Analysis of Kinetics Parameters Controlling Atomistic Reaction Process of a Quasi-Reversible Electrode System

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1. Introduction

For understanding the mechanism of electrolysis it is important to estimate kinetics parameters controlling the atomistic reaction process of metal electrode that is polarized in an electrolyte solution, but it seems not to have been performed satisfactorily. The reason for this is attributed to the fact that because actual electrode reactions proceed quasi-reversibly via consecutive two processes which consist of surface reaction and volume diffusion of ions involved in the reaction, the expression for its current density/overpotential relationship have become complex and not been presented explicitly. This is also related to the subjects of studies concerning the process of deposition or dissolution of atoms in crystal growth or its dissolution.

It is well known that the etch pits having a crystallographic symmetry are formed at dislocation sites of the low indices surfaces of a crystal which was etched under a specified condition (e.g. Gilman et al., 1958; Young, Jr., 1961). The dislocation etch pit is thought to be formed via a nucleation and growth process of two-dimensional pits at the dislocation site or via a spiral dissolution of the surface step which is caused by screw dislocation (Burton et al., 1951; Cabrela and Levine, 1956). Therefore elucidation of its formation mechanism is important for understanding of the surface step motion which is thought to play major role in the dissolution process of a crystal, and dissolution kinetics of crystals in the etch pit formation has been investigated and discussed by some researchers (e.g. Ives and Hirth, 1960; Schaarwächter, 1965; Jasper and Schaarwächter, 1966; Van Der Hoek et al., 1983) so far.

However the research concerning parameters controlling surface step motion in the dissolution of crystals has not been satisfactorily performed. Especially it has not been examined quantitatively except for a few studies (e.g. Onuma, 1991). This is principally due
to the reason that because the dissolution of a crystal proceeds generally via a dissolution reaction of surface atom and diffusion process of the dissolved atom (ion) into interior of solution, it is difficult to experimentally inspect the dissolution kinetics of surface step which depends on both processes. Since the dissolution rate of a metal crystal which is anodically dissolved under polarization in an electrolyte solution can be investigated by measurement of current density, dissolution mechanism of metal crystals has been researched electrochemically (e.g. Despic and Bockris, 1960; Lee and Nobe, 1986). However because of the same reason as the above mention, discussions on the results have become complex and not always contributed to understanding of surface step motion.

Recently, however, it has been proposed by the author that an expression to analyze the relationship between anodic current density and overpotential of a quasi-reversible electrode system including both the consecutive reaction processes is derived explicitly on the basis of an appropriate assumption (Imashimizu, 2010, 2011). According to the analysis, if the anodic and cathodic diffusion-limited current densities are measured for a given quasi-reversible electrode system, we can experimentally determine the kinetics parameters controlling dissolution process of crystals of the metal electrode, by assuming expressions for the activation and concentration overpotentials which are driving forces of surface reaction process and volume diffusion process respectively.

Thus the dissolution rates at dislocation-free and edge dislocation sites of (111) surface when a copper crystal was anodically dissolved in an electrolyte solution are investigated and discussed based on the above thinking, in this chapter. The relationships between anodic current density and overpotential are analyzed and discussed electrochemically by using the method developed for anodic dissolution processes of quasi-reversible electrode as described above. Activation enthalpy, transfer coefficient and surface concentrations of the ions involved in the dissolution process are experimentally estimated, and kinetics parameters controlling anodic reaction of the copper crystal/electrolyte system are quantitatively examined. An expression for the vertical dissolution rate at dislocation site is proposed based on a nucleation model of two-dimensional pit, and the critical free energy change at nucleation is quantitatively examined.

2. Experimental procedures for study of dissolution kinetics of copper crystals

2.1. Preparation of specimens

Single crystals of copper with [111] direction about 10 mm in diameter were prepared from the starting material of re-electrolyzed copper of 99.996 % purity by using the pulling method. They were divided into the cylindrical crystals approximately 15 mm length by a strain-free cutting. A terminal for detection of electric current and potential was soldered to an end surface of the cylindrical crystals. Another end surface was chemically polished so that the deviation of surface orientation from [111] direction is within 8.7×10^{-3} rad, and was further electrolytically polished in a high concentrated phosphoric acid solution. The crystal
specimen was embedded in a Teflon holder with paraffin so that its polished surface is exposed. Then it was supplied for the electrolysis experiment after the boundary portion between the paraffin and the periphery of polished surface was covered with a vinyl seal having a hole 6 mm in diameter (Watanabé et al., 2003).

2.2. Apparatus for potentiostatic electrolysis

Schematic diagram of the electrolytic cell for this experiment is shown in Fig.1 (Watanabé et al., 2003). The crystal specimen was immersed in the electrolyte solution which consists of 5 kmol m\(^{-3}\) NaCl, 0.25 kmol m\(^{-3}\) NaBr and 10\(^{-4}\) kmol m\(^{-3}\) CuCl (Jasper and Schaarwächter, 1966) so that (111) surface of copper is located at approximately 5mm below the surface of electrolyte solution, and was held at a specified temperature. Then, the crystal specimen was set under a constant overpotential, and (111) surface of the crystal was anodically dissolved for a prescribed time, while anodic current density/time curve was recorded. The potentiostatic electrolysis experiments were performed at a range of lower overpotential and at a range of higher overpotential. After that the structure of dissolved surfaces were observed by use of the optical microscope system equipped with lens for interferometry.

![Figure 1. Schematic diagram of electrolytic cell. S: sample; E: electrolyte; WE: terminal for potential and current; RE: saturated calomel electrode; CE: counter electrode of platinum wire; B: salt bridge; I: thermobath; R: Regulator; and T: thermometer.](image)

2.3. Features of anodic current density/time curves and structure of dissolved surface

Figure 2 shows typical anodic current density/time curves which were recorded while the copper crystal was anodically dissolved for 360 s or 600 s at the respective overpotentials. Anodic current density under any condition decreases steeply immediately after start of electrolysis and reaches a nearly constant current density \(i_s\) when it was carried out at an
overpotential lower than about 125 mV as shown by the curve of 87 mV in Fig. 2. Figures 3 (a), (b) and (c) are the optical micrographs of the (111) surfaces which were dissolved for 600 s at overpotentials in a range of 60 mV to 125 mV being held at 298K. The surfaces are rather smooth though etch pits tend to be formed as overpotential increases.

**Figure 2.** Anodic current density/time curves under potentiostatic electrolysis at 298K.

**Figure 3.** Optical micrographs of the surfaces which were anodically dissolved at lower overpotentials of (a): 58 mV, (b): 88 mV, and (c): 108 mV.

On the other hand, the current density reaches a minimum current density $i_{sm}$ that is pointed by arrow after the initial steep decrease when an overpotential higher than about 125 mV was applied. Then it tends to increase gradually along with fluctuating and take a higher steady value as shown by the curve of 176 mV. Figures 4 (a), (b), and (c) are optical
micrographs of the surfaces which were dissolved for 300 s at 156, 166, and 176 mV respectively being held at 298K. One can see that etch pits are significantly formed.

![Micrographs](image)

**Figure 4.** Optical micrographs of the surface that were anodically dissolved under potentiostatic electrolysis at higher overpotentials of (a): 156 mV; (b): 166 mV; (c): 176 mV.

### 2.4. Measurement of anodic current density

#### 2.4.1. Steady anodic current densities at lower overpotentials

The initial steep decrease of current density is principally due to the fact that a diffusion layer of the dissolved atoms (ions) forms in the neighborhood of crystal surface in process of time so as to decrease the undersaturation which is driving force for the dissolution. Therefore an approximately constant current density after its initial steep decrease is thought to be a steady current density $i_s$ which flows accompanying with the consecutive two dissolution processes consisting of surface reaction and volume diffusion of dissolved atoms. This shows that the copper crystal/electrolyte system is a quasi-reversible electrode. Also the steady anodic current densities at lower overpotentials are thought to have only a little influence of formation of dislocation etch pits. Thus we assume that $i_s$ is related to the vertical dissolution rate $v_s$ at dislocation-free site of surface, which is given by the following expression (Schaarwächter and Lücke, 1967):

$$v_s = \frac{\Omega}{ne} i_s$$  \hspace{1cm} (1)

where $e \ [C]$ is elementary charge (electronic charge), $n \ [1]$ the charge number transferred at reaction and $\Omega \ [m^3]$ the atomic volume.

In this experiment, the potentiostatic electrolysis at overpotentials in a range from about 60 mV to 125 mV were carried out for 600 s at each temperature of 268, 283, 298 and 308 K and the relationships between the steady anodic current densities $i_s$ and applied overpotentials $\eta$ were investigated.
2.4.2. Minimum anodic current densities at higher overpotentials

On the other hand, the current density reaches a minimum after an initial steep decrease as shown in the curve of 176 mV in Fig.2 when an overpotential higher than about 125 mV was applied. Then it tends to increase gradually together with fluctuating and take a higher steady value as described in Section 2.3. This is thought to be due to the fact that etch pits remarkably formed at dislocation sites and grew along with time under higher overpotential as shown in Fig.4 (Schaarwächter and Lücke, 1967; Imashimizu and Watanabé, 1983). That is, it is because nucleation and growth of etch pits at dislocation site resulted in an increase of the anodic current density which represents an average dissolution rate of whole surface exposed to electrolyte solution as the areas occupied by etch pits increase. Based on the above knowledge, we assume that the initial minimum current density \(i_{sm}\) under potentiostatic electrolysis at higher overpotentials is approximately equal to a current density that is equivalent to the dissolution rate of dislocation-free site of surface because the contribution to anodic current density of dislocation etch pit formation is thought to be a little in the initial stage of electrolysis. That is, an average value of \(i_{sm}\) was assumed to give the vertical dissolution rate at dislocation-free site of surface approximately as represented by the relation:

\[
v_s = v_{sm} = \frac{\Omega}{n_e} i_{sm}
\]

Thus the electrolysis experiment was carried out for a prescribed time from 60 s to 360 s at each overpotential of 156, 166, 176 and 186 mV keeping the temperature at 298 K and at each temperature of 268, 283, 298 and 308 K under an overpotential of 176 mV. The initial minimum current densities \(i_{sm}\) were obtained from the anodic current density/time curves measured under every condition.

2.5. Measurement of polarization curve and estimation of the diffusion-limited current densities

It needs to estimate activation overpotential \(\eta_a\) and concentration overpotential \(\eta_c\) for analyzing the relationship between anodic current density and applied overpotential as described in Section 1. Thus the polarization curves in a range of overpotential of about -400 mV to 400 mV were measured three times at each temperature of 298K and 308K by the potential step method. Then the anodic and cathodic diffusion-limited current densities were estimated.

2.6. Direct measurement of dissolution rates of surface

2.6.1. Vertical dissolution rate of surface

After the (111) surface of a copper crystal specimen was anodically dissolved at every condition of specified overpotentials and temperatures as described in Section 2.4.2, it was
observed by use of the optical microscope equipped with objective lenses for two-beam interferometry and multiple interferometry.

Figure 5 (a) shows the micrograph of a part of boundary region between the crystal surface exposed to the electrolyte solution and the peripheral portion covered with vinyl seal, which was photographed with two-beam interferometry mode. The vertical dissolution amounts \( s \) of surface shown by the illustration was measured from a deviation of the interference stripes caused by the step which was formed at that boundary region after dissolved. The vertical dissolution amounts \( s \) of surface under each condition was plotted against dissolution time \( t \). The increasing rate \( \dot{s} \) of \( s \) with \( t \) was obtained from the gradient of each linear relationship, and the vertical dissolution rate of surface under every condition was estimated by the \( \dot{s} \).

### 2.6.2. Dissolution rates at dislocation site of surface

Figures 5 (b) and (c) show a pair of micrographs of identical dislocation etch pits formed on dissolved surface which were photographed with optical mode and multiple interferometry mode. In this work, the depth \( d \) of the dark (deep) pits that were formed at positive edge dislocation sites (see Appendix A1) were measured by drawing the vertical cross sections of the pits that is shown by the illustration with use of the micrograph pairs such as Figs.5 (b) and (c). Also the width \( w \) (average distance from center to the three sides of pit) of those dark pits that is shown by the illustration were measured on the micrograph such as Fig.5 (b). Measurements of the depth and width of pit were performed about more than 20 dark pits formed on the surface dissolved under every condition, and the respective average values \( d \) and \( w \) were obtained. The depth \( d \) and the width \( w \) of dark pits were plotted against dissolution time \( t \).

![Figure 5](image)

**Figure 5.** (a): Two-beam interferometry micrograph at the boundary between dissolved and undissolved surfaces; (b): Optical micrograph; (c): Multiple interferometry micrograph of the same view as b.

The increasing rate \( \dot{d} \) of \( d \) with \( t \) was obtained from the gradient of each linear relationship, and the vertical dissolution rate \( v_{ed} \) at edge dislocation site was estimated by

\[
v_{ed} = \dot{d} + v_s, \tag{3}
\]
where $v_s$ means the vertical dissolution rate at dislocation-free site of the surface. Also the increasing rate $w' \, w$ with $t$ was obtained from the gradient of each linear relationship, and the lateral dissolution rate $v_w$ at edge dislocation site was estimated from the relation:

$$v_w = \frac{w}{d} v_{ed} = \frac{w}{d} \left( d + v_s \right) .$$

### 2.7. Analysis of relationship between current density and overpotential

Under potentiostatic electrolysis of the copper/electrolyte system in the present experiment, the copper crystal is thought to be dissolved accompanying an anodic current according to a simple electrode reaction expressed by the following equation (Lal and Thirsk, 1953; Jasper and Scaarwächter, 1966):

$$\text{Cu} + 2\text{Cl}^- \rightarrow \text{CuCl}_2^- + e,$$

where the contribution to current density of reaction of Br ion involved in dissolution process as inhibitor is assumed to be disregarded. The anodic current density $i_s$ flowing steadily at an applied overpotential $\eta$ is generally expressed by a relation:

$$i_s = i_0 \left[ \left( \frac{C_{\text{Cl}^-}}{C^0_{\text{Cl}^-}} \right)^2 \exp \left( \frac{\alpha n e \eta}{kT} \right) - \frac{C_{\text{CuCl}_2^-}}{C^0_{\text{CuCl}_2^-}} \exp \left( -\frac{(1-\alpha) n e \eta}{kT} \right) \right],$$

where exchange current density $i_0$ is represented by

$$i_0 = n e \beta k_s C^0_{\text{Cl}^-} 2^{(1-\alpha)} C^0_{\text{CuCl}_2^-} \nu \exp \left( -\frac{\Delta H_0}{kT} \right) .$$

(Tamamushi, 1967; Maeda 1961). $k_s$ is surface density of kink that is active site at dissolution of surface atom, $\alpha$ the transfer coefficient, $\Delta H_0$ the activation energy (enthalpy) at dissolution of an atom, $\nu$ the atomic frequency, and $\beta$ a supplementary factor of rate constant of electrode reaction. Also, $C_{\text{Cl}^-}$ and $C_{\text{CuCl}_2^-}$ are the surface concentrations of Cl$^-$ and CuCl$_2^-$ ions involved in a steady anodic dissolution, and $C^0_{\text{Cl}^-}$ and $C^0_{\text{CuCl}_2^-}$ the ones in equilibrium state. They are represented as a relative surface density as follows.

If the electrolyte solution contacting with crystal surface contained $X$ ions of $m$ kmol m$^{-3}$, the surface concentration $C_X [1]$ can be expressed by the following relation:

$$C_X = \frac{m \times 10^3 N_A \left[ \text{m}^{-3} \right] \times \xi [\text{m}]}{(bb^*)^{-1} \left[ \text{m}^{-2} \right]},$$

where $N_A$ is the Avogadro constant, $bb^*$ the area occupied by an atom and $\xi$ the thickness of electrolyte solution layer contacting with the crystal surface (Imashimizu, 2011).
The anodic dissolution of copper crystal in this experiment is thought to proceed quasi-reversibly with a surface reaction and volume diffusion of dissolution atom as described in Section 2.4.1. So we assume that the activation overpotential \( \eta_a \) and the concentration overpotential \( \eta_c \) are written by

\[
\eta_a = \eta - \eta_c \quad \text{and} \quad \eta_c = \frac{kT}{ne} \ln \left( \frac{C_{\text{CuCl}_2^-}/C_0^{\text{CuCl}_2^-}}{\left(\frac{C_{\text{Cl}^-}}{C_0^{\text{Cl}^-}}\right)^2} \right),
\]

(9)

where the following relations:

\[
\frac{C_{\text{Cl}^-}}{C_0^{\text{Cl}^-}} = 1 - \frac{i_s}{i_{\text{Cl}^-}}, \quad \frac{C_{\text{CuCl}_2^-}}{C_0^{\text{CuCl}_2^-}} = 1 - \frac{i_s}{i_{\text{CuCl}_2^-}}
\]

(10)

are given, if \( i_{\text{Cl}^-} \) and \( i_{\text{CuCl}_2^-} \) are the anodic and cathodic diffusion-limited current densities of the electrode reaction respectively (Tamamushi, 1967). Thus activation overpotential \( \eta_a \) and concentration overpotential \( \eta_c \) are assumed to be given by Eqs.(9) and (10), when the anodic dissolution of copper crystal proceeds steadily at an applied overpotential \( \eta \) by a quasi-reversible electrode reaction of Eq.(5). Also surface undersaturation \( \sigma \) is defined by

\[
\sigma = 1 - \exp \left( -\frac{n\eta_a}{kT} \right) = 1 - \exp \left( -\frac{n\eta}{kT} \right) \frac{C_{\text{CuCl}_2^-}/C_0^{\text{CuCl}_2^-}}{\left(\frac{C_{\text{Cl}^-}}{C_0^{\text{Cl}^-}}\right)^2}.
\]

(11)

Then, the Eq.(6) is reduced to

\[
i_s = ne\beta_k_s C_0^{\text{Cl}^-} \sigma^{2(1-\alpha)} C_0^{\text{CuCl}_2^-} \alpha \left(\frac{C_{\text{Cl}^-}}{C_0^{\text{Cl}^-}}\right)^2 \sigma \exp \left( -\frac{\Delta H_0 - a\eta \eta}{kT} \right)
\]

(12)

by using Eqs. (7), (9) and (11). Also Eq.(12) leads to the following relation:

\[
i_s \left( \frac{C_{\text{Cl}^-}}{C_0^{\text{Cl}^-}} \right)^{-2} \sigma^{-1} = i_0(T) \exp \left( \frac{a\eta \eta}{kT} \right).
\]

(13)

Thus if the anodic and cathodic diffusion-limited current densities \( i_{\text{Cl}^-} \) and \( i_{\text{CuCl}_2^-} \) are obtained, the experimental relationship of \( i_s/\eta \) would be represented with use of Eqs.(10) and (11) by Eq.(13). Then \( \alpha \) and \( i_0(T) \) would be estimated from the gradient and the constant term of the linear relationship of \( \ln[i_{\text{Cl}^-}/C_{\text{Cl}^-}^2 \sigma^{-1}] \) vs. \( n\eta/kT \). Also \( \Delta H_0 \) would be estimated from the gradient of the linear relationship of \( \ln[i_0(T)] \) vs. \( 1/T \).

On the other hand, concerning the complex term consisting of surface concentrations of \( \text{Cl}^- \) and \( \text{CuCl}_2^- \) ions,
\[
C_{\text{Cl}^-} \cdot 2(1-\alpha)C_{\text{CuCl}_2^-}^0 \cdot \alpha = C_{\text{Cl}^-}^0 \cdot 2(1-\alpha)C_{\text{CuCl}_2^-}^0 \cdot \alpha \left( \frac{C_{\text{Cl}^-}}{C_{\text{Cl}^-}^0} \right)^2 \exp \left( \frac{-\alpha \eta}{kT} \right) 
\]

is lead from Eq.(9). Therefore applying Eq.(14) to Eq.(12) lead to

\[
i_s = ne\beta k_s C_{\text{Cl}^-} \cdot 2(1-\alpha)C_{\text{CuCl}_2^-}^0 \cdot \alpha \sigma \nu \exp \left( -\frac{\Delta H}{kT} \right),
\]

where \(\Delta H\) is given by the relation:

\[
\Delta H = \Delta H_0 - \alpha \eta \sigma.
\]

We can see that Eqs.(15) and (16) are formulae for the steady current density expressed with use of the parameters \(\beta, k_s, C_{\text{Cl}^-}, C_{\text{CuCl}_2^-}, \sigma, \alpha\) and \(\Delta H\) involved in the surface reaction process when the anodic dissolution progresses steadily by a quasi-reversible electrode reaction.

Thus undersaturation \(\sigma\), transfer coefficient \(\alpha\) and activation enthalpy \(\Delta H_0\) for the anodic dissolution reaction of copper crystal/electrolyte system will be estimated from experimental results, and a supplementary factor \(\beta\) and kink density \(k_s\) will be examined by a model of crystal dissolution in this study.

3. Experimental results

3.1. Polarization curves and undersaturation in anodic dissolution

The polarization characteristic of the copper crystal/electrolyte system at 298K is shown in Fig. 6. The anodic and cathodic diffusion-limited current densities \(i_{\text{Cl}^-}\) and \(i_{\text{CuCl}_2^-}\) shown in the diagram were obtained by averaging the values measured three times. Table 1 shows those diffusion-limited current densities obtained from the polarization characteristics measured at 298 and 308 K by a similar method.

Figure 6. Polarization curve of the copper crystal/electrolyte system. \(i_{\text{Cl}^-}\) and \(i_{\text{CuCl}_2^-}\) mean the anodic and cathodic diffusion-limited current densities.
Table 1. Measurements of the anodic and cathodic diffusion-limited current densities $i_{Cl^-}$ and $i_{CuCl_2^-}$.

<table>
<thead>
<tr>
<th>T/K</th>
<th>$i_{Cl^-}$/A m$^2$</th>
<th>$i_{CuCl_2^-}$/ A m$^2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>298</td>
<td>827</td>
<td>-0.0732</td>
</tr>
<tr>
<td>308</td>
<td>1072</td>
<td>-0.156</td>
</tr>
</tbody>
</table>

The undersaturation $\sigma$ were estimated from experimental polarization characteristics such as Fig.6 with use of Eqs.(10) and (11). The diagram that plotted $\sigma$ against $ne\eta/kT$ in a range of $(ne\eta/kT)$ about 0 to 6 is shown in Fig.7. The black dots in the diagram show the values of $\sigma$ which are calculated from the $(i_s/i_{Cl^-})/(ne\eta/kT)$ relationship that was derived by substituting the experimental values $i_0$, $\alpha$, $i_{Cl^-}$ and $i_{CuCl_2^-}$ into Eq.(6).

The experimental relationships of $\sigma/(ne\eta/kT)$ at 298K and 308K approximately consist with each other, and also with the calculated relationship. However, the experimental curves of $\sigma/(ne\eta/kT)$ deviate from the calculated curve in a range of $(ne\eta/kT)$ larger than about 5. This is because the experimental current density includes an increase of current density attributed to significant formation of etch pits at higher overpotentials than about 125 mV. We assumed that the $\sigma/(ne\eta/kT)$ relationship does not almost depend on temperature from the result of Fig.7.

![Figure 7](image-url)

**Figure 7.** Plots of undersaturation $\sigma$ against normalized overpotential $ne\eta/kT$. $\sigma(298)$ and $\sigma(308)$ designate experimental values at 298K and 308K respectively. $\sigma(\text{cal})$ is the calculated one.

### 3.2. Estimations of parameters controlling exchange current density

Figure 8 is the diagram that plotted the steady current densities against overpotentials lower than 127 mV which were measured at 268K, 283K, 298K and 308K. Figure 9(a) is the diagram that plotted $\ln(i_s\sigma^{-1})$ obtained from Fig. 8 against $ne\eta/kT$ at every temperature taking account of $(C\sigma^-/C\sigma^-)^2 \approx 1$. The linear relationships at every temperature in the diagram are drawn so that they have a same gradient given by averaging. The transfer coefficient $\alpha$ was estimated from the gradient of their linear relationships. Then also the exchange current densities $i_s(T)$ at each temperature were estimated from the constant terms of them. Figure 9(b) is the diagram that plotted $\ln(i_s(T))$ against $1/T$. The activation enthalpy for the anodic dissolution reaction of copper crystals was estimated from the gradient of the linear relationship shown in Fig. 9(b).
Figure 8. Plots of steady current densities against overpotentials lower than about 125 mV.

Figure 9. (a): Plots of \( \ln(i_s^2) \) against \( n\eta/kT \); (b): Plot of \( \ln(i_0(T)) \) against \( 1/T \).

Then, because surface concentrations \( C_{0Cl^-} \) and \( C_{0CuCl_2^-} \) of Cl\(^-\) and CuCl\(_2^-\) ions are calculated by Eq.(8) when (111) surface of a copper crystal is in equilibrium with the electrolyte solution consisting of 5 kmol m\(^{-3}\) NaCl and 10\(^{-4}\) kmol m\(^{-3}\) CuCl, the complex term \( C_{0Cl^-}^{(1-\alpha)}C_{0CuCl_2^-}^{\alpha} \) in Eq.(7) giving exchange current density can be evaluated by using the transfer coefficient \( \alpha \) estimated above.

Thus the estimations of parameters controlling exchange current density are summarized in Table 2. The value of \( \beta k_s \) was evaluated by substituting \( i_0, \Delta H_0 \) and \( C_{0Cl^-}^{(2\alpha)}C_{0CuCl_2^-}^{-\alpha} \) into Eq.(7), where the atomic frequency \( \nu = 6.21 \times 10^{12} \text{ s}^{-1} \), elementary electric charge, \( e = 1.602 \times 10^{-19} \text{ C} \) and \( n = 1 \) were assumed.

<table>
<thead>
<tr>
<th>( \alpha )</th>
<th>( \Delta H_0 ) / eV</th>
<th>( i_0 / 10^{-2} \text{ A m}^{-2} )</th>
<th>( C_{0Cl^-}^{(2\alpha)}C_{0CuCl_2^-}^{-\alpha} )</th>
<th>( \beta k_s / 10^{16} \text{ m}^{-2} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.84</td>
<td>0.33</td>
<td>8.2*</td>
<td>2.36\times10^{-6}</td>
<td>1.32</td>
</tr>
</tbody>
</table>

* the value at 298K

Table 2. Estimation of transfer coefficient \( \alpha \), activation enthalpy \( \Delta H_0 \), exchange current density \( i_0 \) at 298K, and a factor \( \beta k_s \) affecting reaction rate constant.
3.3. Anodic dissolution rates at higher overpotentials

3.3.1. Estimation of vertical dissolution rate of surface from anodic current density

Figures 10 (a) and (b) are examples of the anodic current density/time curves which were recorded when the copper crystal was dissolved for 240 s at higher overpotential. The vertical dissolution rate $v_{sm}$ at dislocation-free site of surface under every condition was determined from an average of the initial minimum current densities $i_{sm}$ pointed by arrow of $i/t$ curves (measured for five different dissolution time in a range of 60 s to 360 s under each condition) shown in Fig.10 as described in Sections 2.4.2.

![Figure 10](image1.png)

**Figure 10.** Examples of anodic current density/time curves. (a): The effect of overpotential; (b): The effect of temperature. Initial minimum $i_{sm}$ was obtained in every curve.

3.3.2. Estimation of vertical dissolution rate of surface by direct measurement

Figures 11 (a) and (b) are the diagrams that plotted vertical dissolution amounts $s$ of surface against dissolution time $t$ as described in Sections 2.6.1. The vertical dissolution rates $\dot{s}$ of surface were estimated from the gradient of the linear relationship of $s/t$ shown in Fig.11.

![Figure 11](image2.png)

**Figure 11.** Vertical dissolution amounts of surface vs. dissolution time. (a): Effect of overpotential at 298K; (b): Effect of temperature at 176mV
3.3.3. Estimation of dissolution rates at edge dislocation site by direct measurement

Figures 12 (a) and (b) are the diagram that plotted the depth \( d \) of the dark etch pits which were formed at positive edge dislocation sites on the surface dissolved under each condition against dissolution time \( t \), as described in Sections 2.6.2. Also Figs.13 (a) and (b) are the diagrams that plotted similarly the width \( w \) of the same dark etch pits as the above mentioned against dissolution time \( t \). The increasing rate \( \dot{d} \) of depth \( d \) of etch pit with \( t \) and the increasing rate \( \dot{w} \) of width \( w \) with \( t \) were obtained from the gradient of each linear relationship shown in those results.

![Figure 12](image12.png)

**Figure 12.** Depth of dark etch pit vs. dissolution time. (a): Effect of overpotential at 298K; (b): Effect of temperature at 176mV

![Figure 13](image13.png)

**Figure 13.** Width of dark etch pit vs. dissolution time. (a): Effect of overpotential at 298K; (b): Effect of temperature at 176mV.

3.4. Effects on the dissolution rates of overpotential and temperature

Figures 14 (a) and (b) are the diagrams that plotted the logarithm values of the dissolution rates. It can be seen that the value of \( \log \dot{v}_{sm} \) approximately consists with that of \( \log \dot{s} \). This shows that both \( \dot{v}_{sm} \) and \( \dot{s} \) represent the dissolution rate \( \dot{v}_s \) of dislocation-free site of surface approximately. However, the value of \( \dot{v}_{sm} \) seems to be more exact than that of \( \dot{s} \), because while the former is a quantity related to total dissolution amounts of whole surface, the latter is that related to the dissolved amounts of a part of surface near to boundary between the portion exposed to electrolyte solution and the portion covered by vinyl seal. Thus the dissolution rate \( \dot{v}_s \) of dislocation-free surface was assumed to be given not by \( \dot{s} \) but \( \dot{v}_{sm} \). Then \( \dot{v}_ed \) and \( \dot{v}_w \) in Fig.14 show the values estimated from Eq.(3) and Eq.(4) in which \( \dot{v}_s \) was substituted by \( \dot{v}_{sm} \).
Figure 14. (a): Plots on a logarithmic scale of dissolution rates $v_{sm}$, $v_{ed}$, and $v_w$ against overpotential $\eta$; (b): Similar plots of $v_{sm}$, $v_{ed}$, and $v_w$ against temperature $T$.

It can be seen that both log $v_{sm}$ and log $v_w$ tend to increase rather homogeneously with an increase of $\eta$, from Fig.14 (a). However, the tendency of log $v_{ed}$ are somewhat different and in accelerative. Also, it can be seen from Fig.14 (b) that though log $v_{sm}$ and log $v_w$ tend to similarly increase with an increase of $T$, the tendency of log $v_{ed}$ are somewhat little, compared to the former two. This is seen from the fact that the increasing rate ($\Delta \log v_{ed}/\Delta T = 5.4 \times 10^{-3}$) of the latter is less than that ($\Delta \log v_{sm}/\Delta T = 1.1 \times 10^{-2}$, $\Delta \log v_w/\Delta T = 1.4 \times 10^{-2}$) of the former two.

4. Discussion

4.1. Atomistic dissolution model of crystal surface

4.1.1. Vertical dissolution rate at dislocation-free site of surface

Concerning the dissolution of a crystal, the atomistic model illustrated in schematic diagram of Fig.15 has been proposed (Burton et al., 1951; Schaarwächter, 1965). The dissolution of crystals proceeds via a lateral retreat motion of surface step of an atomic height that is induced by dissolving of surface atom from the kink sites into the solution. The vertical dissolution rate $v_s$ of surface is given by lateral retreat rate $v_h$ and surface density $\tan \theta$ of surface step, which is expressed by the following equation:

$$v_s = v_h \tan \theta = v_h \frac{a}{\lambda},$$

(17)

where $\theta$ is an average inclination of crystal surface to a low index face, $a$ an atomic height of surface step, and $\lambda$ the mean distance between adjacent surface steps. The lateral retreat rate $v_h$ of surface step is expressed by

$$v_h = b \ast k \ast \sigma \ast \nu \ast \exp \left( -\frac{\Delta H_s}{kT} \right),$$

(18)
where $\Delta H_s$ is the activation enthalpy for dissolution of an atom at kink site of surface step, $\nu$ the atomic frequency, $k^*$ the retreat rate constant of surface step, and $b^*$ the unit retreat distance. $\sigma_s$ is surface undersaturation, which is written as

$$\sigma_s = 1 - \exp\left(-\frac{\Delta \mu}{kT}\right),$$

where $\Delta \mu$ is the chemical potential difference of dissolution atom between two phases of a crystal/solution system (Schaarwächter, 1965).

**Figure 15.** Atomistic model for dissolution process of a crystal surface. Atom dissolves from kink site of surface step into solution. K: Kink; S: Surface step; T: Terrace; A: Ad-atom

### 4.1.2. Lateral dissolution rate at edge dislocation site

The dislocation etch pit is thought to be formed via a successive nucleation and growth processes of two-dimensional pits at the dislocation site (Schaarwächter, 1965) or via a spiral dissolution of the surface step which is caused by screw dislocation (Cabrera and Levine, 1956). We discuss the dissolution rate at edge dislocation site of (111) surface of copper crystals, based on a nucleation and growth model of two-dimensional pits (Schaarwächter, 1965) that is illustrated in Fig 16, in the following.

Since the lateral dissolution rate $v_w$ is thought to represent horizontal growth rate of two-dimensional pit nucleated at edge dislocation site of surface, it may be corresponding to the lateral retreat rate $v_h$ of surface step along (111) face. Thus we assume that $v_h$ is given by $v_w$ as shown in the following relation:

$$v_h = v_w.$$  \hfill (20)

### 4.1.3. Vertical dissolution rate at edge dislocation site

On the other hand the vertical dissolution rate at positive edge dislocation site would be examined by the nucleation rate of two-dimensional pit at dislocation site as follows.
According to the classical nucleation theory, if $\Delta G_{ed}^*$ is the critical free energy change at nucleation of a two-dimensional pit at edge dislocation site, a steady state nucleation rate $I$ of two-dimensional pit would be expressed by

$$I = Zr \exp \left( \frac{-\Delta G_{ed}^*}{kT} \right), \quad (21)$$

where $r$ is a separation rate of an atom from an active site of the two-dimensional pit into the solution and $Z$ the Zeldovich factor (Toschev, 1973). Since the separation rate $r$ is assumed to be a similar quantity to the dissolution rate of an atom from kink site of surface, it depends on the surface concentrations of Cl$^-$ and CuCl$_2^-$ ions as known from the Eq. (15) in Section 2.7, and is expressed by

$$r \propto C_{Cl^-}^{2(1-\alpha)} C_{CuCl^-_2}^{\alpha} \nu \exp \left( \frac{-\Delta H_0}{kT} \right). \quad (22)$$

Accordingly, the vertical dissolution rate $v_{ed}$ at edge dislocation site of surface is expressed by

$$v_{ed} = aK_s C_{Cl^-}^{(1-\alpha)} C_{CuCl^-_2}^{\alpha} \nu \exp \left( \frac{-\Delta G_{ed}^* + \Delta H}{kT} \right), \quad (23)$$

where $a$ is the depth of "two-dimensional pit and $K_s$ is an undetermined constant including Zeldovich factor and others (see Appendix A2.).

According to the nucleation theory of dissolution of crystals, $\Delta G_{ed}^*$ is small compared to $\Delta G_s^*$ which is the critical free energy change at nucleation of a two-dimensional pit at dislocation-free site of surface, because of strain energy of dislocation core. It is expressed by

$$\Delta G_{ed}^* = p\Delta G_s^* = p \frac{\pi a \Omega \gamma^2}{\Delta \mu} \quad (24)$$

and
where \( \gamma \) is the interfacial free energy of the crystal and solution at step of the two-dimensional pit, \( G \) the shear modulus and \( q \) and \( \alpha \) the constants (Schaarwächter, 1965).

4.2. Relations between vertical dissolution rate of surface and anodic current density

4.2.1. Expression for dissolution rate of dislocation-free site of surface

When the copper crystal is anodically dissolved by the simple electrode reaction of Eq.(5) the vertical dissolution rate \( v_s \) of dislocation-free surface at lower overpotentials and the \( v_{sm} \) at higher overpotentials would be estimated by Eq.(1) and Eq.(2) respectively as described in Section 2.4. Thus it is experimentally estimated with use of Eqs. (1), (2), and (15) by the following expression:

\[
v_{sm} = v_s = \Omega \beta k_s C_{Cl^-}^{2(1-\alpha)} C_{CuCl^2-}^\alpha \sigma \nu \exp \left(-\frac{\Delta H}{kT} \right) . \tag{26}
\]

According to the dissolution model of crystals, the dissolution rate at dislocation-free site of surface is expressed from Eqs. (17) and (18) by

\[
v_s = \frac{a}{\lambda} b^* k^* \sigma_s \nu \exp \left(-\frac{\Delta H_s}{kT} \right). \tag{27}
\]

Therefore, the following relations are obtained from Eqs.(16), (26) and (27) concerning the rate constant of the lateral retreat rate of surface step and activation enthalpy for the dissolution.

\[
k^* = \beta \frac{b}{\chi_0} C_{Cl^-}^{2(1-\alpha)} C_{CuCl^2-}^\alpha = \beta \frac{b}{\chi_0} C_0^{2(1-\alpha)} C_0^\alpha \left(C_{Cl^-} / C_{CuCl^2-} \right)^2 \exp \left(\frac{ane \eta_c}{kT} \right) \tag{28}
\]

and

\[
\Delta H_s = \Delta H = \Delta H_0 - an \eta_a \tag{29}
\]

Also from Eqs. (11) and (19)

\[
\Delta \mu = n \eta_a \tag{30}
\]

is obtained. It can be seen that the rate constant \( k^* \) of lateral retreat motion of surface step is electrochemically expressed by Eq.(28) and that it increases with an increase of concentration overpotential \( \eta_c \).
4.2.2. Estimation of kinetics parameters controlling the dissolution rate

As mentioned above the dissolution rate \( v_{sm} \) at dislocation-free site of surface under higher overpotentials is expressed by an approximate equation:

\[
v_{sm} = \Omega \beta k_s C_0^{Cl^-} 2^{(1-\alpha)} C_0^{CuCl_2^-} \alpha \sigma v \exp \left( \frac{-\Delta H_0 - \alpha \eta \sigma}{kT} \right),
\]

from Eqs. (14) and (26), where we assumed \( i_s \ll i_{Cl^-} \), that is,

\[
\frac{C_{Cl^-}}{C_0^{Cl^-}} = 1 - \frac{i_s}{i_{Cl^-}} = 1.
\]

Thus concerning the dissolution rate at dislocation-free site of surface which have a constant kink density \( k_s \), a following approximate expression is lead from Eq.(31) (Imashimizu, 2011).

\[
\ln v_{sm} = \ln(\nu \Omega \beta k_s C_0^{Cl^-} 2^{(1-\alpha)} C_0^{CuCl_2^-} \alpha \sigma v) + \ln(\frac{-\Delta H_0 - \alpha \eta \sigma}{kT})
\]

Figures 17 (a) and (b) are the diagrams that plotted the dissolution rate \( v_{sm} \) shown in Figs.14 (a) and (b) on a natural logarithmic scale against \( \eta \) (\( T = 298K \)) and \( 1/T \) (\( \eta =176mV \)) respectively. It can be seen that the values of \( (\eta \sigma/kT) \) and \( ((\Delta H_0 - \alpha \eta \sigma)/k) \) are estimated by comparing the Eq.(33) with gradients of the linear relationships drawn in Figs. 17 (a) and (b), because the overpotential and temperature dependences of \( \sigma \) in the range of 156 mV to 186 mV are assumed to be a little. Thus, \( \alpha \) and \( \Delta H_0 \) were also obtained from overpotential dependence of vertical dissolution rate \( v_{sm} \) of surface at higher overpotentials and temperature dependence of that. The estimations are shown in Table 3, showing \( \alpha \) and \( \Delta H_0 \) are in good agreement with those values in Table 2.

![Figure 17](image-url)

**Figure 17.** Vertical dissolution rate \( v_{sm} \) at dislocation-free site of surface on a natural logarithm scale. (a): The plot against overpotential \( \eta \); (b): The plots against the inverse \( 1/T \) of temperature.
According to atomistic dissolution model of a crystal surface illustrated in Fig.15, the relation of \( \theta = \tan^{-1}(a/\lambda) = \tan^{-1}(v_{sm}/v_{h}) \) is lead from Eq. (17), which represents the inclination angle of surface to (111) face. Since it is approximately given by \( \theta \approx \theta^* = \tan^{-1}(v_{sm}/v_{w}) \) with use of Eqs.(20) and (26), the values of \( \theta^* \) obtained from Fig.14 were plotted against \( \eta \) and \( T \) in Figs.18 (a) and (b). It can be seen that the tendencies of change in \( \theta^* \) against \( \eta \) and \( T \) are not clear and not reasonable. The average value of \( \theta^*_{av} \) is 2.1×10^-2 rad, which is a little large compared to a deviation 8.7×10^-3 rad from [111] direction that was aimed when we prepared the surface of specimen as described in Section 2.1. This is probably attributed to the fact that actual surface exposed to electrolyte solution was slightly spherical as a whole and was having microscopic swells. That is, the variation of their values seems to be due to experimental error. Thus the vertical dissolution rate at dislocation-free site of surface is assumed to be given by retreat rate of the surface steps which preexists on the prepared surface, which gives following relation:

\[
\beta k_s = \frac{1}{\lambda x_0} = \frac{b}{x_0} \cdot \frac{\tan \theta^*}{ab}
\]

Accordingly \( \beta (b / x_0) \) is calculated from \( \beta k_s \) in Table 2 by using Eq. (34), which is shown in Table 3 where assumed \( \theta^* = \theta^*_{av} \) (0.021 rad).

![Figure 18. Inclination of surface to (111) face, which is given by \( \theta^* = \tan^{-1}(v_{sm}/v_{w}). \)](image)

<table>
<thead>
<tr>
<th>( \alpha )</th>
<th>( \Delta H_0/eV )</th>
<th>( v_{sm}/m \text{s}^{-1} )</th>
<th>( \beta b/x_0 )</th>
<th>( v_{ed}/m \text{s}^{-1} )</th>
<th>( \Delta G^*/eV )</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.85</td>
<td>0.33</td>
<td>1.6×10^{-9}†</td>
<td>0.034†</td>
<td>5.7×10^{-9}†</td>
<td>0.16†</td>
</tr>
</tbody>
</table>

†the value at 298K,

**Table 3.** Estimations of kinetics parameters controlling dissolution rate at edge dislocation site of surface of copper crystals.

### 4.3. Vertical dissolution rate at dislocation site

#### 4.3.1. Estimation of the critical free energy change for nucleation of two-dimensional pit

As mentioned in Section 4.1.2 the dissolution rate \( v_{ed} \) at edge dislocation site is expressed by Eq. (23), but if Eq. (14) is applied it is reduced to

\[
 v_{ed} = aK_s C_{\text{Cl}^-}^0 \cdot \frac{2(1-\alpha)}{C_{\text{CuCl}_2}^0} \cdot \left(\frac{C_{\text{Cl}^-}^0}{C_{\text{Cl}^-}^*}\right)^2 \cdot \exp \left( \frac{-\Delta G_{ed}^*/eV + \Delta H_0 - \alpha \eta g \eta}{kT} \right) \cdot \nu.
\]

(35)
Accordingly, if we assume \( \text{Cl}^-/\text{Cl}^0 = 1 \), \( \Delta G_{\text{ed}}^* \) is given by

\[
\Delta G_{\text{ed}}^* = -kT \ln \left( \frac{v_{\text{ed}}}{aK_s C_{\text{Cl}^-}^{1/2} C_{\text{CuCl}_2}^{1/2} \alpha} \right) - \Delta H_0 + \alpha n \eta. \tag{36}
\]

Thus \( \Delta G_{\text{ed}}^* \) under each condition was estimated by Eq. (36) with use of experimental value of \( v_{\text{ed}} \) as well as estimations of \( \alpha \) and \( \Delta H_0 \) which were obtained in Section 4.2.2. The \( \Delta G_{\text{ed}}^* \) estimated with use of two assumed values of undetermined constant \( K_s \) for a specified condition (\( \eta = 176 \text{ mV} \) and \( T = 298 \text{K} \)) are shown together with the values \( \alpha \) and \( \Delta H_0 \) in Table 3, where \( a = 2.09 \times 10^{-10} \text{ m} \) and \( \nu = 6.21 \times 10^{12} \text{ s}^{-1} \) were used.

According to the precedent theoretical study (Schaarwächter 1965), in which the conditions for the formation of visible etch pit at dislocation site were investigated on the basis of a proposed nucleation model, the critical free energy change is estimated to be 0.115 eV. The present estimation of \( \Delta G_{\text{ed}}^* \) approximately consists with that value as shown in Table 3, though the exact value of \( K_s \) can not be evaluated in this study. This is seemed to be reasonable as described in Appendix A2.

On the other hand, however, it was admitted that the value of \( \Delta G_{\text{ed}}^* \) varies with overpotential and temperature as mentioned below.

### 4.3.2. Overpotential and temperature dependences of \( \Delta G_{\text{ed}}^* \)

Figures 19 (a) and (b) are the diagrams that plotted the square root of \( \Delta G_{\text{ed}}^* \) estimated assuming \( K_s = 1 \) by Eq. (36) against \( \eta \) and \( T \) respectively. It can be seen that \( \Delta G_{\text{ed}}^{*1/2} \) is not constant but changes in different manners with increases in \( \eta \) and \( T \). The reason for this is probably that \( \Delta G_{\text{ed}}^{*1/2} \) is proportional to the interfacial energy \( \gamma \) as known from Eqs. (24) and (25).

\[\text{Figure 19. Square root of the critical free energy change for the formation of a two-dimensional pit. (a): The overpotential dependence; (b): The temperature dependence.}\]
It is known that the interfacial energy varies with electrode potential according to so-called electrocapillary curve (Tamamushi, 1967). Therefore, the change in $\Delta G_{ed}^{*1/2}$ with $\eta$ is surmised to be due to the potential dependence of $\gamma$ because the overpotential dependences of the undersaturation $\sigma$ and therefore that of $\Delta \mu = ne\eta_a = -kT\ln(1-\sigma)$ in an overpotential range of 156 to 186 mV are assumed to be a little as described in Section 4.2.2. This is supported by the fact that Fig.19 (a) indicates a quadratic dependence similar to the electrocapillary curve. Also, it is inferred from Fig.19(b) and Eq.(24) that $\gamma$ should increase with an increase in $T$, because $\Delta \mu$ tend to increase with increase in $T$. This is probably attributed to a decrease in specific adsorption of anion accompanied by an increase of interfacial energy with rising of temperature.

The overpotential dependence of log $v_{ed}$ is in accelerative, and somewhat different from that of both log $v_{sm}$ and log $v_{w}$. Also the increasing rate of log $v_{ed}$ with increase in temperature is smaller than that of both log $v_{sm}$ and log $v_{w}$ as shown in Figs.14 (a) and (b). The reason for this seems to be attributed to the overpotential and temperature dependences of the interfacial energy of the electrode surface as mentioned above.

5. Conclusions

Following conclusions were obtained from the results and discussion:

1. The transfer coefficient, activation enthalpy and surface concentrations of the ions which control the dissolution reaction were estimated from measurements of the relationships between steady anodic current densities and applied overpotentials when copper crystals are dissolved in an electrolyte solution under potentiostatic electrolysis.
2. The values of a supplementary factor and kink density affecting rate constant of dissolution reaction were examined.
3. The dissolution rate at edge dislocation site of (111) surface of copper was discussed quantitatively by a nucleation model of two-dimensional pit based on the classical nucleation theory.
4. The present estimation of the critical free energy change $\Delta G_{ed}^{*}$ for nucleation of a two-dimensional pit at edge dislocation site reasonably consisted with the evaluation by the precedent study.
5. The overpotential and temperature dependences of dissolution rate at edge dislocation site were somewhat different from those dependences of dissolution rate at dislocation-free site. The reason for this is probably that $\Delta G_{ed}^{*}$ changes according to the overpotential and temperature dependences of interfacial energy.

6. Appendix

A1. Kinds of dislocation etch pits and their characters

The surface of copper specimen on which some small glass spheres 300 $\mu$m in diameter were dropped beforehand was anodically etched by the present method. Fig.20 (a) is an optical
micrograph of dissolved surface in which Rosseta pattern composed of dark and light etch pits was formed at the portion that was hit by a small glass sphere. This proves that dark and light etch pits are formed at the sites of positive and negative edge dislocations respectively because the six arms of Rosseta pattern are composed of rows of a pair of positive and negative edge dislocations.

**Figure 20.** Optical micrographs for identifications of dark and light pits. The surfaces dissolved by the present method; (a): Rosseta pattern composed of etch pits; (b): a distribution of etch pits. (c): etch pits formed by a chemical etchant in the same portion as that observed in b.

In another experiment, the surface of prepared copper specimen was anodically etched first by the present method, and a distribution of etch pits were observed by the optical microscope. Subsequently after electropolished the etched surface of specimen, the surface was etched for 10 s by a modified Young's etchant prepared by Marukawa (Marukawa, 1967), and the same portion as the previous portion was observed. Figs.20 (b) and (c) are a pair of optical micrographs of the surfaces etched by such two methods. It has been reported by Marukawa that the dark (deep) and light (shallow) pits are formed at screw dislocations and edge dislocations on the surface etched by the modified Young’s etchant respectively. Accordingly it can be seen that the light etch pits are formed at the sites of screw dislocations on the surface that was anodically etched by the present method, by comparing the kinds of etch pits which are observed in these micrographs. Thus Table 4 is obtained concerning dislocation characters related to dark and light etch pits.

<table>
<thead>
<tr>
<th>Etching</th>
<th>Edge dislocation (positive)</th>
<th>Edge dislocation (negative)</th>
<th>Screw dislocation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chemical†</td>
<td>Light</td>
<td>Light</td>
<td>Dark</td>
</tr>
<tr>
<td>Electrolytic†</td>
<td>Dark</td>
<td>Light</td>
<td>Light</td>
</tr>
</tbody>
</table>

*† by modified Young’s etchant, †† present method

**Table 4.** Relations between the dislocation characters and the kinds of etch pits which are formed by two etching methods
In this work, the depth and width of the dark (deep) pits were measured to investigate the dissolution amounts at positive edge dislocation sites.

A2. Estimation of undetermined constant \( K_s \)

As described in the Section 4.1.3, if the separation rate \( r \) of an atom at nucleation of two-dimensional pit is a quantity similar to the dissolution rate of an atom from kink site of surface, it would need to take account of supplementary factor \( \beta \) affecting the exchange current density as a parameter involved in the separation rate \( r \). Then the dissolution rate \( \dot{v}_{ed} \) at edge dislocation site derived from the nucleation rate Eq. (21) is represented afresh by

\[
\dot{v}_{ed} = a \beta Z C_{Cl^-} 2^{(1-\alpha)} C_{CuCl^-}^{\alpha} \nu \exp \left( -\frac{\Delta G_{ed}^* + \Delta H}{kT} \right). \tag{37}
\]

Thus, we assume that the undetermined constant \( K_s \) is approximately given by a relation:

\[
K_s = \beta Z. \tag{38}
\]

We have assumed in the Section 2.7 that the exchange current density \( i_0 \) is given by Eq. (7) for simplification, but to be exact \( i_0 \) should be expressed with use of the activities of the ions involved in the electrode reaction instead of the concentrations. Also, transmission coefficient should be taken account of as pre-exponential factors in Eq. (7). Therefore it is generally hard to estimate \( \beta \) including some unknown factors. However, concerning \( \beta \) of the present electrode reaction, \( \beta \, (b/x_0) = 0.034 \) was estimated experimentally as shown in Table 3. Also it can be seen from an observation of etch pit by optical microscope that surface steps have a structure along a crystallographic direction of the crystal. Accordingly if \( (b/x_0) \) is assumed to be a quantity of 0.02 to 0.2, it would give an estimation of \( \beta = 0.17 \sim 1.7 \).

On the other hand, if we assume the free energy change \( \Delta G_{ed} \) \((j)\) for formation of a two-dimensional pit consisting of \( j \) vacancies at edge dislocation site, it is written as

\[
\Delta G_{ed}(j) = 2\gamma (\pi a \Omega)^{1/2} j^{1/2} - \frac{\Delta \mu}{p'} j \tag{39}
\]

Then the critical size \( j^* \) of two-dimensional pit and the critical free energy change \( \Delta G_{ed}^*(j^*) \) are given by

\[
j^* = p'^2 \pi a \Omega \gamma^2 \Delta \mu^2 \quad \text{and} \quad \Delta G_{ed}^*(j^*) = p'^2 \pi a \Omega \gamma^2 \frac{\Delta \mu^2}{\Delta \mu} \tag{40}
\]

respectively. It can be seen that \( \Delta G_{ed}^* (j^*) \) is expressed by the same relation as Eq. (24), and that the factor \( p' \) has the same contents with Eq. (25), that is, \( p' = p \). Then, Zeldovich factor is expressed from the definition (Toschev, 1973) by

\[
Z = \sqrt{-\frac{1}{2\pi kT} \left( \frac{\partial^2 \Delta G_{ed}(j)}{\partial j^2} \right)_{j^*}} = \sqrt{\frac{1}{4\pi kT} \left( \frac{\Delta \mu^2}{p'^2 \Delta G_{ed}^*} \right)}. \tag{41}
\]
Accordingly, $Z = 0.76$ is estimated, if $p = 0.18$ (Schaarwächter 1965), $\Delta \mu = 0.027$ eV ($\sigma = 0.65$) (Imashimizu, 2011), $\Delta G^{\circ} = 0.12$ eV (Table 3) and $kT = 0.0257$ eV ($T = 298K$) are used. Thus $K_s = 0.13$-1.3 is estimated from Eq. (38), which suggests the reasonability of the assumed value of $K_s$ shown in Table 3.

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**7. References**


