

Full Paper

Multistep Surface Electrode Mechanism Coupled with Preceding Chemical Reaction-Theoretical Analysis in Square-Wave Voltammetry

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Abstract- In this theoretical work, we present for the first time voltammetric results of a surface multistep electron transfer mechanism that is associated with a preceding chemical reaction that is linked to the first electron transfer step. The mathematical model of this so-called “surface CEE mechanism” is solved under conditions of square-wave voltammetry. We present relevant set of results portraying the influence of kinetics and thermodynamics of chemical step to the features of simulated voltammograms. In respect to the potential difference at which both electrode processes occur, we consider two different situations. In the first scenario, both peaks are separated for at least [150 mV], while in the second case both peaks occur at same potential. Under conditions when both peaks are separated for at least [150 mV], the first process can be described with the voltammetric features of a surface CE mechanism, while the second peak gets attributes of a simple surface electrode reaction. When both peaks take place at same potential, we elaborate an elegant methodology to achieve separation of both overlapped peaks. This can be done by modifying the concentration of the substrate “Y” in electrochemical cell that is involved in the preceding chemical reaction. The results of this work can be of big assistance to experimentalists working in the field of voltammetry of metal complexes and drug-drug interactions.

Keywords- Two-step surface reactions; Protein-film voltammetry; CEE mechanism; Kinetics of homogenous chemical reaction; Equilibrium constant

1. INTRODUCTION

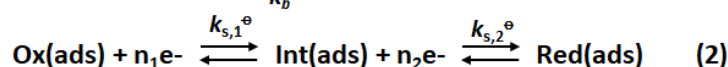
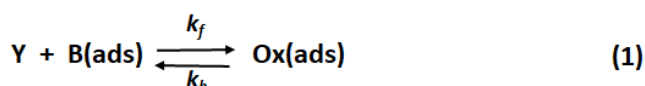
The voltammetric determination of kinetic and thermodynamic parameters that are related to interaction between biomolecules, particularly when these are immobilized on a given substrate, attracts considerable interest in the last 30 years [1-7]. Electrochemical systems in which at least one participant is adsorbed at the working electrode surface are commonly defined as “surface redox reactions” [1, 3, 4, 8]. If voltammetry of lipophilic redox enzymes is considered, then the voltammetric technique that allows studying the redox chemistry of such systems is named “protein-film voltammetry” [5, 8-10]. Relevant experimental examples belonging to surface redox systems are found in antibody-antigen recognition [6], drug-drug interactions [11], recognition of single-stranded DNA [6, 7] and in interactions of many physiological molecules [1, 3, 6]. To get a voltammetric access to parameters that are relevant to thermodynamics and kinetics of surface-active redox substances, we have to monitor the changes in surface concentration of defined substrates analyzed under conditions of applied potential. However, before we consider a suitable voltammetric procedure for such an analysis, we must determine the exact nature of electrochemical mechanism going on in electrochemical cell. The last is of utmost importance in order to apply adequate method for getting access to kinetics and thermodynamics related to defined system of interest. In the last 30 years, the square-wave voltammetry (SWV) is recognized as a powerful pulse voltammetric technique that enables studying the redox mechanisms of many surface redox reactions, and enzymatic reactions as well [1, 3, 5, 8-10]. In addition, it provides elegant means to access relevant kinetics and thermodynamics parameters of important surface redox systems [1-8]. As many surface-active biomolecules exhibit electrochemical activity in a multielectron multi-step fashion [4-7], it was very important to develop theoretical models of such systems under conditions of SWV. In [12-18] we reported on important theoretical models of surface multistep mechanisms associated with follow-up and regenerative chemical reactions under conditions of SWV. In this work, we report for the first time a voltammetric theory of multielectron two-step surface redox mechanism that is associated with preceding chemical reaction under conditions of SWV. The abbreviation of this particular electrode mechanism is a “surface CEE mechanism”. Importance of this particular mechanism is seen in the redox chemistry of many metal-ligand complexes [19] and complex biomolecules as described in [3-7].

2. EXPERIMENTAL

2.1. Mathematical model

Surface electrode mechanism comprising two successive electron transfers, associated with a preceding chemical reaction that is linked to the first electrode step is elaborated theoretically under conditions of square-wave voltammetry. The electrochemical abbreviation

of this complex system is a “surface CEE” mechanism. With term “E” we define a given electron transfer step, while the term “C” stands to describe to a reversible chemical reaction. We assume that all electrochemically active species are strongly and uniformly immobilized to the surface of the working electrode. In the first step (1), we assume occurrence of a reversible chemical reaction between electrochemically inactive species Y and B(ads) that generates the initial electroactive species Ox(ads). The electrochemical transformation of Ox(ads) to Red(ads) takes place in two consecutive one-electron steps (2). Schematically, this mechanism can be described as follows:



Only the species “B(ads)” and “Y” are present in the electrochemical cell before potential is applied. We assume that all immobilized species are uniformly adsorbed (“ads”) at the surface of the working electrode and there are no any lateral interactions between the adsorbed species. “Int(ads)” stands for an electroactive substrate that is formed electrochemically as an intermediate in the first electrode reduction step. With “Red(ads)” we assign the final redox-active species, which is generated electrochemically during the second electrode transfer step from Int(ads). “Y” is a symbol of the substrate that should be present in large excess in the electrochemical system, and it shows no electrochemical activity in the potential region explored in the voltammetric experiment. We assume that “Y” reacts selectively and in chemically reversible manner with B(ads) species, while generating the initial electro-active species Ox(ads). Mathematically, the elaborated surface CEE mechanism in this work can be represented with following equations:

$$t = 0; \Gamma(\text{B}) = \Gamma^*(\text{B}); \Gamma(\text{Ox}) = K_{\text{eq}} \Gamma^*(\text{B}); \Gamma(\text{Red}) = 0 \quad (\text{a})$$

$$t > 0; \Gamma(\text{B}) + \Gamma(\text{Ox}) + \Gamma(\text{Int}) + \Gamma(\text{Red}) = \Gamma^*(\text{B}); K_{\text{eq}} = k_f/k_b; \quad (\text{b})$$

For $t > 0$, the differential equations (c-f) link the changes of surface concentrations with the faradaic current and the kinetics and thermodynamics of preceding chemical step:

$$d\Gamma(\text{B})/dt = k_b\Gamma(\text{Ox}) - k_f\Gamma(\text{B}) \quad (\text{c})$$

$$d\Gamma(\text{Ox})/dt = -I_1/(n_1FS) - k_b\Gamma(\text{Ox}) + k_f\Gamma(\text{B}) \quad (\text{d})$$

$$d\Gamma(\text{Int})/dt = I_1/(n_1FS) - I_2/(n_2FS) \quad (\text{e})$$

$$d\Gamma(\text{Red})/dt = I_2/(n_2FS) \quad (\text{f})$$

The mutual dependence between the electric current, electrical potentials, the surface concentrations of electrochemically active species, and the kinetic parameters relevant to both electron transfer steps is achieved via the Butler-Volmer equations having following forms:

$$(I_1/n_1FS) = k_{s,1}^{\ominus} \exp(-\alpha\Phi_1) [\Gamma(\text{Ox}) - \exp(\Phi_1) \Gamma(\text{Int})] \quad (\text{g})$$

$$(I_2/n_2FS) = k_{s,2}^{\ominus} \exp(-\alpha\Phi_2) [\Gamma(\text{Int}) - \exp(\Phi_2) \Gamma(\text{Red})] \quad (\text{h})$$

In the Supplementary of this work we provide the readers entire MATHCAD working sheet containing all recurrent formulas and parameters needed for simulation of the theoretical voltammograms of the surface CEE mechanism. In all simulation voltammetric patterns, we defined the reduction currents to have positive value (blue color), and negative sign is ascribed to oxidation currents (represented with red color). Net SWV currents are assigned with black color at all simulated patterns. It is important to note that we defined all potentials against the standard redox potential of the first electrode process (defined as “Peak I” in this work). In all calculations, we set the starting potential to defined positive value, and the scan is directed towards negative final potentials.

2.2. Definition of the parameters that exhibit effect to the relevant characteristics of calculated square-wave voltammograms

With the equation $\Psi = \Psi_I + \Psi_{II}$ is defined the net dimensionless current of theoretical SW voltammograms that is as a sum of the particular currents related to the first and the second electrode step, respectively. These particular currents are defined as $\Psi_I = I_1 / [(n_1FSf\Gamma^*)]$ and $\Psi_{II} = I_2 / [(n_2FSf\Gamma^*)]$. Symbol “I” stands for the faradaic electric current of first (I_1) and second (I_2) electrode process, while n_1 and n_2 refer to number of electrons involved in each electrode step. Symbol S stands for the active area of the working electrode, while with f we define the frequency of SW pulses, $f = 1/(2t_p)$. In last equation, t_p is the time of duration of a single potential pulse in SWV. With Γ^* we assign the total surface concentration, which is actually equal to the initial surface concentration of adsorbed B(ads) species. Φ stays for the dimensionless potentials that are defined as $\Phi_1 = nF(E - E_1^{\ominus}/RT)$ and $\Phi_2 = nF(E - E_2^{\ominus}/RT)$, for the first and the second electrode process, respectively. In last two equations, E_1^{\ominus} and E_2^{\ominus} are symbols for the standard potentials of the first and the second electron transfer step, respectively. With “ α ” we assign the coefficient of electron transfer that was set to $\alpha = 0.5$ for both EE steps. T is a symbol of the thermodynamic temperature (T = 298 K in all simulations), R is the universal gas constant, and F is the Faraday constant. The features of theoretical SW voltammograms are also function of several dimensionless parameters. The dimensionless kinetic parameters $K_I = k_{s,1}^{\ominus}/f$ and $K_{II} = k_{s,2}^{\ominus}/f$ portray the effect of $k_{s,1}^{\ominus}$ and $k_{s,2}^{\ominus}$ (i.e. the standard kinetic constants of both EE processes) to the duration of the SW potential pulses. In addition, the major attributes of theoretical SW voltammograms are

affected by a dimensionless chemical parameter K_{chemical} , defined as $K_{\text{chemical}} = \varepsilon/f$. In last equation, $\varepsilon = (k_f + k_b)$ is the cumulative chemical parameter defined as a sum of the first order rate constant k_f and k_b of the forward and backward chemical reactions, respectively. K_{chemical} reflects the overall rate of the chemical step relative to the time-frame of current measurement in SWV. Since we assume to have substrate “Y” that should be present in very large excess in the electrochemical cell, then it holds true that chemical parameter K_{chemical} is of pseudo-first order. It depends on the concentration of substrate “Y” $-c(\text{Y})$ via following relationship: $\varepsilon = [k_f^{\circ}c(\text{Y}) + k_b]$. In the last equation, k_f° is the real rate constant of forward chemical step. In addition, the features of calculated SW voltammograms are function of the equilibrium constant K_{eq} that is defined as $K_{\text{eq}} = k_f/k_b$. The magnitude of chemical equilibrium constant determines the quantity of Ox(ads) that is available to undergo electrode transformation. In all simulations, we set the parameters of the applied bias to following constant values: frequency $f = 10$ Hz, amplitude of the SW pulses $E_{\text{sw}} = 50$ mV, and potential increment $dE = 10$ mV. Additional material for the algorithms used in this work is provided in the *Supplementary* of this work and in our works [1, 13]. For all simulations, we used commercially available software MATHCAD 14.

3. RESULTS AND DISCUSSION

The voltammetric complexity of multistep surface mechanisms can be initially simplified if we assume that both peaks are separated for at least $|150 \text{ mV}|$. Under such circumstances, the voltammetric patterns of a surface CEE mechanism are equivalent to the SW voltammograms simulated for a surface EE mechanism [20], when the equilibrium constant $K_{\text{eq}} > 10$, independent on the magnitude of dimensionless chemical parameter K_{chemical} [1, 21]. In this scenario, the chemical equilibrium is shifted significantly to the right, i.e. towards production of initial electrochemically active species Ox(ads). Under such conditions, one can explore the methods reported in [1, 20] to get access to the kinetic and thermodynamic parameters relevant to both electron transfer steps. When $K_{\text{eq}} \leq 1$, then we witness significant effects of equilibrium constant and dimensionless chemical parameter to the features of calculated SW voltammograms. In Figure 1 we present several calculated SW voltammograms of a surface CEE mechanism portraying the effect of the rate of preceding chemical step. SW voltammograms are simulated for $K_{\text{I}} = K_{\text{II}} = 1.6$, and for magnitude of $K_{\text{eq}} = 0.5$. As expected, magnitude of K_{chemical} shows effect to the features of “Peak I” only, which gets shapes as typical of a surface CE mechanism [1, 21]. For $K_{\text{chemical}} \geq 10$, there is no further influence of chemical reaction rate to the features of simulated SWV patterns, and entire system turns to a two-step surface EE mechanism [20].

Presented in Figure 2a-b is the effect of K_{chemical} to the magnitude of net SWV peak currents and net SWV peak potentials of “Peak I”. Curves are calculated for $K_{\text{I}} = 0.2$, and for three different magnitudes of K_{eq} .

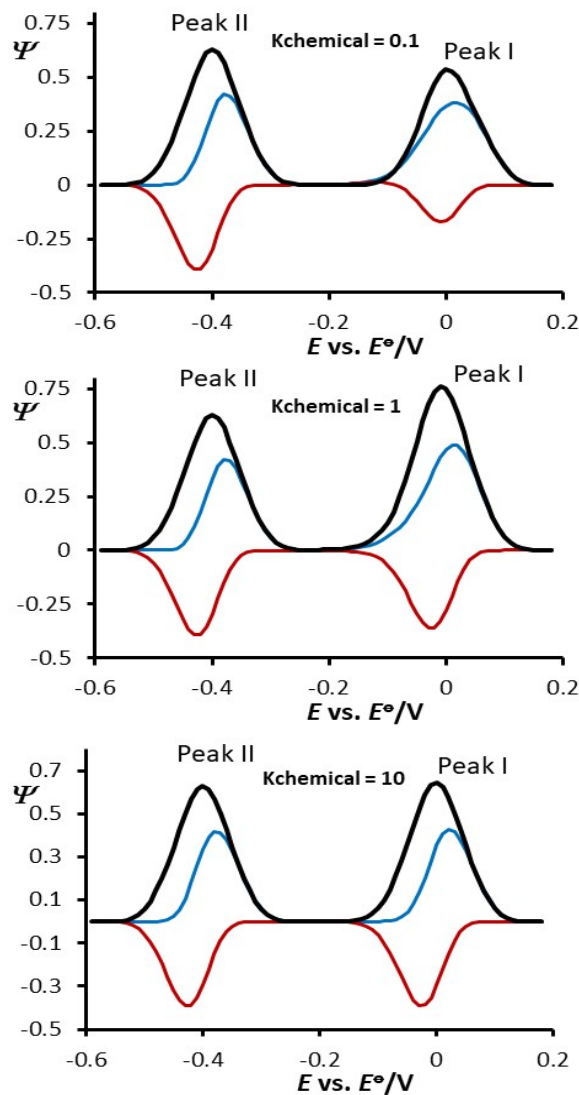


Fig. 1. Influence of the dimensionless chemical parameter K_{chemical} to the features of SW voltammograms simulated for $K_{\text{I}} = K_{\text{II}} = 1.60$, and equilibrium constant of $K_{\text{eq}} = 0.5$. The potential separation between both peaks is set to $|400 \text{ mV}|$. The values of other parameters used in simulations were: temperature $T = 298 \text{ K}$; number of electrons $n_1e^- = n_2e^- = 1$; electron transfer coefficient $\alpha = 0.5$, potential step $dE = 10 \text{ mV}$, frequency $f = 10 \text{ Hz}$. The values of K_{chemical} are given in the charts.

A sigmoidal dependence exists between the net SWV peak current of “Peak I” - $\Psi_{\text{net,p1}}$ and the $\log(K_{\text{chemical}})$, with a linear “kinetic zone” existing roughly in the region $-1 < \log(K_{\text{chemical}}) < 1$ (Figure 2a). Indeed, the region in which we “detect” a linear kinetic effect of $\log(K_{\text{chemical}})$ to $\Psi_{\text{net,p1}}$ depends on magnitude K_{eq} , and it is larger at smaller values of K_{eq} . The shift of the net-SW potential of “Peak I” as a function of logarithm of chemical kinetic parameter also follows a sigmoidal dependence (Figure 2b).

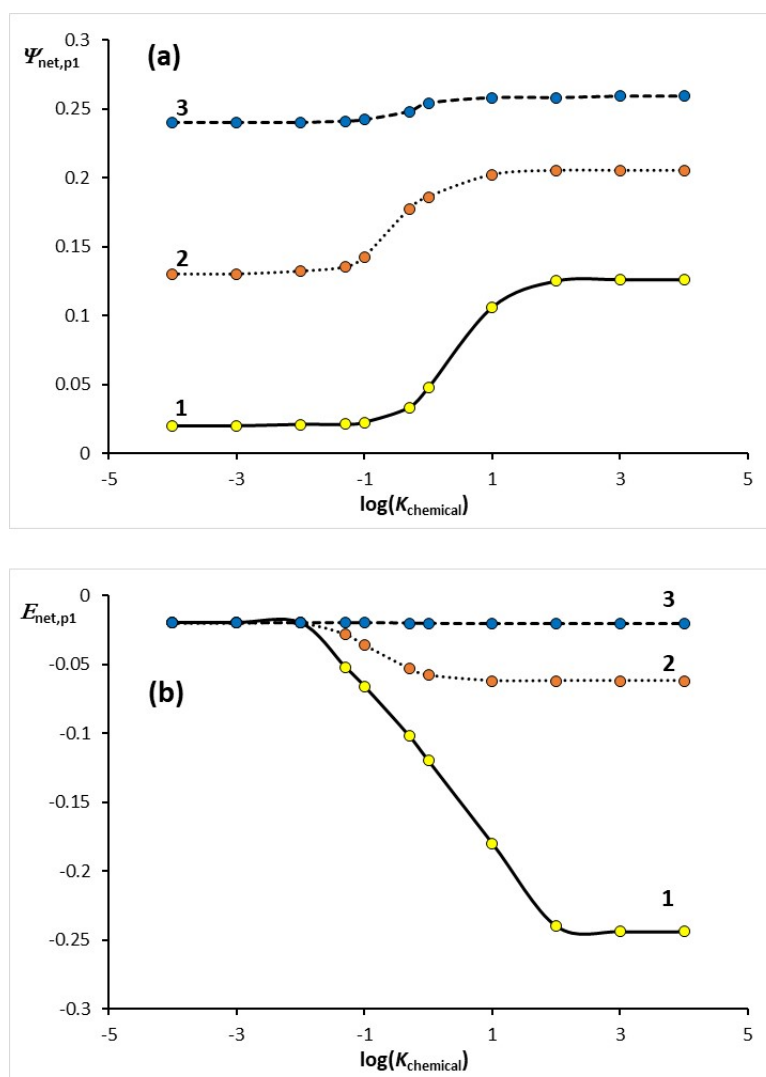


Fig. 2. Effect of dimensionless chemical rate parameter K_{chemical} to the net peak currents $\Psi_{\text{net,p1}}$ (a) and the net peak-potentials $E_{\text{net,p1}}$ (b) of SW voltammograms of “Peak I”. Voltammograms are simulated at $K_{\text{I}} = 0.2$ and $K_{\text{eq}} = 0.1$ (1); 1 (2) and 10 (3). The other parameters used in simulation were same as those in Figure 1.

For $K_{\text{eq}} \leq 1$, we observe linear parts of the dependence $E_{\text{net,p1}}$ vs. $\log(K_{\text{chemical}})$, roughly in the regions $-1 < \log(K_{\text{chemical}}) < 1.5$. The slope of the linear parts of the curves $E_{\text{net,p1}}$ vs. $\log(K_{\text{chemical}})$ is a function of K_{eq} and it is defined as $2.303[RT/F]\log[K_{\text{eq}}/(1+K_{\text{eq}})]$. Therefore, the magnitude of the slope of $E_{\text{net,p1}}$ vs. $\log(K_{\text{chemical}})$ can be explored for the determination of the value of K_{eq} . The dependences between $\Psi_{\text{net,p1}}$ and $E_{\text{net,p1}}$ as a function of K_{eq} are presented in Figure 3a-b. Curves are simulated for $K_{\text{I}} = K_{\text{II}} = 0.2$, and for three different values of K_{chemical} . All curves of the dependence of $\Psi_{\text{net,p1}}$ vs. $\log(K_{\text{eq}})$ have a sigmoidal shape with identical slopes of the linear parts (Figure 3a). For magnitudes of $\log(K_{\text{eq}}) > 2$, we detect a constant value of $\Psi_{\text{net,p1}}$, independent on K_{eq} .

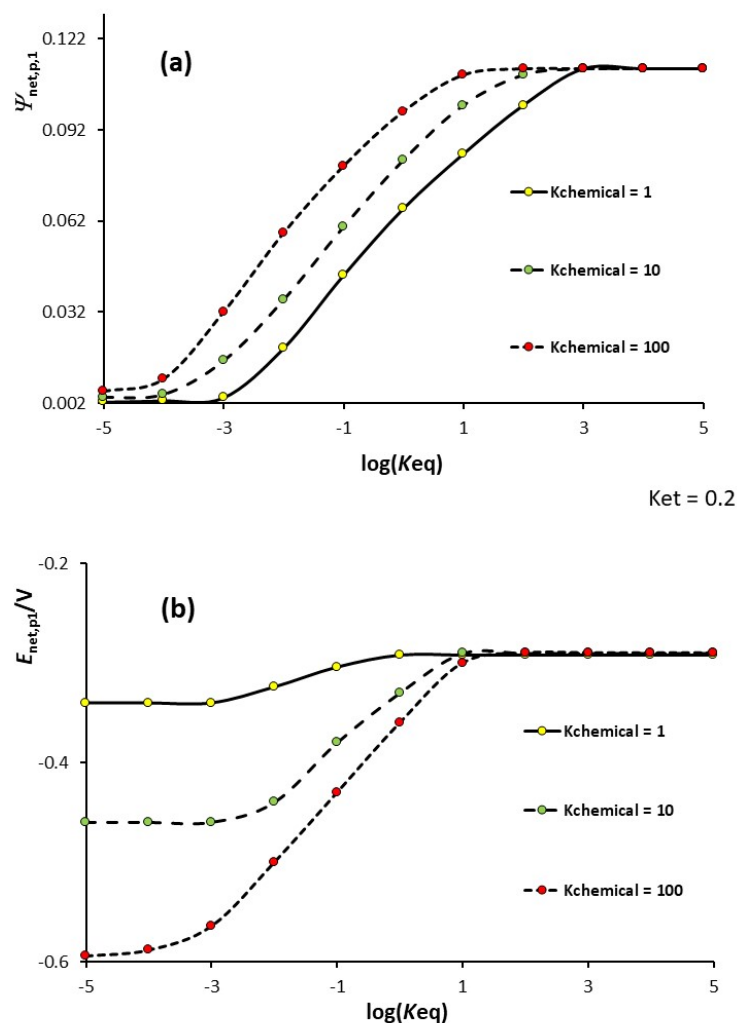


Fig. 3. Influence of the equilibrium constant of preceding chemical reaction K_{eq} to the net peak currents $\Psi_{net,p1}$ (a) and the net peak-potentials $E_{net,p1}$ (b) of SW voltammograms of “Peak I”. Voltammograms are simulated at $K_i = 0.2$ and $K_{chemical} = 1, 10$ and 100 . The other parameters used in simulation were same as those in Figure 1.

This scenario corresponds to significant shift of the chemical equilibrium towards Ox(ads) species, which turns the features of first peak (“Peak I”) from CE to a simple E mechanism [21]. The dependence of the magnitude of net SWV peak potential of “Peak I”- $E_{net,p1}$ as a function of $\log(K_{eq})$ also follows a sort of sigmoidal function (Figure 3b). In general, an increase of K_{eq} generates a shift of $E_{net,p1}$ towards more positive values. For $\log(K_{eq}) > 1$, the net SWV peak potential of “Peak I” is independent on K_{eq} . The slopes of the linear segments of the dependences $E_{net,p1}$ vs. $\log(K_{eq})$ are function of $K_{chemical}$, and they get steeper at higher values of chemical rate parameter. A typical feature of all surface mechanisms in pulse voltammetric techniques is the so-called “quasireversible maximum”. A sort of parabolic dependence of net SWV peak currents and the logarithm of electrode kinetic parameter exists, as a result of the synchronization of the kinetics of electron transfer to the time-frame

of current measurements in SWV [1, 22]. This phenomenon can be used for the estimation of standard rate constant of electron transfer step in a very simple manner [1]. As reported in [21, 23], for a surface CE mechanism, the position of the “quasireversible maximum” is not affected by the kinetics of chemical reaction that precedes the electron transfer step. Therefore, this feature can be used for accurate assessment of both $k_{s,1}^{\ominus}$ and $k_{s,2}^{\ominus}$ independently for both SWV peaks of a surface CEE mechanism.

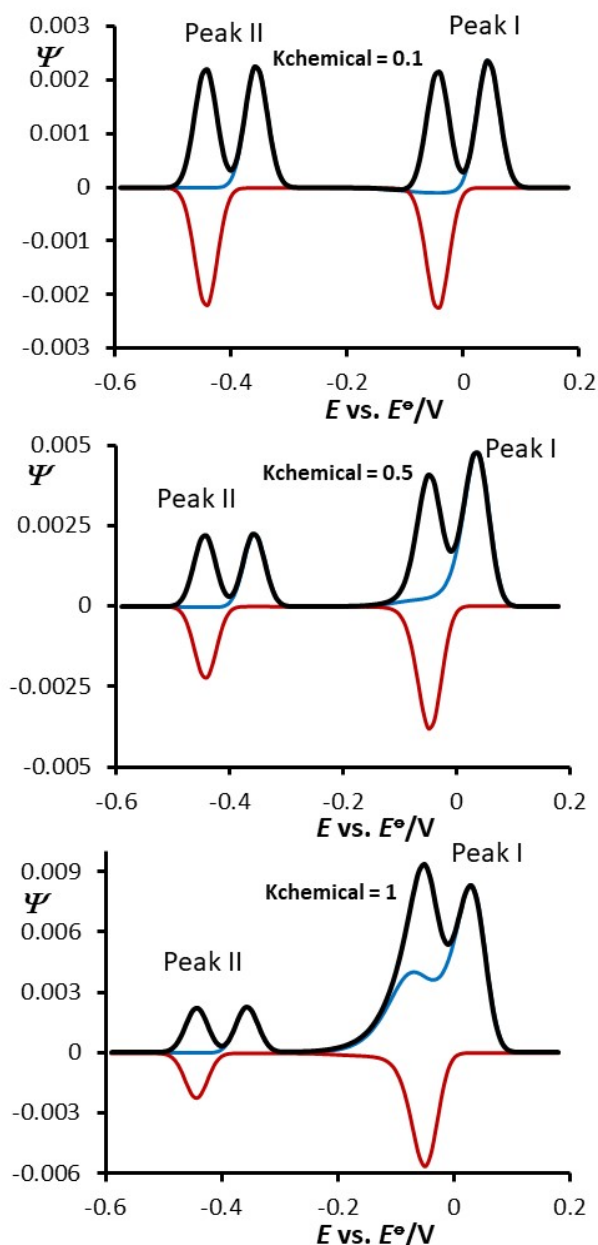


Fig. 4. Influence of the dimensionless chemical parameter K_{chemical} to the features of SW voltammograms in situation of fast electron transfer at both electrode steps. Voltammometric patterns are simulated for $K_{\text{I}} = K_{\text{II}} = 10$, and equilibrium constant of $K_{\text{eq}} = 0.1$. The potential separation between both peaks is set to $|400 \text{ mV}|$. The values of K_{chemical} are given in the charts. Other simulation conditions are same as those in Figure 1.

When the kinetics of electron transfer is very fast (see Figure 4a), then one might witness a phenomenon of so-called “split net SWV peaks” [1, 22, 24]. In such scenario, a very short time-frame at SW potential pulses is needed for the electrochemical conversion of redox active form Ox(ads) to redox form Red(ads). However, an electrochemical conversion of Ox(ads) to Red(ads) also happens in the so-called “dead time” (i.e. the time segment in which faradaic current is not measured) of SW pulses [1, 22]. Therefore, in the small time-frame where electrical current is measured at the defined SW pulses, only very small amounts of Ox(ads) (or Red) will be available for electrochemical transformation. Such a phenomenon will be portrayed in small faradaic currents measured in SWV of surface electrochemical systems that features very fast rate of electrode steps [24]. As the rate of electrode reaction gets faster, then the reduction process shifts towards more positive potentials, while the oxidation process moves to more negative potentials. This will result in a splitting of the net SWV peak in two symmetric peaks, whose characteristics can be used to determine the magnitude of k_s^\ominus as elaborated in [1, 24]. Shown in figure 4b-d is the effect of chemical rate parameter K_{chemical} to the features of calculated patterns in scenario of “split SW peaks” of a surface CEE mechanism. Voltammograms are simulated for $K_{\text{I}} = K_{\text{II}} = 10$, and for $K_{\text{eq}} = 0.1$. As the second process at more negative potentials (Peak II) remains unaffected by K_{chemical} , the features of first SWV process (Peak I) get significantly affected by the value of K_{chemical} . An increase of K_{chemical} from 0.1 to 1 is followed by a rise of all current components of “Peak I”. In same direction, we witness a diminishment of the potential separation between the split patterns of net “Peak I”. For $K_{\text{chemical}} > 2$, the splitting phenomenon vanishes and only a single-featured “peak I” exists at more positive potentials, with intensity that is much higher than that of “Peak II” (not shown). The phenomena portrayed in figure 4 can be used as a qualitative criterion to recognize the surface CEE mechanism, if both SWV peaks are separated for at least |150 mV|.

When both SWV peaks, which represent both electrode reactions of a defined multistep electron transfer process, are separated for at least |150 mV|, then it becomes quite easy to achieve independent estimation of all relevant parameters to both electrode steps [12, 23]. However, when both EE steps occur at a same potential, then the CEE electrode mechanism will be portrayed in a single SW voltammogram that “hides” all processes in it (figure 5a). In such scenario, it is a challenging task to determine whether the obtained single SWV peak is a consequence of a simple two-electron transfer that occurs in one step, or it is due to occurrence of two successive electron transfers. In our recent paper [25], we have shown that in a surface EEC mechanism studied with SWV, there is a simple way to determine occurrence of a two-step successive surface electrode mechanism, when both EE steps take place at equal potential. It has been shown that at an increased rate of the chemical reaction that follows the electron transfer step, expressed via K_{chemical} , can lead to displacement of the second electrode step of a surface EEC_{rev} mechanism to positive potentials. Similar

algorithm can also be applied to recognize a surface CEE mechanism, when both EE processes are defined to take place at a very same potential. In this case, an increased rate of the preceding chemical reaction displaces the first electron transfer process towards more negative potentials (Figure 5b-f). For $K_{\text{eq}} = 0.1$, and $K_{\text{I}} = K_{\text{II}} = 1.6$, a simple separation of both electron transfers (that happen at very same potential) can be achieved for $K_{\text{chemical}} \geq 5$ (see Figure 5e-f). This feature can be explored as additional diagnostic criterion for recognizing the surface CEE mechanism. Therefore, one can use this characteristic for separation of both EE processes of a CEE mechanism, and to analyze them independently, if both happen at the same potential.

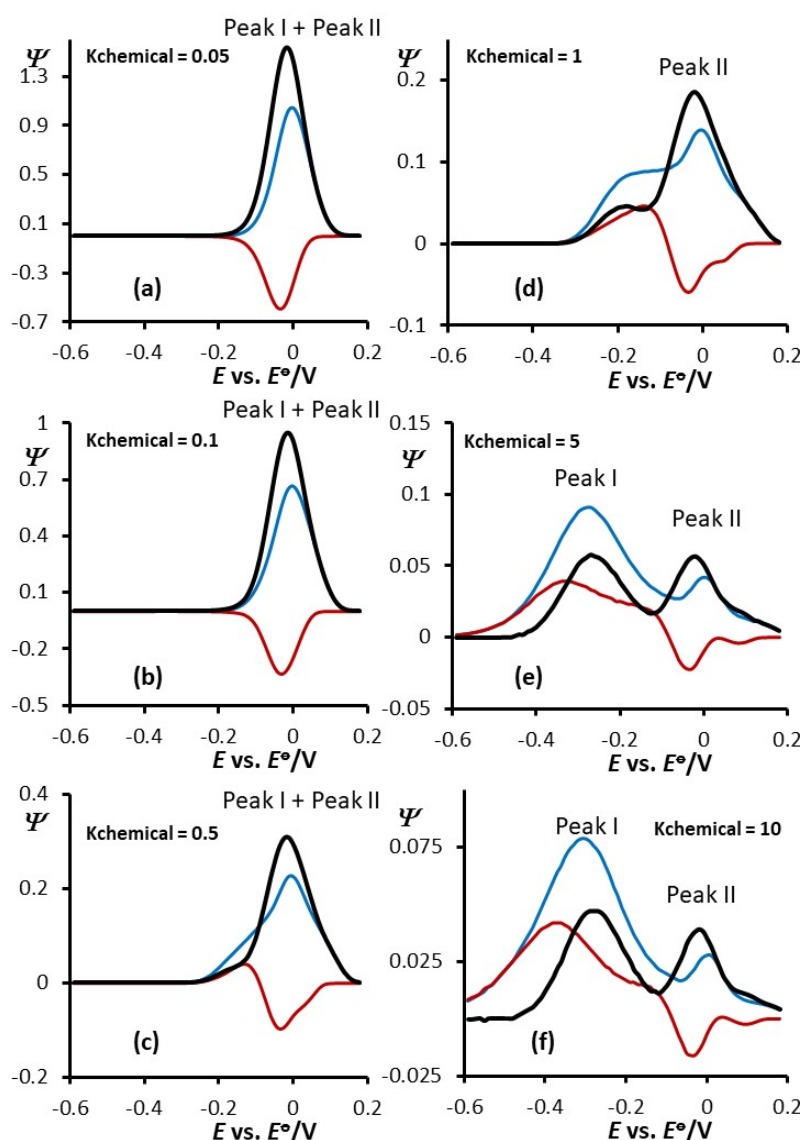


Fig. 5. Influence of the magnitude of dimensionless chemical parameter K_{chemical} to the features of SW voltammograms in situation when both electrode processes take place at same potential. SW voltammetric patterns are simulated for $K_{\text{eq}} = 0.1$, and $K_{\text{I}} = K_{\text{II}} = 1.60$. The values of K_{chemical} are given in the charts. Other conditions are same as in Figure 1.

4. CONCLUSION

The results presented in this work enable a comprehensive study of the voltammetric responses of a surface CEE mechanism as a function of the kinetic rate constant and equilibrium constant of preceding chemical step. From the voltammograms simulated in this work, we can observe that the kinetic and thermodynamic parameters associated to chemical step can exhibit rather complex behavior to simulated SW voltammograms. Even more complex situation is met if both electron transfers take place at same electrode potential. When both electrode steps of a surface CEE mechanism occur at potentials that are at least $|150 \text{ mV}|$ separated from each other, then one can explore independent methodologies to determine the relevant rate constant and equilibrium constant of both electrode steps. For both EE steps, one can explore the phenomenon “quasireversible maximum” [1] and “split net SWV peaks” [24] to get access to the magnitudes of $k_{s,1}^{\circ}$ and $k_{s,2}^{\circ}$. Under such circumstances, the methods reported in [1, 20, 23] can be used to access the kinetics and thermodynamic parameters related to the preceding chemical reaction. For these estimations, one needs the value of electron transfer coefficient that can be evaluated by using the method reported in [26]. If both EE processes occur at same potential, a crucial point is to recognize whether the electrode step is a single electron transfer, or it is a consequence of two one-step reactions. As we elaborated in this work, at given magnitudes of K_{eq} , K_{I} and K_{II} , all relevant characteristics of simulated SWV patterns of a surface CEE mechanism are function of the dimensionless chemical parameter K_{chemical} . For $K_{\text{eq}} \leq 1$, an increase of K_{chemical} leads to a shift of the position of first electrode step towards more negative potentials. As presented in figure 5e-f, at given critical value of K_{chemical} , we can achieve separation of both EE processes defined to take place at same potential. In the section “Mathematical model” of this work, it is explicitly shown that the dimensionless chemical parameter K_{chemical} depends on concentration of “Y”- $c(\text{Y})$ and the frequency f . It is important to mention that f affects in the same time the kinetics of chemical reactions, and the rate of the electrochemical reactions (via K_{I} and K_{II}) of the redox active compounds. Therefore, the frequency analysis in real experiments of such systems will produce a rather complex interplay to all relevant kinetic parameters. So, a more relevant protocol to achieve experimentally the separation of both peaks of SW voltammograms portrayed in figure 5 is to make modification of the molar concentration of substrate “Y”. In such experiments, we should keep at constant values the SW frequency, and also the SW amplitude and the potential step [27]. From the experimental voltammograms obtained by modifying the molar concentration of substrate “Y” only, we can distinguish the elaborated surface CEE from other surface mechanisms (two-step) that are associated with chemical reactions. This is achievable, because all other multistep surface mechanisms coupled with chemical reactions (i.e., ECE, EEC and EEC’ mechanisms) [12-18, 25] show very specific SW voltammetric characteristics, regardless if both EE processes occur at a very same or at quite different potentials.

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