Role of Chemical Steps in the Electrochemical Reduction of Metal Complexes

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Abstract
The role of chemical steps in electrochemical processes involving metal complexes is considered invoking the model of mass transport of chemically interacting particles. The quantitative description is based on the second Fick’s law equations supplemented with the respective kinetic terms. Simple differential equations containing no kinetic terms are obtained for total concentrations of metal, ligand, proton donors and acceptors. Their surface concentrations can be determined ignoring the kinetics of chemical steps and using experimental voltammetric data transformed according to the analytical expressions obtained. In the case of labile systems, linear Tafel plots normalized with respect to the surface concentration of the electrochemically active complex can be constructed. The application of the suggested algorithm is illustrated by the analysis of experimental data obtained for Cu(II) reduction in glycolic acid solutions and hydrogen evaluation in the acetate system.

Keywords
Metal complexes, Electroreduction, Chemical steps, Lability

Introduction
Metal complexes still attract extensive attention, since they are widely distributed in both nature and in practical applications. The electrochemical processes used in such important areas as electroanalysis, electroplating, etc., are of particular note. When solving various problems of theoretical and applied research, one is faced with the specifics of these systems, namely, with the variety of chemical interactions (often rather numerous), attending the electrochemical charge transfer step. The simplest example of the process proceeding through the so-called CE mechanism (chemical step + electrochemical step) could be the reaction $X \rightarrow O + n\ e \rightarrow R$, wherein the electrochemically active substance $O$ is formed from the electrochemically inert substance $X$ during the preceding chemical reaction. By far the most rigorous quantitative characteristics of this mechanism were obtained when the single preceding chemical step is of the first (or pseudo-first) order. When increasing this parameter or the number of chemical steps, analytical relations are no longer available.

Nevertheless, most real electrochemical systems are significantly richer in various interactions [1]. Consideration of these phenomena in general, is a rather complex task that requires involvement of non-linear differential equations and the use of the kinetic characteristics which are not always available. In this connection, there is a need for the acceptable simplifications of the problem and finding the most favorable ways to analyze the experimental data.

In this paper, we analyze the ways of taking into account the kinetics of chemical interactions, the applicability of which would not be limited in the number of system components and chemical steps. To solve the certain problems, such feature as the system lability is involved. Application of the results of theoretical analysis is illustrated by the data obtained in the experimental study of some systems.

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Theoretical Notes

Analysis of experimental voltammetric data usually assumes the use of kinetic equations that contain not only the kinetic parameters of the charge transfer step, but also the surface concentrations of species taking part in the electrochemical reaction. To determine these quantities, it is necessary to apply an adequate model in which the regularities of mass transfer of chemically interacting particles would be taken into account. As we will see from the following, it becomes necessary to discuss in more detail some concepts that are involved in this theoretical model.

Mass-Transport of chemically interacting species

Let us suppose that the reduction of the Electrochemically Active Complex (EAC)

$$M_L^{n+} + ne \rightleftharpoons M + pL$$

(1)

Proceeds in the system under linear diffusion conditions. Concentrations of the components, as time, $t$, and coordinate, $x$, functions, can be determined by solving differential equations, representing the second Fick’s law and supplemented with terms that take into account the kinetics of chemical interactions. Detailed formulation of this problem is given below in the Appendix.

We focus on the interesting, in our view, conclusion that follows from the analysis. It is noteworthy that a very simple differential equation

$$\frac{\partial c(x,t)}{\partial t} = D \frac{\partial^2 c(x,t)}{\partial x^2},$$

(2)

containing no kinetic terms, is valid for the total concentration of metal, $c_{M}$, ligand, $c_L$, or proton donors and acceptors, $c_p$. Its usage is tempting, since solutions of such equations supplemented with corresponding initial and boundary conditions have been well-expounded in literature. The initial condition (at $t = 0$) is commonly treated as an equilibrium state with constant (bulk) concentrations of species over the whole area of the system. The same is valid for the respective boundary conditions at $x \geq \delta$ (outside the $\delta$-thick diffusion layer). Another boundary condition follows from the first Fick’s law and correlates the flux of the EAC with the current density. When Eq. (2) deals with the total metal concentration ($c = c_M$), there is no necessity to preset which species is electrochemically active. In any case, the current density

$$i = -nFD \left. \frac{\partial c_M(x,t)}{\partial x} \right|_{x=0},$$

(3)

Where $n$ is a number of electrons involved in the reaction (1) and $F$ is a Faraday constant.

Since the ligand is supposed to be electrochemically inert,

$$\frac{\partial c_L(x,t)}{\partial t} \bigg|_{x=0} = 0.$$

(4)

The problem formulated in such a way has the known analytical solution:

$$\Delta c_M(t) \equiv c_{M,b} - c_{M,t}(t) = \frac{1}{nF\sqrt{D}} \int_0^t \frac{i(t-u)}{\sqrt{u}} \psi(u) du,$$

(5)

Where the variable $t$ in the time-dependent current density $i(t)$ is replaced by the auxiliary variable $(t - u)$. The function

$$\psi(u) = 1 + \sum_{m=0}^{\infty} (-1)^m \exp \left( -\frac{m^2 \delta^2}{Du} \right).$$

(6)

Takes into account the existence of $\delta$-thick diffusion layer. We want to point out the importance of the equation (5): since it contains no kinetic characteristics of chemical steps, this relationship is suitable for determining the total surface concentration both in labile and inert systems. However, no information concerning individual components of the system follows from above equations. Therefore, it is necessary to formulate extra interrelations between their concentrations.

Lability of electrochemical systems

When chemical steps are sufficiently fast, concentrations might be treated as close to equilibrium ones. Then these quantities might be defined by the respective equilibrium constants. To determine whether such an approach is justified, it is necessary to consider the system lability. Formulation of lability criteria is based on comparing limiting fluxes controlled by finite rates of dissociation of complex species to their limiting diffusional fluxes [2]. As a result, this criterion depends on many parameters including diffusion coefficients, the rate constants of chemical steps, the thickness of diffusion and reaction layers, the size of micro-electrodes, etc. [2-11]. Again, strict analytical expressions have been derived mainly for the first-order chemical step. This obstacle can be circumvented to a certain extent by using Eq. (5).

From the standpoint of lability, we can distinguish two limiting cases concerning the limiting currents ($i_d$) that occur in the region where the surface concentration of the EAC approaches zero. If the system is highly labile, the loss in the surface concentration of the EAC, caused by the process (1), is easily compensated by fast EAC formation during the corresponding chemical stages. Then, the concentrations of all particles containing metal ions tend to zero in the $i_d$ region, that is, $c_{M,b} \rightarrow 0$. In this case, the maximum value of $\Delta c_{M,b}$ determined by equation (5), can reach the total metal concentration in the bulk of solution:
\[ \Delta c_M(t)_{\text{max}} = c_{M,b}. \]  

(7)

Hence, the fulfillment of condition (7) is an indispensable sign indicating a labile character of the system. In this case, the relations between surface concentrations might be determined on the basis of step-wise, \( K_j \) or cumulative, \( \beta_j \), stability constants of complexes defined as

\[
K_j = \frac{k_j}{k_{-j}} = \frac{[ML^{n+}_{j}]}{[ML^{n+}_{j-1}][L]},
\]

(8)

\[
\beta_j = \frac{[ML^{n+}_{j}]}{[M^{n+}][L]^j},
\]

(9)

Where \( k_j \) and \( k_{-j} \) are the respective rate constants (see Appendix).

A completely different situation arises when the system is highly inert. In this case, extremely slow chemical stages are not able to restore the loss of the EAC at the electrode surface. As a result, only the EAC surface concentration varies and drops to zero in the region of the limiting current. Then, the condition (7) changes to

\[
\Delta c_M(t)_{\text{max}} = c_{EAC,b}.
\]

(10)

Though chemical reactions are supposed to be very slow, their kinetics should be taken into account in the certain cases, e.g., when the equilibration time of the electrochemical system is estimated. However, in this limiting case of lability, the charge transfer step can be analyzed taking no account for preceding chemical steps.

**Concept of the diffusion layer thickness**

To obtain the surface \( c_s \) from experimental \( i(t) \) data transformed according to the Eq. (8), it is necessary to estimate the values of two parameters: the average diffusion coefficient and the thickness of the Nernst-type diffusion layer. According to this model, diffusion takes place in a layer of stagnant solution close to the electrode|solution interface. It has been established by means of different methods that similar structures actually form under the natural convection conditions. Their thickness makes up 0.02-0.05 cm; however, it is rather difficult to define strictly the value of \( \delta \) because it depends on various factors.

This quantity can be eliminated under the steady-state conditions, when \( \partial c/\partial t = 0 \). Then the equation

\[
c_s = c_b \left( 1 - \frac{i}{i_d} \right),
\]

(11)

Can be used that follows from the well-known relation between the limiting current density, \( i_d \), and \( \delta \):

\[
i_d = \frac{nFDc_b}{\delta}.
\]

(12)

In the case of labile systems, the quantity \( c \) in Eq. (11) assumes the value of the total concentration \( c_{M,b} \). Differently, \( c = c_{EAC,b} \) when the system is highly inert.

Certain complications arise in the case of time-dependent (non-steady-state) electrolysis. In particular, it was established that the \( \delta \) varies with the potential sweep rate \( v \) in the Linear Potential Sweep (LPS) voltammetry. The empiric regularity of \( \delta \sqrt{v} \) was const previously determined for the Fe(CN)\(_6\)^{3-}/Fe(CN)\(_6\)^{4-} system [12] and was found to be also valid for Cu(II) solutions containing glycolic and gluconic acids [13,14]. The attempts have been made to substantiate this regularity theoretically [15-18]. Though the models used support a linear dependence between \( \delta \) and \( \sqrt{v} \), nevertheless the \( \delta \) values following from these theoretical models are one order of magnitude lower than experimental quantities. More realistic data follow from another approach [19] based on Levich’s view of microscopic chaotic motion that takes place in the viscous sub-layer adjacent to any wall or electrode surface. The \( \delta = 230 \mu m \) was found from analysis of chronoamperometric transients obtained for one-electron reversible oxidation of Fe(CN)\(_6\)^{3-}; this result is in line with the respective data given in Figure 1. Slopes of lines, which approximate \( \delta \cdot v \) dependences in logarithmic coordinates, range from 0.48 to 0.54.

**Concept of the effective diffusion coefficient**

It was accepted in differential equations given in the Appendix that diffusion coefficients of all the particles are equal. If the size of complexes or ligands is similar, such an assumption is justified. However, it is known that H\(^+\) and OH\(^-\) ions distinguish themselves from other by their greater mobility, which is determined by the
so-called structural diffusion (Grotthuss mechanism) involving a shift of structural units as a result of rearrangement of the protons without a considerable shift of water molecules themselves.

Theories of various electrochemical analysis methods (polarography, chronopotentiometry, RDE, etc.) state that the thickness of the diffusion layer depends on $D$. Hence, if diffusion coefficients of certain particles differ, thickness of their diffusion layers must be different. When applying such images to the systems of metal complexes, contradictions arise, which are especially obvious in the case of labile complexes. Then, individual $\delta_{j}$ with its own $D_{j}$ should exist for each particle. Let us imagine that the certain species has an increased mobility. Then, with diffusion mass transport going on, their excess will appear in a certain region of the solution, causing the respective shifts of chemical equilibria. As a result, some part of more mobile particles will have to react with other components of the solution. Formally the same result can be obtained by assigning a lesser $D$ to these particles. If we investigate the particles of the reduced mobility, on the contrary, their $D_{j}$ should be increased. Thus, the mechanism, which allows one and the same effective $D$ to be assigned to all the particles, appears in the labile systems. This mechanism was studied in more detail in [20]; similar conclusions were also drawn in investigating combined $\text{H}^{+}$ and $\text{HSO}_{4}^{-}$ diffusion [21,22].

The effective $D$ depends on the individual diffusion coefficients of complex species and their molar fractions in the solution [6,8,23]. To estimate its value, the following relation is usually used:

$$D^m = \sum_{j} D^m_{j} \chi_{j},$$

(13)

Where $\chi_{j}$ is a molar fraction of the certain complex species. The index $m$ depends on the experimental conditions; it is equal to 1 (steady-state electrolysis), $\frac{1}{2}$ (polarography), $\frac{3}{2}$ (RDE voltammetry), etc. Other comparable expressions are also available. For instance, when $\text{ML}^{n}_{b}$ complexes prevail in the system, the equation

$$D^m = \frac{D_0 + D_N \beta_{N} [L]^N}{1 + \beta_{N} [L]^N},$$

(14)

is obtained [23]. Note that the analysis of $D$ dependence on $[L]$ was used to determine $\beta_{N}$.

The above concept seems to be acceptable in the case of labile systems. The most convenient is to obtain this effective $D$ from the experimental data (limiting currents, transition times, etc.), which are determined by the diffusion characteristics. However, the use of this approach in the case of inert systems remains questionable.

**Determination of kinetic parameters**

Thermodynamics allows us to treat any metal-containing species as a possible EAC. However, there is a lot of experimental data (see, e.g. [1]) testifying that the EAC is liable to contain less ligand particles than the more stable complex predominating in the solution. The removal of ligands from its inner coordination sphere promotes adsorption and a decline in the potential barrier of electron transfer. Thus, kinetic factors, and the activation energy first of all, should be responsible for the electrochemical activity of complexes. Special methods have been suggested and successfully applied for establishing the EAC composition [1,24].

Often a single EAC is found in a certain potential region. Then the kinetic equation of the process (1) takes the form

$$i = i_{0} \left[ \frac{[L]^p}{[L]^b} \exp\left(\frac{\alpha_{a} n F}{RT} \eta\right) - \frac{[\text{ML}^{n+}_{p}]_s}{[\text{ML}^{n+}_{p}]_b} \exp\left(-\frac{\alpha_{c} n F}{RT} \eta\right) \right],$$

(15)

Where $i_{0}$ is an exchange current density, $\alpha_{a}$ and $\alpha_{c}$ are anodic and cathodic charge transfer coefficients respectively ($\alpha_{a} + \alpha_{c} = 1$). At sufficiently high cathodic overvoltages ($\eta_{c} = -\eta$), i.e. when $\eta_{c} >> RT / nF$,

$$\ln |i| = \ln i_{0} + \ln \left[ \frac{[\text{ML}^{n+}_{p}]_b}{[\text{ML}^{n+}_{p}]_s} \right] + \frac{\alpha_{a} n F}{RT} \eta_{c}.$$  

(16)

Kinetic parameters are usually obtained from experimental data, transformed according to the Tafel-type equation. However, it is necessary to point out that the dependence of $\ln |i|$ vs. $\eta_{c}$ given by Eq. (16), is not linear, as is usually the case, since the EAC surface concentration is not constant and depends on $i$ or $\eta$. This relation can be linearized through the replacement of the quantity $i$ with the normalized current density, viz.

$$i_{\text{norm}} = \frac{[\text{ML}^{n+}_{p}]_s}{[\text{ML}^{n+}_{p}]_b}.$$  

(17)

Again, we have 2 cases when the EAC surface concentration (the value of $[\text{ML}^{n+}_{p}]_s$) is obtained relatively easy. For labile systems, we use the $c_{ML}$ quantity established with Eq. (5), and material balance equations together with expressions (9) for respective stability constants. Remember that $c_{ML} = c_{MLL}$ when the ligand is electrochemically inactive. If the system is highly inert, the $c_{\text{EAC}}$ is obtained immediately with Eq. (5), considering that this relationship reflects only changes in EAC surface concentration. Further analysis of Normalized Tafel Plots (NTP) is usual.

**Experimental examples**

Electrochemical experiments were carried out using the usual electrochemical equipment: a potentiostat/galvanostat and a conventional three-electrode cell. Linear Potential Sweep (LPS) voltammograms were recorded at a potential sweep rate ranging from 0.01 to 0.2 V s$^{-1}$. In
all cases, cathodic scans were applied, starting with the open-circuit potential.

To prepare the working electrodes, a 1 cm² platinum disc was coated with 5-7 μm thick copper in the solution containing (g dm⁻³): CuSO₄·5 H₂O - 250, H₂SO₄ - 50. A polycrystalline layer with well-exhibited crystallographic edges and faces was formed. Detailed characteristics of surface morphology are given [25]. The electrode potential was measured in reference to the Ag | AgCl | KCl (sat) electrode and was converted to the standard potential. The data for iₚ were obtained in Cu(II)-free solutions.

**Figure 2: LPS voltammograms obtained for Cu(II)-glycolic acid system at different potential sweep rates.** Dependencies of the left and right peak currents (iₚ₁ and iₚ₂, respectively) vs. \( \sqrt{\nu} \) are given in the inset.

Experiments carried out at different pH showed [26], that the rate-determining step might be the transfer of the first electron on the Cu(II) aqua-complex. Then the kinetics of Cu(II) reduction is given by the equation [1]

\[
i = 2i_{01} \left\{ \exp \left( \frac{2 - \alpha_{c1} F}{RT} \eta \right) \frac{[Cu^{2+}]_s}{[Cu^{2+}]_b} \right\} \exp \left( -\frac{\alpha_{c1} F}{RT} \eta_c \right)
\]

and the expression for NTP becomes as follows:

\[
\ln |i| = \ln 2i_{01} + \ln \frac{[Cu^{2+}]_s}{[Cu^{2+}]_b} + \frac{\alpha_{c1} F}{RT}\eta_c,
\]

Where \( i_{norm} = |i| \frac{[Cu^{2+}]_b}{[Cu^{2+}]_s} \).

Experimental data, obtained at different \( \nu \) and transformed into NTP, fall on one general line (Figure 4). The values of kinetic parameters match the similar data reported in [27] for 0.1 M Cu(ClO₄)₂ solutions.

**Figure 3: An experimental voltammogram (curve 1, ordinate to the left) and its transform according Equation (8) (curve 2, ordinate to the right).**

Acetic acid system—an extension of the theoretical model

The approach considered in the present paper is also applicable to the hydrogen evolution that often attends the electrodeposition of metals. When the plating solutions contain such ligands as hydroxy acids, these substances can generate hydrated protons (H₃O⁺ ions) establishing the conditions for the CE mechanism discussed above. Then, hydronium ions are formed additionally in chemical reactions, which precede the electron transfer step.
Considerable amount of data, accumulated in this area, created the preconditions to use the simplest solutions of weak acids for experimental verification of certain regularities following from theoretical models. For this purpose, we chose acetic acid solutions due to the relative simplicity of homogeneous processes in them and availability of sufficiently reliable kinetic data.

Investigations of hydrogen evolution in acetate solutions were carried out at 3 < pH < 5 by the LPS voltammetry using copper and platinum electrodes [28, 29]. It was found that voltammograms display current peaks, which take a shape characteristic of other electrochemical processes including the reduction of metal complexes. It was also established that LPS current peaks vary linearly with \(\sqrt{v}\) and with the total concentration of proton donors and acceptors that is in line with the theoretical predictions given above.

For interpretation of the experimental data, we appealed to the model of CE mechanism involving two consecutive steps: the preceding chemical reaction \(HA \rightleftharpoons H^+ + A^-\) and the charge transfer \(H^+ + e \rightarrow \frac{1}{2} H_2\). Procedures, described above, were successfully used in the analysis of experimental data, provided that the behavior of hydronium ions is similar to that of M\(^{n+}\) and acetate-anions play role of ligand.

Surface concentrations of individual species (acetic acid molecules HA, hydronium ions H\(^+\) and acetate anions A\(^-\)) were obtained using material balance equations and the expression of the equilibrium constant:

\[
K_a = \frac{k_1}{k_2} = \frac{[H^+][A^-]}{[HA]}, \tag{18}
\]

To obtain the bulk concentration of hydronium ions from pH, the activity coefficient \(\gamma_{H^+} = 0.7\) was used. These simulations and other computations that follow were performed with \(K = 2.7 \times 10^{-5}\), \(k_1 = 3.46 \times 10^6\) s\(^{-1}\) and \(k_2 = 1.28 \times 10^{11}\) dm\(^3\) mol\(^{-1}\) s\(^{-1}\) [30]. As a result, the linear normalized Tafel plots were obtained. An example of such NTP is shown below in the inset of Figure 5 (the data for pH 3). It can be assumed that the data obtained at different \(v\) fall on one general line.

Further investigations have shown that unusual double current peaks are observed at 1.5 < pH < 2.5 (Figure 6). An extra current maximum emerges at -0.8 V and becomes detectable already at pH 2.5. The shape of these maxima is distorted. In appearance they resemble the superposition of two peaks, located quite close to each other.
It was of interest to find out to what extent the approach used is applicable in this more complicated case. We performed the necessary procedures to verify whether the system is labile enough. An example of the experimental voltammogram and its transform by Eq. (5) is shown in Figure 5. The function $\Delta c_i(t)$ increases with time approaching a characteristic plateau, which is the limit of $\Delta c(t)$ and should not exceed the volume concentration $c_{uls} = 34.3$ mM. Indeed, the plateau of $\Delta c_i(t)$ function is located at this value (curve 2 in Figure 5) when the effective $D = 1.1 \times 10^{-5}$ cm$^2$s$^{-1}$. This quantity is very close to $D = 1.2 \times 10^{-5}$ cm$^2$s$^{-1}$ given in [31] for 25°C. Further rise in this curve, as well as in the experimental voltammogram (curve 1), is associated with the beginning of a new process, viz. the discharge of water molecules.

An example of normalized Tafel plots obtained at pH 1.9 is shown in the inset of Figure 5. Again, the data, obtained at different $\nu$, are close enough. Although the transformed data slightly meander with $E$, we found it possible to apply a single fitting line that turned out to be almost parallel to the NTP obtained at pH 3.0. Kinetic parameters following from NTP analysis are also close: $\alpha_c = 0.72 \pm 0.04$ and the effective $i_c = 0.2-0.4$ nA cm$^{-2}$.

Though the literature on hydrogen evolution is very rich, kinetic data relating to the copper electrode are rather slender. The value of $\alpha = 0.7$ was detected for perchloric acid solutions over a wide region of temperatures (158-300 K) [32]. For less acidic acetate solutions, we determined that $\alpha = 0.64$, and the exchange current density of the order of 1 nA [29]. Similar $\alpha _c$ values were also obtained [1] for glycolic, malic, tartaric and gluconic acid solutions containing 0.3-0.5 M Na$_2$SO$_4$ as a supporting electrolyte. Formation of significant amount of HSO$_4$ ions, which are additional proton donors, could be one of the reasons which led to the higher exchange current densities reaching ~10 µA cm$^{-2}$. Analysis of the RDE voltammetric data, obtained for 0.03 M H$_3$BO$_3$ solution (pH 5.7) and performed on the basis of Koutecky-Levich equation [33], produced $i_c = 3.46 \times 10^{-7}$ A cm$^{-2}$ and $\alpha_c = 0.5$ (the latter value follows from the cathodic Tafel constant equal to 120 mV per decade).

To compare quantitatively experimental and theoretical voltammograms, we performed simulations using the entire set of differential equations. Since the rate constants $k_1$ and $k_2$ were employed, no assumptions on system lability were necessary. An example of results, obtained with standard rate constant $k_2 = 2 \times 10^{9}$ cm$^{-1}$s$^{-1}$ and $\alpha_c = 0.62$ is shown in Figure 7 (values of $D$, $K_s$, $k_s$, and $k_2$ are given above). A high cathodic overvoltage of irreversible process arises from low $k_1$ values. Peak heights and their ratio are also different.

Fitting procedures made it possible to obtain a reasonable agreement between simulated and experimental data, an example of which is shown in the inset of Figure 7. Here, $k_1 = 4 \times 10^{-12}$ cm$^{-1}$s$^{-1}$ and $\alpha_c = 0.75$. These parameters are in line with kinetic parameters obtained from NTP analysis. It was also found that the results of simulation are highly dependent on the pH: a change of pH by 0.1 leads to changes in the simulated voltammograms exceeding the reproducibility of experimental data. The best agreement between the data in the inset is obtained when the simulation for pH 1.97 is performed. It should be mentioned that quite a large amount of initial data $(D, K_s, k_s, k_i, k_1, k_2, \alpha_c)$ was used. Each of them has been identified with a certain error. Besides, experimental errors associated with measurements of current-voltage characteristics and pH should be taken into account. All this bearing in mind it seems possible to conclude that the CE model quite adequately renders the electrochemical behavior of the acetate system at low pH.

**Concluding Remarks**

To take into account the role of chemical stages in electrochemical processes involving metal complexes, a mass transfer model of chemically interacting particles is needed. This problem is solved using a system of differential equations reflecting Fick’s second law and supplemented with terms that take into account the kinetics of the chemical stages. Simple differential equations containing no kinetic terms are obtained for total concentration of metal, ligand, as well as proton donors and acceptors. Their surface concentrations can be determined from experimental voltammetric data transformed according to the analytical expressions obtained. In so doing, the kinetics of chemical steps may be ignored.
To determine the surface concentration of EAC and the products of its transformation, it is necessary to formulate additional relationships between these quantities. In the case of labile systems, the material balance equations are used in conjunction with the expressions for the constants of the corresponding equilibria. The data obtained make it possible to construct the Tafel dependences, normalized by the EAC surface concentration.

The results obtained in the analysis of experimental data obtained for a number of real systems indicate the adequacy of the suggested model.

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Appendix

Let us consider a relatively simple system—the solution containing complexes $\text{ML}^{n+}_j$ ($j \in [0, N]$) and uncharged ligand $L$ that is capable of forming coordination bonds with $M^{n+}$ ions. Chemical reactions occurring in the system can be presented in general form as follows:

$$\text{ML}^{n+}_{j-1} + L \rightleftharpoons \text{ML}^{n+}_j.$$  \hspace{1cm} (A1)

The rates of forward and backward chemical reactions, given by (A1) ($w_j$ and $w_{-j}$ respectively), can be expressed by the relationships:

$$w_j = k_j [\text{M}^{n+}_{j-1}][L],$$ \hspace{1cm} (A2)

$$w_{-j} = k_{-j} [\text{M}^{n+}_j],$$ \hspace{1cm} (A3)

Where $k_j$ and $k_{-j}$ are the respective rate constants. When the solution contains an excess of supporting electrolyte, it is possible to neglect the effects of ion migration. Then, to describe the mass transport in the diffusion layer, the second Fick’s law supplemented by kinetic terms should be written for all mobile species. When expressing the kinetic terms, it is necessary to take into account that each $\text{ML}^{n+}_j$ complex takes part in two chemical steps. When one of them is given by Eq. (A1), the next step, $\text{ML}^{n+}_j + L \rightleftharpoons \text{ML}^{n+}_{j+1}$, is characterized by rates $w_{j+1}$ and $w_{-(j+1)}$. Then, the differential equations describing a linear diffusion of metal-containing species take the following form:

$$\frac{\partial [\text{ML}^{n+}_j]}{\partial t} = D \frac{\partial^2 [\text{ML}^{n+}_j]}{\partial x^2} + w_j - w_{-j} - \left(w_{j+1} - w_{-(j+1)}\right).$$ \hspace{1cm} (A4)

In contrast to metal complexes, the ligand $L$ takes part in all chemical steps and the differential equation for this species assumes the form:

$$\frac{\partial [L]}{\partial t} = D \frac{\partial^2 [L]}{\partial x^2} + \sum_{j=1}^{N} \left(w_{-j} - w_j\right).$$ \hspace{1cm} (A5)

Note: in the above records, the kinetic terms ($w_j$ or $w_{-j}$) are equal to zero at $j < 1$ and $j > N$. Besides, the same effective diffusion coefficient $D$ is attributed to all species. The validity of this approach is discussed below.

The set of $(N+2)$ differential equations has a characteristic feature. A linear combination of equations written for metal-containing species yields a simple relationship

$$\frac{\partial c_M}{\partial t} = D \frac{\partial^2 c_M}{\partial x^2},$$ \hspace{1cm} (A6)

Containing no kinetic terms. Here $c_M$ is a total metal concentration, which is determined as a sum of concentrations of metal-containing species, i.e.

$$c_M = \sum_{j=0}^{N} [\text{ML}^{n+}_j].$$ \hspace{1cm} (A7)

A similar expression

$$\frac{\partial c_L}{\partial t} = D \frac{\partial^2 c_L}{\partial x^2}$$ \hspace{1cm} (A8)

is also obtained for a total ligand concentration

$$c_L = \sum_{j=0}^{N} j[\text{ML}^{n+}_j] + [L].$$ \hspace{1cm} (A9)

Nothing essentially does not change when the system is more complicated, for example, if it contains protonated ligands $\text{LH}^{m+}_j$. Then their concentrations should be included in the respective material balance equations and the system of differential equations should be supplemented by corresponding relations, considering the kinetics of equilibria involving $\text{LH}^{m+}_j$, $\text{H}^+$ and $\text{OH}^-$ species [1]. Similar extra procedures should be made if protonated complexes...
are formed in the system. In all these cases, we again obtain the relations (A6) and (A8). However, it should particularly be noted that an account for new chemical interactions with hydronium ions results in the emergence of a new relation:

\[ \frac{\partial c_{H}}{\partial t} = D \frac{\partial^{2} c_{H}}{\partial \alpha^{2}} \]  

(A10)

Where \( c_{H} \) is the total concentration of proton donors and acceptors. In contrast to the values of the \( c_{M} \) or \( c_{L} \), which have clear physical meaning, the quantity \( c_{H} \) is in a sense artificial. This characteristic follows from the set of the differential equations and represents the sum of concentrations involving: hydrated protons and their donors (with positive sign) and proton acceptors (with negative sign). The OH\(^{-}\) ion is a typical proton acceptor since it is capable of binding hydronium ions.