltammetry and corresponds to the maximum growth rate of Ag(I) oxide on silver in 0.1 M KOH [18].

3. Change of alkaline solution in a cell in a sealed argon atmosphere for the removal of soluble oxidation products of silver or zinc.

4. Potentiodynamic cathodic polarization at a scan rate of 1 mV s$^{-1}$ from open-circuit potential $E_0$ to $E_c$.

In a separate series of experiments, after the completion of stage 2, the dependence of the open-circuit potential $E_0$ from time was recorded for 20 min, after the solution was changed (stage 3) and the cathodic potentiodynamic reduction of the oxidation products remaining on the electrode surface was performed (stage 4).

For the determination of the role of super-equilibrium vacancies before anodic oxide formation, Table 1. Symbols and the geometric surface of Ag-Zn alloys (α-phase) with different concentration of zinc $N_{\text{Zn}}$

<table>
<thead>
<tr>
<th>$N_{\text{Zn}}$, at%</th>
<th>5</th>
<th>10</th>
<th>15</th>
<th>20</th>
<th>30</th>
</tr>
</thead>
<tbody>
<tr>
<td>Symbol</td>
<td>Ag5Zn</td>
<td>Ag10Zn</td>
<td>Ag15Zn</td>
<td>Ag20Zn</td>
<td>Ag30Zn</td>
</tr>
<tr>
<td>Surface, cm$^2$</td>
<td>0.40</td>
<td>0.44</td>
<td>0.43</td>
<td>0.42</td>
<td>0.48</td>
</tr>
</tbody>
</table>
formation, the potential-controlled and charge-controlled SD of Ag15Zn and Ag30Zn alloys in 0.01 M HNO₃ + 0.09 M KNO₃ (pH 2.16) solution, prepared on double distilled water from chemically pure reagents and deaerated by sparging with chemically pure argon was performed. The sequence of stages was as follows:

1. Immersion of the working electrode for 240 s in acid solution under protection potential $E_c = –1.1$ V, eliminating the ionization of both components for the reduction of traces of oxides and standardization of the surface.

2. Anodic selective dissolution for 60 s at potentials $E_1^{SD} = 0.4$ V, $E_2^{SD} = 0.5$ V or $E_3^{SD} = 0.6$ V. Potential values were selected based on the results of anodic voltammetry (Fig. 1) and correspond to the thermodynamic stability region of the electropositive component of the alloy.

On the alloys obtained as a result of stages 1 and 2, the oxide was formed with the subsequent determination of the current efficiency (stages 1–4).

The current efficiency $\Psi$ of the anodic oxidation process in all experiments was calculated as the ratio of the cathodic charge of film reduction $Q_c$ to the anodic charge of its formation $Q_a$:

$$\Psi = \frac{Q_c}{Q_a}. \quad (1)$$

The determined current efficiency was used for the calculation of the thickness of the formed oxide layer:

$$L = \frac{\Psi M q_a}{z F \rho}, \quad (2)$$

where $M$ and $\rho$ are the molar mass and density of the oxide; $q_a$ is the density of the anodic charge; $z$ is the number of electrons involved in oxide formation; $F = 96485$ C mol⁻¹ is Faraday constant.

The control of surface morphology and composition was carried out using scanning electron microscope JSM 6380LV with microanalysis system INCAx-sight 250 (JEOL, Japan)⁷. Images were obtained using the secondary electron mode with an accelerating voltage of 20 kV.

3. Results and Discussion

3.1. Anodic oxidation of Ag–Zn alloys without preliminary selective dissolution

On chronoamperograms obtained in deaerated 0.1 M KOH on silver and Ag–Zn alloys at a potential $E = 0.56$ V, a sharp and then smoother decrease of current was observed (Fig. 2a). At times greater than 10–20 s, the current density reached an almost constant value.

According to (2), for the formation of an Ag₂O film with an estimated thickness of 34 nm, assuming 100% current efficiency, it is necessary to accumulate a charge density of 20 mC cm⁻². Fig. 2a shows that with an increase in the initial concentration of zinc in the alloy, the polarization time required for the accumulation of a given charge density decreased.

With times not exceeding 5–10 s, linear sections in the coordinates, criterion for diffusion-limited processes were observed (Fig. 2b). A similar situation observed for pure silver and its alloys with gold allowed the authors of the study [19] to draw a conclusion about the solid-phase-diffusion control of the anode process. It should be assumed that on silver-zinc alloys, the anode process is controlled by mass transfer along the phase of the growing oxide as well.

In the general case, the recorded current and charge characterize not only the formation of Ag₂O oxide

$$2Ag + 2OH^- = Ag_2O + H_2O + 2 e^-,$$  \quad (3)

but also the transition of silver ions into the solution:

$$Ag + 2OH^- = Ag(OH)_2^- + e^-.$$  \quad (4)

In addition, zinc oxidation, accompanied by the formation of an oxide or hydroxide film on the...
surface of the electrode and ions in the electrolyte solution is possible:

\[
\text{Zn} + 2\text{OH}^- = \text{ZnO} + \text{H}_2\text{O} + 2\text{e}^-; \quad (5)
\]
\[
\text{Zn} + 2\text{OH}^- = \text{Zn(OH)}_2 + 2\text{e}^-; \quad (6)
\]
\[
\text{Zn} + 4\text{OH}^- = \text{Zn(OH)}_4^{2-} + 2\text{e}^-; \quad (7)
\]

In solutions with a high pH value (> 10), soluble products are formed:

\[
\text{ZnO} + \text{OH}^- + \text{H}_2\text{O} = \text{Zn(OH)}_3^-; \quad (8)
\]
\[
\text{ZnO} + 2 \text{OH}^- + \text{H}_2\text{O} = \text{Zn(OH)}_4^{2-}. \quad (9)
\]

3.2. Measurement of open-circuit potential

Certain information regarding the nature and electrochemical behaviour of the oxide or hydroxide phase formed in 0.1 M KOH can be obtained using chronopotentiometry in open-circuit mode, carried out after the end of potentiostatic polarization (Fig. 3). When \( N'_{\text{Zn}} \leq 10 \text{ at.\%} \) the open-circuit potential \( E_0 \) of oxidized alloys varies slightly over time and remains close to the values characteristic of Ag(I) oxide formed on silver. When concentration of zinc in the alloy increases, the potential \( E_0 \) shifts to less positive values. At the same time, a rather sharp decrease of potential over time was observed. Chemical dissolution of Ag(I) oxide can be the reason for a decrease in \( E_0 \):

\[
\text{Ag}_2\text{O} + 2 \text{OH}^- + \text{H}_2\text{O} = 2 \text{Ag (OH)}_2^-.
\]

3.3. Selective dissolution of Ag-Zn alloys

Chronoamperograms obtained on Ag15Zn and Ag30Zn alloys in 0.01 M HNO_3 + 0.09 M KNO_3 at various \( E^{\text{SD}} \) potentials had a similar shape. Typical current transients are shown in Fig. 4a for Ag15Zn alloy. During the anodic polarization, the current dropped to almost zero, reflecting the almost complete dissolution of Ag(I) oxide.
complete dezincification of the surface layer of the alloy. The presence of linear sections in the Cottrell coordinates (Fig. 4b) allows applying SD chronoamperometry theory for the calculation of the diffusion coefficient and concentration of vacancies [15, 16]:

\[
I(t) = z_A F N_A D_A^{1/2} S_g \left[ 1 - e^{\frac{z_A F \eta}{RT}} \right].
\]  

(11)

Here: \( I \) – current, \( t \) – time, \( z_A \) – the number of electrons involved in the oxidation of the electronegative component, \( N_A \) – volume concentration of atoms of component A in the alloy, \( D_A = D_v N_v \) – diffusion coefficient of the atoms of component A in the surface layer of the alloy, \( D_v \) – diffusion coefficient of vacancies, \( N_v \) – concentration of vacancies, \( S_g \) – geometric surface area of the electrode, \( V_m \) – molar volume of the alloy, \( R = 8.314 \text{ J K}^{-1} \text{ mol}^{-1} \) – universal gas constant, \( T = 298 \text{ K} \) – temperature, \( \eta \) – overvoltage.

The values of equilibrium vacancy concentrations equal to \( 2.55 \times 10^{-5} \) and \( 1.05 \times 10^{-5} \) at.\% in Ag15Zn and Ag30Zn alloys, respectively, were obtained by thermodynamic calculations using data on the vacancy formation energy and their diffusion coefficients in individual metals [16].

Zinc diffusion coefficients \( D_v \) calculated according to (11) from the slopes \( i_t^{-1/2} \)-dependencies for each of the alloys increased with increase in \( E^{SD} \) (Table 2). According to the solid-phase-diffusion mechanism of SD, the reason for this is an increase in the concentration of superequilibrium vacancies (Table 2), e.g., the concentration of vacancy defects in the surface layer.

### 3.4. Anodic oxidation of Ag-Zn alloys after selective dissolution

Chronoamperograms obtained in 0.1 M KOH on alloys with different concentration of vacancy defects in the surface layer were similar in shape to chronoamperograms obtained on alloys without preliminary SD (not shown in the figures). Linear sections were also preserved in the criterion coordinates, indicating the transport nature of the limitations of the anode process in

| Table 2. Diffusion coefficients of zinc \( D_v \) and concentration of superequilibrium vacancies \( N_v \) in the surface layer of Ag-Zn alloys at different potentials \( E^{SD} \) |
|------------------|----------------|----------------|----------------|
| **Alloy**       | **\( E^{SD}, \text{V} \)** | **\( D_v \times 10^{18}, \text{cm}^2 \text{s}^{-1} \)** | **\( N_v \times 10^4, \text{at.\%} \)** |
| Ag15Zn          | 0.40           | 3.8            | 1.0            |
|                 | 0.50           | 8.1            | 2.1            |
|                 | 0.60           | 12.2           | 3.2            |
| Ag30Zn          | 0.40           | 0.7            | 0.09           |
|                 | 0.50           | 1.1            | 0.1            |
|                 | 0.60           | 2.7            | 0.4            |

Fig. 4. Current transients of Ag15Zn in original (a) and Cottrell coordinates (b) at different \( E^{SD} \) in 0.01 M HNO\(_3\) + 0.09 M KNO\(_3\).
an alkaline environment. Most likely, the nature of the anodically formed products did not undergo significant changes during the transition from alloys without SD to alloys after SD.

3.5. The cathodic reduction of insoluble products of alloys oxidation

The cathodic voltammograms obtained on silver and alloys with an equilibrium concentration of vacancies after the end of anodic oxidation and change of solution in the cell are shown in Fig. 5a.

Considering the equilibrium potentials of processes (3)–(7), we can conclude that the maximum cathode current at potentials $E_{Ag(I)} = 0.32–0.36$ V corresponds to the reduction process of Ag(I) oxide. A sharp increase in the cathode current associated with the onset of hydrogen emission was observed at potentials near $–1.0$ V, which was significantly higher than the equilibrium potentials of formation/reduction of zinc oxide or hydroxide ($–1.3$ V). Thus, if the products of zinc oxidation were also present in the oxide film on the alloys, then it was impossible to detect their presence using cathodic voltammograms.

When the cathodic reduction was preceded by a 20-minute stage of potential measurement in open-circuit mode, then the shape of voltammograms in general did not change (Fig. 5b). However, the peak of reduction of the Ag(I) oxide sharply decreased in amplitude, and it was absent on the Ag20Zn and Ag30Zn alloys. Indeed, on these alloys $E_0$ value was below $E_{Ag(I)}$ Potentials of the reduction of Ag$_2$O. It should be assumed that the major volume of the formed oxide was already chemically dissolved according to (10) over the period of recording of the open-circuit potential.

If anodic oxidation was preceded by SD of alloys (stages 1‘ and 2‘), then $E_0$ and $E_{Ag(I)}$ potentials on cathodic voltammograms were shifted to more positive values characteristic for pure silver (Fig. 5c). The cathodic peak of reduction of the Ag(I) oxide was reliably fixed, and the absolute values of the maximum current were much higher than on alloys not subjected to preliminary SD.

5.6. The current efficiency of the anodic formation of Ag(I) oxide

The results of calculating the current efficiency according to (1) and the thickness of the Ag$_2$O films corrected to this value by (2) formed on silver and alloys of the Ag-Zn system after various stages of research are summarized in Tables 3 and 4. Cathodic charge $Q_c$ characterizing the reduction of Ag(I) oxide was calculated according to the results of cathodic voltammetry (Fig. 5a, b, c) in the potential region from $E_0$ up to $E = 0.20$ V.

The current efficiency of anodic oxide formation on alloys without SD (Table 3) was lower than on silver, and decreased with increasing zinc concentration in the alloy. Accordingly, the thickness of the Ag(I) oxide also decreased from 32 to 5.8 nm during the transition from Ag to Ag30Zn. The decrease in current efficiency with increasing zinc concentration can be explained by an increase in the porosity of the oxide film. As a result, the possibility of dissolution of silver through the pores increased, the fraction of the current used for the formation of silver ions entering the solution

![Fig. 5. Cathodic voltammograms of silver and Ag-Zn alloys after the anodic polarization at $E_a = 0.56$ V and change of the solution: (a) without an open-circuit stage; (b) after the open-circuit stage and after the preliminary selective dissolution of Ag15Zn (c)](image-url)
increased, and the fraction of the current used for the formation of the film decreased.

After a 20 min exposure in open circuit mode, the current efficiency values of $\Psi'$ calculated from (1) and oxide thickness $L^*$ determined by (2) further decreased (Table 3), which confirms the assumption of the chemical dissolution of silver oxide in an alkaline medium.

The current efficiency of anodic oxide formation on alloys subjected to preliminary SD (Table 4) was lower than that on silver, but higher than that on alloys without preliminary SD. As the potential of SD increased, i.e., increased concentration of vacancy defects in the surface layer of the alloy, the current efficiency and thickness of the Ag(I) oxide decreased.

3.7. Morphology of anodically formed films

Comparison of the results of energy-dispersive microanalysis conducted on the alloys before the start of their anodic oxidation and after its completion demonstrated an increase in the oxygen concentration in the surface layer (Table 5), which confirms the presence of an oxide layer on the surface.

The thickness of the oxide intended for SEM was increased to 120 nm, taking into account the obtained current efficiency values. Typical SEM images of the oxidized surface for Ag15Zn and Ag30Zn without SD (Fig. 6a, b) and for the same alloys, but subjected to preliminary SD at the potential $E_{SD}$ (Fig. 6 c, d) are presented. On the surface of all alloys, individual particles of the oxide phase can be observed; the particle shape was close to spherical. Depending on the initial concentration of zinc in the alloy and the conditions for preparing the surface for oxidation, the particles size and quantity on the unit of the electrode surface changed.

On alloys not subjected to preliminary SD, a decrease in the particle diameter of the formed oxide increased, and the fraction of the current used for the formation of the film decreased.

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oxide with increasing zinc concentration was revealed (Table 6). Probably this situation was associated with a decrease in the interatomic distance of Ag-Ag from 0.409 to 0.403 nm with increasing $N_{Zn}$ from 0 to 30 at.% [23]. At the same time, an increase in the oxide particles quantity on the unit of the electrode surface was observed.

Upon transition to alloys subjected to preliminary SD, the particles diameter decreased, and the particles quantity on the unit of the electrode surface increased (Table 6). In this case, a decrease in the particle size of the Ag(I) oxide probably was associated with an increase in the concentration of vacancy defects in the surface layer of alloys. It is not possible to trace a clear trend of decreasing oxide particles diameter or increasing their quantity on the unit of the electrode surface with increasing zinc concentration on alloys after SD.

### 4. Conclusions

1. Ag(I) oxide is the main product of the anodic dissolution in an aqueous alkaline medium of homogeneous polycrystalline alloys of the Ag-Zn system ($\alpha$-phase), as well as these alloys, but subjected to preliminary selective dissolution of zinc from their surface layer in an acidic medium.

2. Silver-zinc alloys with enriched to almost pure silver, but with a structurally-disordered surface layer, are characterized by increased current efficiency of the anodic oxide formation process, as well as a higher thickness of Ag(I)

#### Table 6. Diameter $d$ of Ag$_2$O particles and their quantity $P$ on the unit of the electrode surface

<table>
<thead>
<tr>
<th>Stage</th>
<th>Parameter</th>
<th>Electrode</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Ag</td>
</tr>
<tr>
<td>Without SD</td>
<td>$d$, nm</td>
<td>350</td>
</tr>
<tr>
<td></td>
<td>$P$, $\mu$m$^{-2}$</td>
<td></td>
</tr>
<tr>
<td>$E_1^{SD}$</td>
<td>$d$, nm</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>$P$, $\mu$m$^{-2}$</td>
<td>–</td>
</tr>
<tr>
<td>$E_2^{SD}$</td>
<td>$d$, nm</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>$P$, $\mu$m$^{-2}$</td>
<td>–</td>
</tr>
<tr>
<td>$E_3^{SD}$</td>
<td>$d$, nm</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>$P$, $\mu$m$^{-2}$</td>
<td>–</td>
</tr>
</tbody>
</table>
oxide in comparison with equilibrium alloys of the same chemical composition. At the same time, these characteristics turn out to be noticeably lower in comparison with compact polycrystalline silver.

3. Ag(I) oxide, anodically formed on silver and silver-zinc alloys (α-phase) of not only the initial composition, but also after the selective dissolution of zinc from their surface layer, is chemically unstable in an alkaline medium.

4. The diameter of the anodically formed Ag(I) oxide particles decreases, and their quantity on the unit of the electrode surface increases with an increase in the concentration of vacancy defects in the surface layer of the alloy.

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Conflict of interests
The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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