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Two-Layer Formation of Passivating Films on Cobalt in Neutral Solution

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ABSTRACT

Layer structure and thickness of passivation oxide films on cobalt have been investigated at stationary and transient states in solution of 0.30 mol dm\(^{-3}\) boric acid-0.075 mol dm\(^{-3}\) sodium borate mixture of pH 8.4 by means of chemical analysis and in situ ellipsometry combined with electrochemical measurements. A two-layer structure composed of an inner CoO and an outer Co\(_3\)-O\(_4\) is found in a potential range more positive than 0.75V (vs. the hydrogen electrode at the same solution). The nonstoichiometry changes from \(\Delta = 0\) to 0.33 with increasing anodic potential. The outer Co\(_3\)-O\(_4\) layer thickness increases linearly with potential, whereas the inner layer thickness is not a simple function of potential, but depends on the film formation procedure. A step-wise potential increase gives an initial rapid growth of the outer Co\(_3\)-O\(_4\) layer followed by a gradual thickening of the inner CoO layer. The transient film growth behavior is interpreted in terms of a field-assisted ion diffusion mechanism, and the self-diffusion coefficient in the inner CoO layer is estimated to be \(D/cm \sec^{-1} = 6.5 \times 10^{-21} \sim 2.6 \times 10^{-20}\).

The previous papers (1-4) have shown that the passive oxide film on cobalt is potential dependent in neutral boric acid-sodium borate solution of pH 8.4. At potentials more negative than 0.86V, referred to the hydrogen electrode in the same solution (HESS), a slightly soluble oxide film of CoO 2 \(\sim\) 3 nm thick is formed under stationary potentiostatic conditions (2-3). At more positive potentials a two-layered film is formed, consisting of an inner CoO and an outer Co\(_3\)-O\(_4\) layer (2-3). The nonstoichiometry \(\Delta\) of the outer layer depends on the anodic potential, changing from \(\Delta = 0\) at 1.15V to \(\Delta = 0.33\) at 1.4V. The outer layer thickness increases linearly with the potential, whereas the inner layer thickness is not a simple function of potential. In the present paper the kinetics and mechanism of electrochemical formation of two-layered oxide films on cobalt were studied by using a potential step technique associated with the ellipsometric measurements.

Experimental

The specimens were prepared from cobalt rods of 99.99% purity (Johnson-Matthey Limited) into polycrystalline sheets 0.8 mm thick with the surface area of 5 or 10 cm\(^2\). The specimen pretreatment was the same as that described in the previous papers (1-4). The electrolytes were prepared from doubly distilled water and analytical grade reagents. Anodic formation of the passive films on cobalt was performed in a mixture of 0.30 mol dm\(^{-3}\) boric acid and 0.075 mol dm\(^{-3}\) sodium borate solutions at pH 8.4. Cathodic reduction of the passive films was conducted in a mixture of boric acid and sodium borate solutions at pH 6.5. A boric acid solution at pH 5.3 and a mixture of chemical analysis and in situ ellipsometry combined with electrochemical measurements. A two-layer structure composed of an inner CoO and an outer Co\(_3\)-O\(_4\) is found in a potential range more positive than 0.75V (vs. the hydrogen electrode at the same solution). The nonstoichiometry changes from \(\Delta = 0\) to 0.33 with increasing anodic potential. The outer Co\(_3\)-O\(_4\) layer thickness increases linearly with potential, whereas the inner layer thickness is not a simple function of potential, but depends on the film formation procedure. A step-wise potential increase gives an initial rapid growth of the outer Co\(_3\)-O\(_4\) layer followed by a gradual thickening of the inner CoO layer. The transient film growth behavior is interpreted in terms of a field-assisted ion diffusion mechanism, and the self-diffusion coefficient in the inner CoO layer is estimated to be \(D/cm \sec^{-1} = 6.5 \times 10^{-21} \sim 2.6 \times 10^{-20}\).

The ellipsometer used was of horizontal type with a quarter-wave plate placed on the side of incident beam. The fast axis of the plate was fixed at a 45° orientation. Measurements were made by the standard null method at a wavelength of \(\lambda/nm = 546\) and at an incidence angle of \(\varphi_1 = 73.30^\circ\) (3, 5).

All measurements were made at 20.0° ± 0.2°C. The potential of cobalt electrodes was measured in reference to a saturated calomel electrode (SCE) and converted to a scale of the hydrogen electrode in the same solution (HESS).

The amount of cobalt ion dissolved into solutions was analyzed by a colorimetric nitroso-R-salt method.

Result

Two-layer structure of passivation films.—Figure 1 illustrates the results of ellipsometric measurements (Fig. 1a), the amount of dissolved Co(II) ion from the passive film, \(W_d\), and the decay of potential, \(\phi\), (Fig. 1b) during a galvanostatic cathodic reduction of a passive film on cobalt. The passive film was formed by 1 hr oxidation at a constant potential in the solution of pH 8.4 and the reduction was performed at a constant current in the solution of pH 6.5. The loci of polarizer azimuth \(P\) and analyzer azimuth \(A\) are directly plotted in Fig. 1a, where \(P_o\) and \(A_o\) correspond to the oxide-free reference surface of cobalt; \(P_o = 28.6^\circ\) and \(A_o = 4.43^\circ\) determine the complex refractive index of metallic cobalt, \(n_{Co} = 2.83-3.86i\) (2, 3).

The loci \(P_o\) vs. \(A_o\) exhibit two breaks at the points marked on \(Q_o^\circ\) and \(Q_{M}^\circ\), which appear to correspond to two breaks observed in \(W_d\) vs. \(Q_o\) and \(\phi\) vs. \(Q_o\) curves. In view of the previous results (3, 4), it is found that \(Q_o^\circ\) is the cathodic charge required for the nonstoichiometry \(\Delta\) of the outer Co\(_3\)-O\(_4\) layer to decrease to \(\Delta = 0\), and that \(Q_{M}^\circ\) is the charge required for the outer layer to be completely reduced to Co\(_2^+\) ions. Thus, the reaction for \(Q_o^\circ\) and the reaction for \(Q_{M}^\circ\) are respectively represented by reactions (A) and (B)

\[
\text{Co}_3\text{O}_4 + \frac{8\Delta}{3} \text{H}^+ + \frac{8\Delta}{3} e^- \rightarrow (1 - \frac{\Delta}{3})\text{Co}_2\text{O}_4 + \frac{4\Delta}{3} \text{H}_2\text{O} \quad \text{[A]} 
\]

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**References**


**Key words:** metals, anode, ellipsometry, passivity.
Theoretical 8P-SA loci calculated for two-layered films to reach that of direct one-step oxidation for 1 hr. It is required for the inner layers. More than 6 hr of oxidation at 1.24V starting from the oxide-free cobalt electrode. Figure 3 shows a 8A-SP curve observed after a potential step from 1.04 to 1.24V is applied to a passivated cobalt that has been oxidized at 1.04V (V) for 1 hr. The filled circle in Fig. 3 indicates the values of 8P and 8A observed after direct 1 hr oxidation at 1.24V (2nd step). In Fig. 1 the theoretical lines are also drawn for various thicknesses of the outer and the inner layers. Figure 2 shows the 8A vs. 8P loci during the spontaneous dissolution of a passivation film in solutions of 0.5 mol dm\(^{-3}\) NH\(_4\)OH-0.5 mol dm\(^{-3}\) NH\(_4\)Cl mixture (A) and of 0.3 mol dm\(^{-3}\) boric acid (\(\bigcirc\)). The break marked by n\(^{\text{M}}\) in the 8P-8A loci corresponds to a point where the potential decays from the first to the second wave.

\[
\text{CoO}_2 + 8H^+ + 2e \rightarrow \text{Co}^{2+} + 4H_2O \quad \text{[B]}
\]

The third stage of film reduction after Q\(^{\text{M}}\) is the reduction of the inner CoO layer. To determine the complex refractive indexes of the layers, the best fit approach was made between the theoretical \(\delta \Phi\) vs. \(\delta \Psi\) loci for different complex refractive indexes and the experimental ones by taking into account the approximate film thickness estimated from the film charge in anodic oxidation and in cathodic reduction. The complex refractive indexes thus determined for the outer layer are \(n_{\text{CoO}_3} = 3.2-0.95i\) and \(n_{\text{CoO}_4} = 3.2-0.5i\), and for the inner layer \(n_{\text{CoO}} = 2.3-0.1i\) (2, 3). In Fig. 1 the theoretical lines are also drawn for various thicknesses of the outer and the inner layers.

Figure 2 shows the 8A vs. 8P loci during the spontaneous dissolution of the passive film in two different solutions of 0.5 mol dm\(^{-3}\) NH\(_4\)OH-0.5 mol dm\(^{-3}\) NH\(_4\)Cl mixture at pH 9.6 and of 0.30 mol dm\(^{-3}\) boric acid at pH 5.3, where no composition change of the solutions of 0.5 mol dm\(^{-3}\) NH\(_4\)OH-0.5 mol dm\(^{-3}\) NH\(_4\)Cl mixture (A) and of 0.3 mol dm\(^{-3}\) boric acid (\(\bigcirc\)). The break marked by n\(^{\text{M}}\) in the 8P-8A loci corresponds to a point where the potential decays from the first to the second wave.

Transient behavior.—To elucidate the layer growth, ellipsometric transient measurements of the film were carried out in two-step oxidation, in which, after renewal of the solution containing no dissolved cobalt ion, a potential step was applied to a preoxidized cobalt electrode. Figure 3 shows a 8A-8P curve observed after a potential step from 1.04 to 1.24V is applied to a passivated cobalt that has been oxidized at \(\Phi = 1.04V\) (V) for 1 hr. The filled circle in Fig. 3 indicates the values of 8P and 8A observed after direct 1 hr oxidation at 1.24V starting from the oxide-free cobalt surface. It is noted that more than 6 hr of oxidation is required for the 8P-8A value of two-step oxidation to reach that of direct one-step oxidation for 1 hr. By superimposing on the results of Fig. 3 the theoretical 8P-8A loci calculated for two-layered films with the complex refractive indexes, \(n_{\text{CoO}} = 2.3-0.1i\) and \(n_{\text{CoO}_4} = 3.2-0.5i\), as a function of layer thickness, the thickness of the both layers can be evaluated as a function of time during oxidation. In Fig. 4 and 5 the thickness change thus estimated is plotted together with the current density (C.D.) change. In Fig. 4 the thickness for 1.24V oxidation at which gaseous oxygen evolves on the electrode was estimated using the complex refractive index of n\(^{\text{M}}\) = 3.2-0.5i for the outer CoO\(_3\) layer. It is seen that initially the outer layer increases accompanying no appreciable growth of the inner layer and that the inner layer begins to thicken after the outer layer grows to a certain thickness comparable to the thickness formed by direct one-step oxidation. It is noted that the inner layer formed by two-step oxidation is thinner than that formed by direct one-step oxidation. This is due to the anodic deposition of a Co(III) film from Co(II)
ions in aqueous solution produced prior to the film formation, as is shown later.

When the potential is stepped down (Fig. 5), an initial cathodic current causes a decrease of the outer layer thickness, producing a slight increase of the inner layer thickness, and then a definite decrease of the inner layer thickness occurs toward a steady-state value of 4.0 nm without yielding any change in the outer layer thickness after the C.D. turns to be anodic.

The preferential growth of the outer layer, which occurs in an initial period of potential step oxidation (Fig. 4), also takes place during galvanostatic oxidation, as shown in Fig. 6. In this case the anodic C.D. of 1 μA cm⁻² was applied to a passivated cobalt electrode that has been oxidized at Φ = 1.03V for 1 hr. Before the galvanostatic oxidation, the film has consisted of the inner layer of 1.1 nm and the outer layer of 0.9 nm. Application of a small anodic current first causes a rapid increase of the outer layer with a decrease of the inner layer and then the film of a single Co₃O₄ layer (n = 3.2-0.5i) with a linear rise of potential.

Finally, the film transforms to CoO₂ (n = 3.2-0.95i) without producing any increase of the inner layer.

In Fig. 7 the change in the ΔA vs. ΔP loci is illustrated for the second step of oxidation at different potentials in the range of two-layer formation, starting from a passivated cobalt electrode that has been oxidized at 1.04V for 1 hr. The ΔP-ΔA value corresponding to the film formed by direct one-step oxidation for 1 hr lies on the extrapolated growth line of two-step oxidation. At potentials more positive than 1.34V the locus of the second step oxidation even after 22 hr (ΔP does not reach the value of the passivated electrode directly oxidized in one step for 1 hr (●). The second step oxidation at 1.54V gives rise to transformation of Co₃O₄ to CoO₂ in an initial
period, as suggested by a change in the complex refractive index from $n = 3.2-0.5i$ to $3.2-0.95i$.

Comparisons of the thickness and the final C.D. are made in Fig. 6 between direct 1 hr oxidation and two-step 22 hr oxidation. In this figure the thickness and the C.D. in the potential region where the film of a single CoO layer is formed is also plotted. In the potential region for a two-layer film formation the final anodic C.D. in two-step oxidation is one-tenth times as small as that in direct one-step oxidation. Difference in C.D. between the potential regions of CoO and CoO$_2$ formation can be seen in two-step oxidation, but becomes indistinct in direct one-step oxidation. The thickness of the outer layer does not depend on whether the film is formed by one-step or two-step oxidation and increases nearly linearly with potential except for the film formed above 1.4V. The inner layer thickness, however, differs in different oxidation procedures. It appears that the inner layer thickness of two-step oxidation is larger at potentials below 1.2V and smaller at potentials above 1.3V than that of direct one-step oxidation. From the above results it follows that the thickness of the outer layer is determined only by the oxidation potential, whereas the thickness of the inner layer is dependent on the oxidation procedure.

Anodic deposition of Co(II) ion in solution.—The previous work (1) has shown that the amount of Co(II) ion dissolved in solution was 0.5 $\mu$g cm$^{-2}$ in 5 sec oxidation and 0.03 $\mu$g cm$^{-2}$ in 1 hr oxidation at constant potential 1.34V starting from the film-free surface. This means that the Co(II) ion dissolved in solution in the initial period of anodic passivation is anodically deposited on the passivated cobalt. Another evidence on the anodic deposition is given from an experiment in which Pt electrode is potentiostatically oxidized for 1 hr in the boric acid-sodium borate solution of pH 8.4 containing Co(II) ion of concentration 1.03 mg dm$^{-3}$. Figure 9 shows the amount of Co(II) ion anodically deposited from the solution to form a cobalt oxide film on Pt, which was estimated from the decrease of Co(II) ion concentration in the solution after potentiostatic 1 hr oxidation at constant potential. The amount of Co(II) deposited was also evaluated from the amount of Co(II) ion dissolved from the film on Pt by galvanostatic reduction at 10 $\mu$A cm$^{-2}$ in the boric acid-sodium borate solution of pH 6.5. The anodic oxidation of Co(II) ion in the solution to form the film takes place at potentials more positive than 1.2V, which is in agreement with the potential at which an abrupt increase of the inner layer thickness as well as the outer layer nonstoichiometry commences.

Discussion

Potential distribution in the passivation film.—We consider the two-layered structure of an inner CoO and an outer Co$_3$O$_4$ shown in Fig. 10 and the potential distribution in the film shown in Fig. 11. If the reactions between the phases are in equilibrium at all the interphases, the irreversible potential differences in CoO (II phase) and in Co$_3$O$_4$ (III phase) are given by

$$\Delta\phi_{II} = \phi_{CoO}/CoO_4 - \phi_{CoO}/CoO$$

$$\Delta\phi_{III} = \phi_{CoO}/Co_3O_4 - \phi_{CoO}/CoO$$

where $\phi_{CoO}/CoO_4$, $\phi_{CoO}/CoO_4$, and $\phi_{CoO}/Co_3O_4$ are the potentials of electrode systems consisting of Co/CoO, CoO/CoO$_2$, and Co/CoO$_2$ respectively. If we assume the potential of $\phi_{CoO}/CoO$ to be 0.14 ± 0.04V vs. HESS, thermodynamically estimated by Göhr (6), and the potential of $\phi_{CoO}/CoO_4$ to be 0.75V, experimentally established by the authors (1,4,7), the irreversible potential difference in the CoO phase is

$$\Delta\phi_{II}/V = 0.61 ± 0.06$$

It is this potential difference which creates the driving force for growth of the CoO layer. It follows from Eq. [2] that the irreversible potential difference in the outer Co$_3$O$_4$ phase can not be defined as a con-
constant value, but varies depending on the nonstoichiometry $\Delta$.

Thermodynamically, we can describe the outer layer composition as a function of electrode potential. The layer structure may also be thermodynamically described, but a number of kinetic parameters are required for the determination of the composition profile in depth.

A kinetic model for two-layer formation.—There have been only a few theoretical studies on the two-layer growth on metals. Yurek, Hirth, and Rapp (8) and Shaw and Rolls (9) have derived kinetic equations of two-layer growth for relatively thick oxide scales on metals. We propose here a kinetic mechanism of two-layer growth in anodic oxide films, which explains the observed transient behavior of anodic oxide film on cobalt. For simplification we assume that the diffusing ion through the film toward the solution is Co(II) ion and that the inner and the outer layers are CoO and Co$_3$O$_4$, respectively. Further, we assume the reaction scheme shown in Table I, where subscripts I, IIa, IIb, IIIa, IIIb, and IV indicate respectively the locations in the electrode system illustrated in Fig. 10.

The reaction rate is expressed in terms of mol flux of cobalt(II) ion $k(x=a,b,...)/mol-Co(II)$ ion sec$^{-1}$ cm$^{-2}$.

From the mass balance of transferring ions, the following relationship can be obtained among the reaction rates

$$k_a = k_b$$

$$k_b = k_2 + k_3$$

and

$$k_c = k_2 + k_3$$

We now introduce the following quantities for the reaction rates

$$i_{II}/A-cm^{-2} = 2Fk_b$$

$$i_{II}/A-cm^{-2} = 2Fk_0$$

$$i_{diss}/A-cm^{-2} = 2Fk_t$$

$$i_{depos}/A-cm^{-2} = 2Fk_h$$

where $i_{II}$ and $i_{III}$ correspond respectively to the ionic C.D. of diffusing Co(II) ion through the inner CoO and the outer Co$_3$O$_4$ layers, and $i_{diss}$ and $i_{depos}$ to the dissolution C.D. and the deposition C.D. of these ionic C.D.'s can be used to describe the three experimentally measurable quantities, $i_{II}$, $L_{coo}$, and $L_{co3o4}$ by taking the mass balance given by Eq. [4], [5], [6], and [7] into account. From reactions [a], [b], [g], and [h], the anodic C.D. that can be measured is given by

$$i_a = \frac{4}{3} i_{III} - \frac{1}{3} i_{diss} + \frac{1}{3} i_{depos}$$

Table I. Reaction scheme for growth of CoO/Co$_3$O$_4$ two-layered film on Co

<table>
<thead>
<tr>
<th>Reaction</th>
<th>Rate (mol-Co ion sec$^{-1}$ cm$^{-2}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>[a] $Co_{III}^{2+} + 2e^{-} \rightarrow Co_{II}^{2+}$</td>
<td>$k_a$</td>
</tr>
<tr>
<td>[b] $Co_{II}^{2+} \rightarrow Co_{III}^{2+}$</td>
<td>$k_0$</td>
</tr>
<tr>
<td>[c] $Co_{II}^{3+} \rightarrow Co_{II}^{2+} + e^{-}$</td>
<td>$k_c$</td>
</tr>
<tr>
<td>[d] $Co_{III}^{2+} + 3Co_{III}^{2+} + 2e \rightarrow 4Co_{II}^{2+}$</td>
<td>$k_d$</td>
</tr>
<tr>
<td>[e] $Co_{III}^{2+} \rightarrow Co_{III}^{2+} + e^{-}$</td>
<td>$k_e$</td>
</tr>
<tr>
<td>[f] $Co_{II}^{3+} \rightarrow Co_{II}^{2+} + e^{-}$</td>
<td>$k_f$</td>
</tr>
<tr>
<td>[g] $Co_{III}^{2+} + 4/3$H$<em>2$O $\rightarrow 1/3Co</em>{III}^{2+}Co_{III}^{2+} + 8/3$H$^+$</td>
<td>$k_g$</td>
</tr>
<tr>
<td>[h] $4Co_{III}^{2+} + 4/3$H$<em>2$O $\rightarrow 1/3Co</em>{III}^{2+}Co_{III}^{2+} + 8/3$H$^+$</td>
<td>$k_h$</td>
</tr>
</tbody>
</table>

Fig. 11. A model of potential distribution in the two-layered passive film on cobalt. Dotted lines indicate the possible potential distribution at equilibrium for Co/CoO and Co/CoO/Co$_3$O$_4$ electrodes.

Since the inner layer increases in thickness by reaction [d], the growth rate can be described from Eq. [6] and [7] in terms of $i_{III}$ and $i_{III}$

$$\frac{dL_{co3o4}}{dt} = 4(i_{III} - i_{III})L_{co3o4}$$

where $L_{co3o4}/nm cm^2 C^{-1}$ is the thickness of Co$_3$O$_4$ equivalent to 1 C cm$^{-2}$ of Co(II) ion. The outer layer grows by reactions [g] and [h], and its thickness decreases by reaction [d]

$$\frac{dL_{coo}}{dt} = 2F(k_g + k_h - 3k_d)L_{co3o4}$$

where $L_{co3o4}/nm cm^2 C^{-1}$ is the thickness of Co$_3$O$_4$ equivalent to 1 C cm$^{-2}$ of Co(II) ion. Values of $L_{coo}$ and $L_{co3o4}$ were experimentally obtained; $L_{coo}/nm cm^2 C^{-1} = 9.1 \times 10^2$ and $L_{co3o4}/nm cm^2 C^{-1} = 7.7 \times 10^3$ (2, 3). From Eq. [13] and [14] it is evident that the growth of one layer is strongly dependent on the ionic C.D. of the diffusing Co(II) ion in the other layer.

It is assumed that the ionic flux through the layer is a function of electric field strength $E = \Delta \Phi/L$ following the Cabrera–Mott theory (10, 11).

$$i_{II} = i_{II} \exp (B_{II} \Delta \Phi/L_{in})$$

$$i_{III} = i_{III} \exp (B_{III} \Delta \Phi/L_{out})$$

where $i_{II}0$ and $i_{III}0$ indicate the ionic self-diffusion C.D. at zero field, and $B_{II}$ and $B_{III}$ are constants given.
Correspondingly, the thicknesses are determined at the stationary state, where
\( \frac{dL_{\text{in}}}{dt} = 0 = \frac{dL_{\text{out}}}{dt} \), and \( \frac{dL_{\text{in}}}{dt} = 0 \), the ionic transfer across every interphase is equal to the ionic diffusion through every layer
\[
\frac{dL_{\text{in}}}{dt} = (i_{\text{III}} - i_{\text{dis}}) L_{\text{CoO}}
\]
where \( i_{\text{III}} \) represents the anodic C.D. at the stationary state. Correspondingly, the thicknesses are determined by Eq. [15] and [16]. As \( \Delta \Phi_{\text{III}} \) remains equal to the irreversible potential difference given by Eq. [1] or [3] irrespective of the anodic potential, the thickness of the inner layer, \( L_{\text{in}} \), is determined by the dissolution rate, \( i_{\text{dis}} \), of the outer layer. It follows therefore that the inner layer, even if it is soluble, will grow to a certain thickness in the presence of the protective outer layer. Since the outer layer sustains an increasing potential difference \( \Delta \Phi_{\text{III}} \) with anodic potential \( \Phi_{\text{III}} \), its thickness will increase with anodic potential at the stationary state
\[
\Delta \Phi_{\text{III}} = \Delta \Phi_{\text{II}} - \Delta \Phi_{\text{I}}
\]
In our experimental results the thickness of the outer layer satisfies the above prediction for both one-step and two-step oxidation films, as shown in Fig. 8, whereas the thickness of the inner layer behaves differently depending on the formation procedure. It is thus suggested that the outer layer can approach the stationary thickness more quickly than the inner layer.

**Anodic deposition of Co(II) in solution.**—The difference in the inner layer thickness between one-step oxidation film and two-step oxidation film is caused by the anodic deposition reaction [h] in Table I, which occurs only in the one-step oxidation. As shown in Fig. 8 and 9, the potential at which the anodic deposition from Co(II) ion commences is in agreement with the potential at which an abrupt change of the inner layer thickness is observed. A question is now raised why the inner CoO layer does become thick, when Co(II) ions in the solution are deposited to form the outer layer of higher valence oxide. As the outer layer thickness increases by anodic deposition from Co ion, the rate of Co(II) ion diffusion in the outer layer decreases because of a decreasing electric field intensity in the layer according to Eq. [16] and eventually becomes negligible compared with the ionic diffusion rate in the inner layer
\[
i_{\text{II}} >> i_{\text{III}} = 0
\]
From Eq. [18] therefore the inner layer growth rate is
\[
\frac{dL_{\text{in}}}{dt} = 4i_{\text{III}} L_{\text{CoO}}
\]
which means that almost all Co(II) ions flowing through the inner layer to the CoO/CoO₄ interphase go into reaction [d] to form a new CoO layer at the expense of the outer CoO₄ layer. In this case the inner CoO layer may thicken without any diffusion of Co(II) ion through outer CoO₄ layer which is a barrier against ion flow. By two-step oxidation, because of the absence of dissolved Co(II) ions in the solution, the film growth requires the diffusion of Co(II) through both the layers and hence the film growth rate is small compared with that in one-step oxidation where the anodic deposition is involved.

**Inner layer growth.**—As shown in Fig. 4, when an anodic potential step was applied to a passivated cobalt, the outer layer was observed to grow first in an initial period of time until its thickness reached a steady value, where \( \frac{dL_{\text{out}}}{dt} = 0 \). The initial growth of the outer layer was followed by a gradual increase of the inner layer thickness in the later period of time while the outer layer thickness remained constant. Therefore, for the later stage of film growth, we obtain by substituting \( \frac{dL_{\text{out}}}{dt} = 0 \) and \( i_{\text{steps}} = 0 \) into Eq. [13] and Eq. [14] the following equations
\[
\frac{dL_{\text{in}}}{dt} = \frac{(i_{\text{II}} - i_{\text{dis}})}{L_{\text{CoO}}}
\]
Equation [23] means that the dissolution rate of the outer layer plays a large role in determining whether the inner layer increases or decreases after the outer layer reaches the stationary thickness. From Eq. [18] and the result shown in Fig. 4 and 5, it follows that the potential step from 1.04 to 1.24 V causes the inner layer to grow because \( i_{\text{II}} > i_{\text{dis}} \) and that the reverse potential step brings about a decrease of the inner layer because \( i_{\text{II}} < i_{\text{dis}} \).

The growth rate of the inner layer can be described from the results of Fig. 4 and from Eq. [24] by assuming the Cabrera-Mott relation of Eq. [15]. Figure 12 shows the relationship between the logarithm of anodic C.D. \( i_{\text{II}} \) and the inverse of the inner layer thickness during the second step oxidation at 1.24 V after the outer layer reaches the stationary thickness. From the slope and intercept of the curve, \( B_{\text{II}} \Delta \Phi_{\text{II}}/cm = 4.5 \times 10^{-7} \) and \( i_{\text{II}}/A\cdotcm^{-2} = 1.2 \times 10^{-8} \). If we assume \( \Delta \Phi_{\text{II}} = 0.61 \) (V) from Eq. [3], \( B_{\text{II}}/cm\cdotV^{-1} = 7.1 \times 10^{-7} \). From the value of \( B_{\text{II}} \) Eq. [17] gives \( a_{\text{II}}/cm = 1.8 \times 10^{-8} \) at \( Z_{\text{II}} = 2 \), and \( a_{\text{II}}/cm = 3.8 \times 10^{-9} \) at \( Z = 1 \). By comparing the estimated \( a_{\text{II}} \) values with the distance between the neighboring Co ions in CoO, calculated to be 0.30 nm from the lattice constant of 0.426 nm, it is likely that the migrating species is monovalent cobalt vacancy. The ionic conductivity \( \kappa \) at low fields may be written as
\[
\kappa = \left| \frac{\partial \Phi_{\text{III}}}{\partial (\Delta \Phi_{\text{III}}/L_{\text{in}})} \right| = i_{\text{III}}/L_{\text{CoO}}
\]

![Fig. 12. Relation between inverse of the inner layer thickness and the anodic C.D. in logarithmic scale. Data correspond to the results of \( \Phi = 1.24 \) V oxidation in Fig. 4.](image)
and is estimated to be $\kappa/S \text{ cm}^{-1} = 8.5 \times 10^{15}$. Furthermore, the self-diffusion constant $D$ of moving species in the inner layer may be estimated using Einstein-Nerst relation (12), as follows

$$D = \frac{i_{\text{H}} B_{\text{H}} RT}{2zF^2} \cdot \frac{1}{C_{\text{Co}}}$$  \[26\]

where $C_{\text{Co}}/\text{mol cm}^{-3}$ is the number of mol of CoO in unit volume and is calculated to be $C_{\text{Co}}/\text{mol cm}^{-3} = 0.086$ from the lattice constant. We estimate that $D/\text{cm}^2 \text{ sec}^{-1} = 6.5 \times 10^{-21}$ for $Z = 2$ and $D/\text{cm}^2 \text{ sec}^{-1} = 2.6 \times 10^{-20}$ for $Z = 1$. No comparable self-diffusion constant can be found in the literature and the estimated values in this work are $10^{20}$ times as large as the value extrapolated from the diffusion data for CoO obtained at high temperature above 1200 K by Richardson et al. (13) and by Chen et al. (14). The discrepancy may be attributed to a large concentration of lattice defects in the anodic oxide film.

**Conclusion**

The following conclusions may be drawn;

1. Cobalt is covered with a two-layered oxide film consisting of an inner CoO and an outer Co$_3$O$_4$ or Co$_9$O$_{24}$ in the passive potential region more positive than $\phi/V$ vs. HESS = 0.75 in neutral boric acid-sodium borate solution at pH 8.4.

2. A growth model is presented for electrochemical two-layer formation, which explains the observed dependence of film thickness on the potential and on the time of oxidation.

3. The outer layer reaches within 1 hr the stationary thickness which increases nearly linearly with the electrode potential, and the inner CoO layer requires much longer oxidation time before the stationary thickness is reached. The inner layer thickness that grows in a few hours' oxidation depends not only on the potential but also on the oxidation procedure. It is revealed that the growth of the inner CoO layer is influenced by the ionic diffusion in the outer layer.

4. Anodic deposition of a Co$_3$-nO$_4$ layer (0 $\leq$ $\Delta$ $\leq$ 0.33) from Co(II) ion in solution of pH 8.4 can take place at Co(II) ion concentrations as small as 1.03 mg dm$^{-3}$ in the potential region more positive than 1.2V. Therefore, the anodic oxide film on cobalt becomes thick in the presence of Co(II) ion in solution.

5. The ion migration in the anodic oxide film is likely to be assisted by high electric field and represented by the rate equation $i = i_0 \exp (B\phi/L)$ for each individual layer.

6. The self-diffusion coefficient in the inner CoO layer is estimated to be $D/\text{cm}^2 \text{ sec}^{-1} = 6.5 \times 10^{-21}$ $\sim$ $2.6 \times 10^{-20}$.

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**REFERENCES**

4. N. Sato and T. Ohtsuka, This Journal, 125, 1735 (1978).