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Kinetics of the Hydrogen Electrode Reaction

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It is well recognized that the standard Butler–Volmer equation is lacking in an adequate description of the kinetics of the hydrogen electrode reaction over the complete range of potentials for the alkaline as well as the acid electrolytes. Further, it is unable to explain the asymmetry in current vs potential observed in the hydrogen evolution reaction (HER) vs the hydrogen oxidation reaction (HOR). In fact, even kinetic descriptions via two-step mechanisms (Volmer–Tafel, or Heyrovsky–Tafel) are individually applicable only in limited potential ranges. We present an approach that provides explicit rate expressions involving kinetics of all the three steps (Tafel–Volmer–Heyrovsky) simultaneously, as well as more limiting rate expressions based on two-step pathways. The analysis is based on our recently developed graph-theoretic approach that provides accurate rate laws by exploiting the electrical analogy of the reaction network. The accuracy of the resulting rate expressions, as well as their asymmetric potential dependence, for both HOR and HER is illustrated here based on step kinetics provided in the literature for Pt catalyst in 0.5 M NaOH solution.

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Due to their practical significance, the hydrogen oxidation reaction (HOR) and its reverse, the hydrogen evolution reaction (HER), are by far the most extensively investigated of electrocatalytic reactions. Further, despite being among the simplest of such reactions, their mechanism and kinetic understanding is still incomplete. The significance of dual-pathway kinetics has recently been shown for the HOR on Pt electrode. However, no general rate expression exists that can simultaneously account for these alternate pathways in terms of the accepted three-step mechanism, namely, the Tafel, Volmer, and the Heyrovsky steps. Further, no realistic first-principles prediction of step kinetics yet exists for the hydrogen electrode reaction, although there is now great interest in ab initio predictions as well as in their experimental validation.

Were an accurate rate expression for HOR/HER in terms of its three-step kinetics available, it would not only be revealing, allowing fundamental questions to be answered, such as those posed recently by Gasteiger et al., but when available, it could utilize the first-principles predictions of step kinetics to construct a comprehensive picture of this important and intriguing reaction system, including the elucidation of parallel pathways and the dominant steps. A thorough understanding of HOR and HER would also serve as a yardstick for understanding other electrocatalytic reactions.

Following up on our earlier work, we present here a detailed analysis of the hydrogen electrode reaction based on a graph-theoretic approach we recently developed, namely, the reaction route (RR) graph approach. It involves a topological analysis of alternate pathways as walks on the RR graph, coupled with a kinetic analysis based on Kirchhoff’s laws of (i) flux (i.e., mass conservation) at nodes interconnecting reaction steps represented as branches or edges and of (ii) potential (i.e., state property of thermodynamic potentials) applicable to cycles among mechanistic steps in the RR graph. This procedure is completely analogous to electric circuit analysis, thus allowing the direct use of the electrical analogy in kinetic analysis. Although our earlier work was limited to numerical analysis, here we exploit the electrical analogy to provide explicit rate expressions. The results presented below confirm the importance of two dominant two-step pathways on Pt electrode in alkaline media at different electrode potentials. Further, the complete rate expression provided here that involves all the three mechanistic steps, namely, the Tafel, Volmer, and Heyrovsky steps, is needed for an adequate description over the complete range of potentials of interest for HER and HOR.

Our methodology to deduce a steady-state rate expression for the hydrogen electrode reaction is, thus, based on the RR graph and its electrical analog, along with the notion of intermediate reactions (IRs) for the formation of an intermediate species. It further interrelates the two standard approaches of kinetic analysis, namely, the rate-determining step (RDS)/quasi-equilibrium (QE) approach of Langmuir–Hinshelwood–Houben–Watson (LHHW) and the quasi-steady-state (QSS) analysis of Bodenstein. The resulting rate expression accurately portrays the kinetics of HOR/HER over a broad range of potentials, including their asymmetric potential dependence in both alkaline and acidic media, although in the discussion below we simply use the step rate constants reported for Pt in an alkaline electrolyte. The case of acidic electrolyte, of special interest in fuel cells, will be described in a subsequent publication.

Reaction Mechanism, Network, and Step Kinetics

RRs or pathways. — The hydrogen electrode reaction has been investigated over a long period of time due to its technological and fundamental significance. The most common and well-accepted mechanism involves the Tafel, Volmer, Heyrovsky steps, which have adequately explained the overall reaction (OR) kinetics. This three-step mechanism involves only a single reaction intermediate, H S, where S represents an unoccupied catalyst surface site. However, more recently, other intermediates have been proposed. For instance, intermediates such as adsorbed water (H 2O·S) and adsorbed hydroxyl (OH·S) have been shown to exist on Pt surfaces by Völkening et al., Bedurftig et al., and Rossmeisl et al. Thus, Nørskov et al. consider the OH·S and O·S species to calculate the effect of molecular water on adsorption. Additional intermediates, of course, imply additional elementary steps in the mechanism and would alter the site balance. Nonetheless, because our purpose here is to elucidate an approach that adequately describes the kinetics of the hydrogen electrode reaction, we simply adopt the standard Tafel–Volmer–Heyrovsky mechanism shown below to avoid being distracted by the additional complexities of a more detailed mechanism, which are left for future work.

\[
\begin{align*}
\sigma_{\text{H}_2} & : \text{H}_2 + 2\text{OH}^- \rightleftharpoons \text{H}_2\text{O} + \text{H}^- + \text{e}^- \\
\sigma_{\text{H}_2\text{O}} & : \text{H}_2\text{O} \rightleftharpoons \text{H}_2\text{O} + \text{OH}^- + \text{H}^- + \text{e}^- \\
\sigma_{\text{OR}} & : \text{H}_2 + 2\text{OH}^- \rightleftharpoons 2\text{H}_2\text{O} \\
\end{align*}
\]

Equation 1 above describes the HOR in an alkaline electrolyte. In an acidic electrolyte, however, the corresponding mechanism is

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The Volmer step, $s_V$, above describes the electrochemical consumption of the key intermediate, namely, the surface atomic hydrogen, H$_S$, whereas the nonelectrochemical Tafel step, $s_T$, and the electrochemical Heyrovsky step, $s_H$, describe the generation of H-S from molecular H$_2$. As indicated by the stoichiometric numbers $\sigma_{s_V}$ above, when these steps are combined in a manner that eliminates the intermediate H-S, the OR results. Clearly, this can be accomplished in more than one way, which represents the alternate ORs or pathways. The OR and the elementary steps for the HER are simply the reverse of those written above for the HOR.

Further, there are other ORs or linear combination of steps that do not eliminate all of the species, both intermediate and terminal. Such ORs are called inter- or intermediate, so that $\sigma_{s_V}$ or $\sigma_{s_T}$ may vary widely, this does not present a problem. Other linear combinations or ORs that do not eliminate all of the intermediate species and produce the so-called ORs are called intermediate reaction routes (IRRs). For example, $(-1)s_H + (+1)s_T = IR$, that is

\begin{align*}
    &s_V: H - S + H_2O \rightleftharpoons H_2O^+ + S + e^- \\
    &s_T: H_2 + 2S \rightleftharpoons 2H-S \\
    &s_H: H_2 + S + H_2O \rightleftharpoons H_2O^+ + H-S + e^- \\
    \end{align*}

which is an intermediate reaction that describes the formation of an intermediate, namely, H-S from the terminal species water and electron, as well as the vacant sites, S. We will discuss IRRs further later on.

More generally, we define an IRR as a linear combination, $\sum_{i} \sigma_{s_{i}}s_{i} = IR$, in which all the intermediate species except the one of interest $I_k$ (along with a reference intermediate, say surface site, S) are eliminated. $\Sigma_{i} \sigma_{s_{i}}s_{i} = IR$, the IRRs produced by these IRRs may be written generally as, $\Sigma_{i} \sigma_{s_{i}}s_{i} = (-1)\gamma_{s}S + \Sigma_{i} \sigma_{s_{i}}s_{i}T_{i} = 0$, where $\gamma_{s}$ is positive because the intermediate $I_k$ is considered as a product in the IRR.

Further, there are other ORs or linear combination of steps that eliminate all of the species, both intermediate and terminal. Such ORs are called empty reaction routes (ERs) or cycles, as they produce a “zero” OR (i.e., the stoichiometric coefficients of all the species are zero). In fact, because subtracting one FR from the other, e.g., $FR_{VT} - FR_{HT}$, would eliminate all species, it can provide an ER, namely, $(+1)s_V + (-1)s_H + (+1)s_T = 0$, is an ER, $ER_V$:

\begin{align*}
    &s_V: H - S + OH^- \rightleftharpoons H_2O^+ + S + e^- \\
    &s_H: H_2 + S + OH^- \rightleftharpoons H_2O^+ + H-S + e^- \\
    &s_T: H_2 + 2S \rightleftharpoons 2H-S \\
    \end{align*}

In general, thus, $ER_V \sum_{i} \sigma_{s_{i}}s_{i} = 0$ represents an ER.

Finally, let us consider another linear combination of FRs, e.g., $FR_{VT} + FR_{VT} - FR_{HT}$, which simply results in another FR, namely, $FR_{VT} + (+3)s_V + (+2)s_T + (-1)s_H = OR$. In fact, because new ORs may be obtained simply by linearly combining others, one may, in principle, obtain an infinite set, if no further restrictions are placed on their definition.

This is avoided by the concept of “directness” proposed by Milner, i.e., the number of steps involved in an OR must be minimal. In other words, a “direct” FR must not contain any cycles or ERs. Such a direct FR contains no more than $q + 1$ steps, whereas a direct ER contains no more than $q + 2$ steps, selected from among the given mechanism of $p$ steps. Here, $q$ is the number of linearly independent intermediate species, typically less than $n$ less than the number of intermediates due to intermediates conservation, e.g., site conservation. For the three-step ($p = 3$) HOR/HER reaction mechanism considered here, H-S is, in fact, the only independent intermediate, so that $q = 1$ simply.

For the hydrogen electrode reaction, thus, a direct FR must not contain more than any of the two of the three steps, whereas a direct ER must be restricted to less than three steps. As a result, the FR obtained above from the combination of others, i.e., $FR_{VT} + (+3)s_V + (+2)s_T + (-1)s_H = OR$ is not an appropriate FR as it contains three elementary steps. In fact, it has embedded in it a cycle, namely, $ER_V:(+1)s_V + (-1)s_H + (+1)s_T = 0$, which if subtracted from it, results in $FR_{VT}$.

With this restriction on path length, a finite and unique set of FRs and ERs results, as listed in Table I for the above HOR/HER mechanism. As per the Horiuti–Temkin theorem, furthermore, an independent RR set is any set of $\mu = p - q = 3 - 1 = 2$ RRs, which may include both FRs and ERs, so long as they include among them all of the steps in the mechanism. Moreover, the number of linearly independent ERs is given by $n - (q + 1) = 3 - 2 - 1 = 1$ for the reaction mechanism considered. Thus, a set of two linearly independent RRs may be readily determined by identifying one independent ER and one FR by a simple inspection of the HER/HOR mechanism, as done above, thus avoiding the stoichiometric enumeration of FRs used for complex mechanisms.

Let us consider the $FR_{VT}$ and $ER_V$, mentioned above as the independent set of RRs for the considered HOR/HER mechanism, from which the remaining set of unique FRs can be obtained. Thus, a linear combination of $FR_{VT}$ and $ER_V$ results in $FR_{VT}$ and $FR_{HT}$ (Table I).

These RRs or pathways may, in fact, be simply traced as walks on the RR graph of a mechanism for an OR. The construction of
We start by assembling the ERs, only ER 1 in this case, the construction of the RR graph is straightforward and is illustrated twice. This is a consequence of the fact that the mechanism is walks between the TNs. In fact, every RR in the graph is involved RR/H$_2$O by simply connecting the OR/H$_2$O into a cycle graph. It is further noted that there exists nonunit stoichiometry across which the OR is connected. These RR graphs further follow Kirchhoff’s laws of flow graphs rooted in the species conservation principle along with the state property of thermodynamic functions, which provide the topological constraints on the individual step rates. Of course, Kirchhoff’s laws are central to the analysis of electrical networks. For this reason, RR graphs are completely analogous to electrical networks. In fact, a QSS rate expression of Ohm’s law form, i.e., OR rate = driving force/overall network resistance, can be derived by exploiting the analogy to electrical circuits. This property is used below to obtain a general rate law for the kinetics of HOR/HER.

With the independent set of RRs (i.e., FR$_\text{H}_{\text{OR}}$ and ER$_j$) at hand, the construction of the RR graph is straightforward and is illustrated in Fig. 1. We start by assembling the ERs, only ER$_1$ in this case, into a cycle graph. It is further noted that there exists nonunit stoichiometric ($\sigma_{ij} = +2$) numbers in some of the RRs in the unique set (Table I), i.e., in FR$_\text{VT}$ and in FR$_\text{H}$. This implies that every elementary reaction step $s_j$ as well as the OR must occur twice in the RR graph, which must furthermore be symmetrical.$^{20,21}$ This can be accomplished by fusing two ER$_j$s, as shown in Fig. 1a. Next, the remaining RR of the independent set, namely, FR$_\text{H}_{\text{OR}}$, can be included in the graph by simply connecting the OR (also twice) across TNs, yielding the final RR graph (Fig. 1b). All of the four unique RRs (Table I) for HOR can be traced on the resulting RR graph as walks between the TNs. In fact, every RR in the graph is involved twice. This is a consequence of the fact that the mechanism is nonminimality, i.e., the elementary steps are involved more than once in a FR. Nonetheless, the affinity (or any other thermodynamic potential change across it) and the rate of a step (e.g., $\eta_{\text{H}}$, $v_{\text{H}}$, or $s_j$) remain unchanged regardless of their placement because of the network symmetry. For HER, the FR walks are simply in the opposite direction.

The other characteristic that an appropriate RR graph must satisfy is that the connectivity at the intermediate nodes (INs) and the TNs must be consistent with the QSS condition for the intermediate and terminal species. Thus, INs interconnect mechanistic steps $s_j$ only, with the incidence of steps being consistent with the QSS conditions for intermediates or their linear combination, along with the condition of minimality. The TNs, however, interconnect mechanistic steps $s_p$ to the OR, with the incidence of steps being consistent with QSS equations for terminal species as well as with minimality of incidence.$^{23,24,26}$

Because there is only one linearly independent intermediate here, H·S, its QSS condition is

$$Q_{H·S}: (-2)r_{\text{OR}} + (+1)r_{\text{V}} + (-1)r_{\text{H}} = 0$$

[6]

which is consistent with the connectivity, $\sum_{j}m_{ij}s_j = 0$, where the incidence coefficient $m_{ij} = +1$, if a branch leaves the node $j$, and $m_{ij} = -1$, if a branch is coming into the node $j$, of the only one IN (although present twice) in the RR Graph, that is

IN$_1$: $(-2)s_{\text{OR}} + (+1)s_{\text{V}} + (-1)s_{\text{H}} = 0$

[7]

Similarly, the QSS condition for the terminal species (H$_2$O, OH$^-$, H$^+$, e$^-$) for the HOR in alkaline electrolyte (Eq. 1) are

$$Q_{\text{H}_2\text{O}}: (+2)r_{\text{OR}} + (+1)r_{\text{V}} + (+1)r_{\text{H}_2\text{O}} = 0$$

$$Q_{\text{OH}^-}: (-1)r_{\text{OR}} + (+1)r_{\text{V}} + (+1)r_{\text{OH}^-} = 0$$

$$Q_{\text{H}^+}: (+1)r_{\text{OR}} + (+1)r_{\text{V}} + (+1)r_{\text{H}^+} = 0$$

$$Q_{\text{e}^-}: (-2)r_{\text{OR}} + (+1)r_{\text{V}} + (-1)r_{\text{H}^+} = 0$$

The QSS condition for OH$^-$, e$^-$, and H$_2$O is the same and is represented by the TN$_i$ (represented twice) in the RR graph, namely

$$\text{TN}_i: 2\text{OR} - s_{\text{V}} - s_{\text{H}} = 0$$

[9]

Thus, the resulting RR graph satisfies all of the conditions imposed on the RR graph, i.e., all nodes are balanced in that they satisfy the QSS conditions of one or more surface intermediates (in case of INs) and of one or more terminal species (in case of TNs). Further, all the RRs can be traced as walks or paths on the RR graph. The network includes the commonly considered Volmer–Heyrovsky and Volmer–Tafel pathways, along with the not so common Heyrovsky–Tafel pathway.$^3$ Thus, this is an appropriate RR graph for the hydrogen electrode reaction.

Finally, a curious observation by Gennero de Chialvo and Chialvo$^{12,26}$ that two distinct sets of alternate parameters provide identical HOR/HER kinetics can be explained simply from the topology of the RR graph. Thus, it is clear from the symmetry of the RR graph in Fig. 1b that the Volmer and the Heyrovsky steps can be interchanged without affecting the properties of the circuit. As a result, interchanging the kinetic parameters of the Volmer and the Heyrovsky steps does not alter the current density vs overpotential predictions, as found by Gennero de Chialvo and Chialvo$^{12,26}$, even though it changes the dependence of surface coverage on $\eta$ from $\theta_{\text{H}}(\eta)$ to $1 - \theta_{\text{H}}(\eta)$.

We will use the RR graph in Fig. 1b below for deriving a QSS rate law based on its electrical analog, which would include the flux along all the three pathways (FRs), so that one need not select a pathway individually for kinetic analysis, as is the usual practice. Earlier, we had used this RR graph for a numerical QSS kinetic analysis.$^3$

**Step kinetics.**—The net rate of a generic elementary step, $r_p = \bar{r}_p - f_{p0}$, may be written as
where the symmetry factor is assumed to be

\[ \tilde{r}_p = \tilde{k}_p \prod_{i=1}^{\infty} a_{i}^{\sigma_{i}} = \tilde{\omega}_p \prod_{i=1}^{\infty} \theta_{i}^{\sigma_{i}} \]

where \( \theta_{i} \) is the (unknown) activity of intermediate species \( I_{k} \) (\( k = 0,1,2, \ldots, q \)), \( a_{i} \) is the (known or specified) activity of terminal species \( T_{i} \) (\( i = 1,2, \ldots, n \)), \( \sigma_{i} \) is the stoichiometric coefficient of \( I_{k} \) in reaction step \( s_{p} \) as a reactant and as a product is \( \sigma_{i} \), whereas that for \( T_{i} \) is \( \sigma_{i} \) and \( \sigma_{i} \), respectively. It is, further, useful to club together, in the above mass-action kinetics, the product of the known rate parameters and activities of terminal species into reaction weights, \( \omega_{p} \), leaving behind the rates explicitly in terms of the unknown intermediate concentrations and known \( \omega_{p} \).

The thermodynamic transition-state theory gives the rate constants of the forward and reverse steps as

\[ \tilde{k}_p = \kappa \frac{k_{B} T}{h} \exp\left(-\frac{\Delta G_{p,0}^{\pm}}{R T}\right) \]

\[ \tilde{k}_{p} = \kappa \frac{k_{B} T}{h} \exp\left(-\frac{\Delta H_{p,0}^{\pm}}{R T}\right) \]

where the Gibbs free energy of activation involves electrostatic potential as well, \( \Delta G_{p,0}^{\pm} = \Delta G_{p,0}^{\pm} - \beta_{p} (\nu_{p} e^{-F \Phi_{0}}) \), as per the linear free energy relation. Further, use of the relation \( \Delta G_{p,0}^{\pm} = \Delta H_{p,0}^{\pm} - T \Delta S_{p,0}^{\pm} \) provides, e.g., the forward rate constant as

\[ \tilde{k}_p = \kappa \frac{k_{B} T}{h} \exp\left(-\frac{\Delta G_{p,0}^{\pm}}{R T}\right) \exp\left(-\frac{\Delta H_{p,0}^{\pm}}{R T}\right) \exp\left(\frac{\beta_{p} \nu_{p} e^{-F \Phi_{0}}}{R T}\right) \]

\[ \tilde{k}_{p} = \tilde{\lambda}_{p} \exp\left(-\frac{\Delta H_{p,0}^{\pm}}{R T}\right) \exp\left(\frac{\beta_{p} \nu_{p} e^{-F \Phi_{0}}}{R T}\right) \]

where the symmetry factor is assumed to be \( \tilde{\lambda}_{p} = 1/2 \) for an elementary reaction, \( \tilde{\lambda}_{p} = \tilde{\lambda}_{p} \), the rate constant corresponding to equilibrium electrode potential \( \Phi_{0} \) and \( \Phi_{p} \) is the overpotential potential. The "standard" (for unit activities) electrode potentials (denoted by superscript o) for the hydrogen electrode reaction are, of course, \( \Phi_{H_{2}O}^{o} = 0.000 \) V for acidic electrolytes and \( \Phi_{H_{2}O}^{o} = -0.828 \) V for alkaline electrolytes. Further, \( \tilde{k}_{p,0} \) and \( \tilde{k}_{p,0} \) in the above are rate constants corresponding to equilibrium electrode potential \( \Phi_{0} \), that is

\[ \tilde{k}_{p,0} = \tilde{\lambda}_{p} \exp\left(-\frac{\Delta G_{p,0}^{\pm}}{R T}\right) \exp\left(\frac{\beta_{p} \nu_{p} e^{-F \Phi_{0}}}{R T}\right) \]

and

\[ \tilde{k}_{p,0} = \tilde{\lambda}_{p} \exp\left(-\frac{\Delta H_{p,0}^{\pm}}{R T}\right) \exp\left(\frac{\beta_{p} \nu_{p} e^{-F \Phi_{0}}}{R T}\right) \]

This clearly shows the nature of the pre-exponential factor, \( \tilde{\lambda}_{p} = \kappa (k_{B} T/h) \exp(\Delta S_{p,0}^{\pm} / R T) \), the activation energy that explains the temperature dependence in the usual Arrhenius form with the activation energy being related to the enthalpy of activation in the absence of potential, \( \tilde{E}_{p,0} = \Delta H_{p,0}^{\pm} \), and the potential dependence via the usual Butler–Volmer form. Clearly, both temperature and potential have a substantial effect on the rate constant of an electrochemical step. However, for the nonelectrochemical steps, e.g., the Tafel step, the potential dependence is clearly zero, and the rate constant simplifies to the Arrhenius expression,

\[ \tilde{k}_{\text{H,0}} = \tilde{\lambda}_{p} \exp(-\tilde{E}_{p,0} \Phi_{0} / R T) \]
**Figure 2.** (Color online) The equivalent electrical circuit for the three-step HER mechanism.

Kinetic Analysis via Kirchhoff’s Network Laws

The electrical analogy.— Any quantitative network involving flow (e.g., reaction network or a piping network) must be consistent with two basic laws (Kirchhoff’s laws) of networks, i.e., (i) conservation with respect to the nodes along with (ii) the thermodynamic constraint, i.e., path independence of thermodynamic potentials, etc., pressure drop in a piping system. As another example, Gibbs free energy, or enthalpy or entropy change over the network elements of a process graph must add up to that of the overall process, and it must be zero for a cycle. Thus, the corresponding laws in electrical networks are Kirchhoff’s current law (charge balance at nodes) and Kirchhoff’s voltage law (voltage drops add up to zero in a cycle).

Further, because the use of these network laws is well ingrained in the electrical circuit analysis, it is conceptually useful, although not essential, to draw an analogy of the RR graphs to electrical networks. In fact, an RR graph can be directly converted into an equivalent electric circuit, thus facilitating its analysis by allowing use of the vast array of techniques available for electric circuit analysis. Thus, each branch in the RR graph may be replaced by its equivalent impedance, or “resistance” $R_p$, for the steady-state analysis, whereas the branch representing the OR is replaced by a “voltage” source, $A_{OR}$, i.e., the affinity (or $-\Delta G_{OR}$, the Gibbs free energy change) of the OR. On the other hand, the branch voltage in electrical networks is equivalent to $A_p$ ($A_p = -\Delta G_p$), i.e., the reaction step affinity.

Finally, to complete the electrical analogy, the step kinetics may also be cast in the form of Ohm’s law. This step is not necessary, of course, because Kirchhoff’s laws apply for nonlinear elements, e.g., diodes as well as resistors. It results from the following definitions for the net rate of a reaction step $s_p$ and its affinity $A_p$: 23,24,26

$$r_p = \bar{r}_p - \bar{r}_p, \quad A_p = \frac{\bar{r}_p}{RT} = \ln \frac{\bar{r}_p}{r_p} \quad (p = 1, 2, \ldots, P) \quad [18]$$

The step resistance, $R_p$, is defined as the mean value of the $1/r_p$ between its limiting values, i.e., $r_p$ and $\bar{r}_p$. 23

$$R_p = \frac{1}{r_p - \bar{r}_p} \int_{r_p}^{\bar{r}_p} 1 \, dr = \frac{\ln(r_p/\bar{r}_p)}{r_p - \bar{r}_p} = \frac{A_p}{r_p} \quad (p = 1, 2, \ldots, P) \quad [19]$$

which, thus provides a linear relationship between $A_p$ and $r_p$, in the form of Ohm’s law. This, however, does not represent a linearization of kinetics but may be construed as simply defining the step resistance, which clearly is not a constant but rather changes with reaction conditions, especially temperature and potential. Thus, using its definition $R_p = \ln(r_p/\bar{r}_p)/(r_p - \bar{r}_p)$, the step resistance may be calculated from step kinetics. The resistance remains unchanged whether a step is written for HOR or HER, i.e., whether the reaction is proceeding in the forward or reverse direction.

**Kirchhoff’s Flux Law.**— The RR graph follows Kirchhoff’s potential law (KPL), 23,24,26 which for the $g$th FR is

$$\sum_{p=1}^{P} A_p = A_{OR} \quad \text{or} \quad \prod_{E_{OR}} K_{p}^{E_{OR}} = K_{OR} \quad [20]$$

whereas for an ER

$$\sum_{p=1}^{P} A_p = 0 \quad \text{or} \quad \prod_{E_{ER}} K_{p}^{E_{ER}} = 1 \quad [21]$$

where use has been made of Eq. 18, i.e., the De Donder relation, in obtaining the second set of relations.

Thus, KPL provides an important thermodynamic consistency check on the given kinetic parameters. For example, the KPL relation for ER, i.e., $A_V - A_H + A_T = 0$, implies, with the help of the relation $A_p/RT = \ln(r_p/\bar{r}_p)$,

$$\left(\frac{r_p}{\bar{r}_p}\right)^{\frac{1}{T}} = \left(\frac{1}{T}\right)^{\frac{1}{T}} \quad 1 \quad \text{or} \quad \frac{\omega_{V}}{\omega_{H}} = \frac{\omega_{H}}{\omega_{T}} = \frac{\omega_{T}}{\omega_{H}} \quad 1 \quad [22]$$

The calculated or experimentally determined rate constants must be consistent with these constraints. Alternatively, not all rate constants need to be predicted, some may be found from KPL relations. The affinities of the elementary reaction steps $s_p$ in a FR are related to the OR affinity via a similar reaction, i.e., Eq. 20

$$A_{OR} = A_V + A_H = 2A_V + A_T = 2A_H - A_T \quad [23]$$

The data in Table II are, in fact, consistent with these two requirements.

**Kirchhoff’s Current Law.**— The Kirchhoff’s flux law (KFL), analogous to the QSS analysis, 12,26 applies at each node, 23,24,26 i.e., $\sum_{p=1}^{P} r_{p}^{\mu} \rho_{\mu} = 0$. Thus, at the IN

$$\text{IN}_{i} : (-(2)r_{H} + (+1)r_{V} + (-1)r_{T})_{i} = 0 \quad [24]$$

The use of the step kinetics (Eq. 15) in this, thus, allows one to determine the unknown site fraction $\theta_{H5}$ from

$$2(\omega_{T} - \omega_{H})\theta_{H3}^{2} + \left(4\omega_{T} + (\omega_{V} + \omega_{H}) + (\omega_{H} + \omega_{H})\right)\theta_{H5}^{2}$$

$$- (2\omega_{T} + \omega_{V} + \omega_{H}) = 0 \quad \text{Eq. 15}$$

The solution to which is
The other root of the quadratic equation does not provide a value between 0 and 1.6 This may be used to obtain the surface coverage of adsorbed hydrogen for a given set of kinetic parameters, using which the step rates as well as step affinities and step resistances may be calculated.

The OR rate can next be calculated from the application of KFL at the TN, namely

\[ \Theta_{\text{TS}} = \frac{1}{4(\Delta \tau - \Delta \tau)} \left[ -4\Delta \tau + (\Delta \tau + \Delta \tau) + (\Delta \tau + \Delta \tau) \right] + \sqrt{4\Delta \tau + (\Delta \tau + \Delta \tau) + (\Delta \tau + \Delta \tau)}^2 + 8(\Delta \tau - \Delta \tau)(2\Delta \tau + \Delta \tau + \Delta \tau) \]  

[26]

We have shown earlier that such a representation of the reaction rate is entirely consistent with the numerical results obtained from the conventional KFL/QSS analysis discussed in the last section. Unfortunately, the step resistances in Eq. 29 as defined above by Eq. 19 involve step kinetics including the activity of the intermediate species \( \Theta_{\text{TS}} \), which is, of course, not known a priori, the determination of which is, in fact, the key goal of kinetic analysis. We have, thus, recently proposed an alternate Ohm's law representation of Eq. 28 of the form:

\[ 2r_{\text{OR}} = E_{\text{OR}} \frac{R_{\text{OR}}}{R} \]  

[30]

in which the network resistance \( R_{\text{OR}} \) of a form similar to Eq. 29 as described below, can, in fact, be determined a priori. Here, the thermodynamic driving force is defined as

\[ E_{\text{OR}} = 1 - \zeta_{\text{OR}} \]  

[31]

whereas the reversibility of the OR

\[ \zeta_{\text{OR}} = \frac{r_{\text{OR}}}{i_{\text{OR}}} = \exp(-A_{\text{OR}}) = \frac{1}{R_{\text{OR}}} \prod_{j=1}^{n} a_{i,j} \]  

[32]

Furthermore, because this is a thermodynamic property using KPL, Eq. 20 for the \( A_{\text{OR}} \), along with Eq. 18 for step affinities and the definition of step reversibility, \( \zeta_{\text{OR}} = \frac{r_{\text{OR}}}{i_{\text{OR}}} = \exp(-A_{\text{OR}}) \), we have

\[ R_{\text{OR}} = \frac{R_{\text{R}}R_{\text{R}} + R_{\text{R}}(R_{\text{R}} + 4R_{\text{R}})}{2(R_{\text{R}} + R_{\text{R}} + R_{\text{R}})} \]  

[29]

**Ohm's Law Kinetics**

The overall resistance of a reaction network may be calculated in terms of branch resistances using standard electrical circuit methods.38 For the HOR/HER, the overall rate may be written as

\[ 2r_{\text{OR}} = A_{\text{OR}} R_{\text{OR}} \]  

[28]

where, for the circuit shown in Fig. 2, the overall resistance \( R_{\text{OR}} \) can be obtained by employing, e.g., a \( \Delta - \gamma \) conversion utilized in electrical circuits.5,6 The rate \( r_{\text{OR}} \) on the left side of Eq. 28 has been doubled because the network involves the OR twice (Fig. 2). The overall network resistance for HOR/HER may be shown to be equal to 

![Image](https://www.jes.org)
\[ z_{OR} = \prod_{p=1}^{q+1} \left( \frac{\alpha_{i_p}}{\alpha_{o_p}} \right)^{q_{i_p}} \]  

In fact, because the intermediate species get cancelled in an FR, there results

\[ z_{OR} = \prod_{p=1}^{q+1} \left( \frac{\alpha_{i_p}}{\alpha_{o_p}} \right)^{q_{i_p}} \]  

i.e., it can be written in terms of the step weights and, hence, the OR reversibility, \( z_{OR} \), is a known quantity for a given set of reaction conditions.

For the HOR, thus, for the different FRs (Eq. 3)

\[ z_{OR} = \frac{\alpha_{o_2} \alpha_{o_1}}{\alpha_{i_2} \alpha_{i_1}} = \frac{\alpha_{i_2} \alpha_{i_1}}{\alpha_{o_2} \alpha_{o_1}} = \frac{\alpha_{i_2} \alpha_{i_1}}{\alpha_{o_2} \alpha_{o_1}} \]  

The OR reversibility in the above relations may be written as follows by combining Eq. 16 with, e.g., the first of the relations in Eq. 35. Thus

\[ z_{OR} = e^{-4\Phi} \]  

As mentioned above, \( R'_{OR} \) in the above relations is the total network resistance, which may be written in terms of step resistance in the same form as Eq. 29, that is

\[ R'_{OR} = \frac{R'_{R_0} + R'_{R_1}(R'_{R_1} + 4R'_{R_0})}{2(R'_{R_1} + R'_{R_0})} \]  

where \( R'_{R_0} \) is defined as the resistance of the step \( s_0 \) when it is considered as the RDS, with all other steps at QE, i.e., \( R'_{R_1} = R'_{R_1}(1 - z_{OR}) \) when the entire affinity drop of the OR occurs across the RDS step, that of the QE steps approaches zero. Thus

\[ R'_{R_0} = \frac{1}{R'_{R_0}} \]  

The bullet in the superscript denotes the step as the RDS. Thus, \( \Pi''_s (\mathcal{L}) \) is the rate (current) of the branch (resistor) \( s_0 (R_{R_0}) \) if all other resistors in the circuit were short-circuited, i.e., if the entire motive force \( E_{OR} (\mathcal{A}_{OR}) \) occurred across a chosen step (resistor) \( s_0 (R_{R_0}) \), which, of course, would be the maximum step rate (current) in the step (resistor) for the given motive force.

By the same token, because the driving force (i.e., affinity, \( A_j \)) drop across the remaining steps is virtually zero, they may be considered to be at QE. The RDS and QE hypothesis (also called pseudoequilibrium hypothesis) go hand in hand. It is thus, possible to explicitly determine \( R'_{R_0} \) a priori following the LHHW algorithm.47-49

Equation 38, hence, provides an explicit rate expression for the QSS rate. We have shown before that this alternate form of Ohm’s law kinetics provides exact results for the linear kinetics mechanisms (i.e., step kinetics linear in intermediates),73-49 whereas it provides an approximate, albeit, accurate results in other cases (i.e., when step kinetics are nonlinear in intermediates). Furthermore, Eq. 38 is in a form that is readily amenable to comprehension as well as pruning via comparison of resistances. We next show how to obtain the \( R'_{R_0} \) employing the LHHW approach along with the notion of IRRs.

**LHHW methodology for reaction resistance, \( R'_{R_0} \)**— Recalling that an IRR, \( \Sigma_{jR_i} \mathcal{G}_{jR_i} = \mathcal{R}_i \), is an RR in which all intermediate species, except the given species \( I_k \) (along with a reference intermediate, say surface site, \( S \)), are eliminated, resulting in the \( \mathcal{R}_i \gamma_i I_k + (-\gamma_i)S + \sum_{j \neq k}^{n_i} (\gamma_j)I_j = 0 \). From KPL, the affinity of the \( \mathcal{R}_i \), thus, is

\[ \sum_{j \neq k}^{n_i} \sigma_{j \mathcal{R}_i} = \mathcal{A}_{\mathcal{R}_i} \]  

Using the definition of step reversibility, as before, the IR reversibility

\[ z_{IR_k} = \prod_{\mathcal{R}_i} (z_{\mathcal{R}_i}^{\mathcal{G}_{\mathcal{R}_i}}) = \prod_{\mathcal{R}_i} \left( \frac{\alpha_{o_i}}{\alpha_{i_i}} \right)^{\mathcal{G}_{\mathcal{R}_i}} \]  

Using this in the step kinetics in terms of step weights and noting that all intermediates but \( I_k \) and the vacant surface site \( S \) are eliminated by the stoichiometric numbers chosen to produce the IR

\[ z_{IR_k} = \left( \frac{\theta_{o_k}}{\theta_{i_k}} \right) \prod_{\mathcal{R}_i} \left( \frac{\alpha_{o_i}}{\alpha_{i_i}} \right)^{\mathcal{G}_{\mathcal{R}_i}} \]  

Further, if we select all the steps \( s_j \) in IRR, \( \Sigma_{jR_i} \mathcal{G}_{jR_i} = \mathcal{R}_i \), such that it does not include the step \( s_j \) under consideration as RDS or all the selected steps are among the QE steps, \( z_{IR_s} = 1 \), we have

\[ \mathcal{R}_{R_0}^{\mathcal{G}_{\mathcal{R}_i}} = \left[ \prod_{\mathcal{R}_i} \left( \frac{\alpha_{o_i}}{\alpha_{i_i}} \right)^{\mathcal{G}_{\mathcal{R}_i}} \right]^{\gamma_i} \]  

We use the notation \( \mathcal{G}_{\mathcal{R}_i}^{\gamma_i} \) to represent site fraction of \( I_k \) when \( s_j \) is considered as the RDS. Finally, the site fractions thus calculated are used in the site balance, \( 1 = \sum_{k=1}^{T} \mathcal{G}_k^{\gamma_k} \mathcal{R}_k^{1} \) written in the form
In terms of the kinetics of all of the three accepted steps, the exchange current density of the RDS of the HER/HOR mechanism here. Thus, with step \( s_x \) as per Eq. 42, can be evaluated a priori.

The general approach discussed above is utilized below for the specific case of HER/HOR mechanism here. Thus, with step \( s_V \) as the RDS

\[
\frac{1}{R_T^*} = \frac{1}{\rho_T} = \frac{1}{\omega_T (0_{R,T}^*)^2} \tag{48}
\]

The corresponding intermediate reaction for H-S is \( \text{IR}_{\text{H-S}} \). Using Eq. 46, we thus have \( 0_{R,T}^* = 0_{\text{H-S},V}^* = \omega_V/\omega_e \). Next from Eq. 47, \( 1/\rho_T^* = 1 + \omega_V/\omega_e \). Finally, using this in Eq. 48 along with Eq. 16

\[
i = \frac{v_{\text{OR},e}^2 F}{e^{2\theta}} \frac{(\omega_{\text{H-L},0})^2}{\omega_{\text{V},0}} \left[ 1 + \frac{\omega_{\text{H-L},0}}{\omega_{\text{V},0}}(\omega_{\text{V},0} + \omega_{\text{V},0}e^{-2\theta}) \right] \left( e^{2\theta} - e^{-2\theta} \right) \tag{54}
\]

Although complex-looking, note that this is the first explicit expression available in the literature that provides the kinetics of the hydrogen electrode reaction in terms of the kinetics of all of the three accepted steps (Tafel–Volmer–Heyrovsky) considered together.

Further, when equilibrium is brought about by changing the hydrogen electrode overpotential to zero, i.e., as \( \theta \rightarrow 0 \), then the net electrode current density \( i \rightarrow 0 \), but the current density in each direction \( i \) and \( i \rightarrow im \). The above equation then provides the exchange current density

\[
im = \frac{v_{\text{OR},e}}{F} \omega_{\text{V},0} \omega_{\text{H-L},0} \left[ 1 + \frac{\omega_{\text{H-L},0}}{\omega_{\text{V},0}}(\omega_{\text{V},0} + \omega_{\text{V},0}e^{-2\theta}) + \frac{\omega_{\text{H-L},0} + \omega_{\text{V},0}e^{-2\theta}}{\omega_{\text{V},0}} \right] \left( e^{2\theta} - e^{-2\theta} \right) \tag{55}
\]

For the kinetic data provided in Table II, this relation provides an exchange current density of \( im = 1.7 \times 10^{-4} \) A cm\(^{-2} \) for HER on Pt in 0.5 M NaOH at \( T = 296 \) K. This value compares well with that predicted using the correlation provided by Gennero de Chialvo and Chialvo based on an extension of the Temkin development for a single RR. Many others have also suggested \( im \) to be \( \sim 10^{-4} \) A cm\(^{-2} \) on Pt for alkaline electrolytes.

The last two expressions can be further combined to alternately express the current density in terms of exchange current density
Although the above expressions are approximate, they are highly accurate as shown in comparison with the QSS numerical results over the entire range of potentials of interest for both HOR and HER, as shown in Fig. 3. Moreover, they quite nicely explain the asymmetry between the kinetics in the HER region vs that in the HOR region (Fig. 3).

**Limiting Cases of Dual-Step Kinetics**

When one of the two-step mechanisms (Volmer–Heyrovsky or Volmer–Tafel) is dominant, the third step may be removed from the RR graph and the corresponding simplifying $R'_{OR}$ computed for the hence reduced circuit (Fig. 5). The Heyrovsky–Tafel mechanism is not considered here further based on the QSS results in Fig. 3. Thus, the overall resistances of the two-step pathways involved in HER/HOR are

\[
R'_{VH} = \frac{R_V + R'_H}{2}
\]

\[
R'_{VT} = 2R'_V + R'_T/2
\]

which may be used for $R_{OR}$ in Eq. 38-40, i.e., in $r_{OR} = E_{OR}/(2R'_{OR})$, as in Eq. 38-40, i.e., in $r_{OR} = E_{OR}/(2R'_{OR})$.

**Volmer–Heyrovsky mechanism.**—For the Volmer–Heyrovsky mechanism, using the expressions for the Volmer and Heyrovsky resistances in Eq. 57 and rearranging

\[
\dot{i}_{VH} = \frac{v_{OR,e}F\omega_{V,0}\omega_{H,0}(e^{\phi} - e^{-\phi})}{(\omega_{V,0} + \omega_{H,0})e^{\phi} + (\omega_{H,0} + \omega_{V,0})e^{-\phi}}
\]

This rate expression can, in fact, be derived via the KFL/QSS analysis as well, which provides an explicit solution in this case because the kinetics for both the Volmer and Heyrovsky steps are linear in the unknown surface intermediate concentration.

\[
i = i_0 \left\{ (\dot{\omega}_{V,0} + \dot{\omega}_{V,0})^2 + (\dot{\omega}_{H,0} + \dot{\omega}_{H,0}) \left[ \frac{4\dot{\omega}_{V,0}\dot{\omega}_{V,0} + \dot{\omega}_{V,0}}{\dot{\omega}_{V,0} + \dot{\omega}_{V,0}} \right] \right\} \left[ 1 + \frac{\dot{\omega}_{H,0}(\dot{\omega}_{V,0}e^\phi + \dot{\omega}_{V,0}e^{-\phi}) + (\dot{\omega}_{H,0}e^\phi + \dot{\omega}_{H,0}e^{-\phi})}{(\dot{\omega}_{V,0}e^\phi + \dot{\omega}_{V,0}e^{-\phi})^2 + (\dot{\omega}_{H,0}e^\phi + \dot{\omega}_{H,0}e^{-\phi})} \right] e^{2\phi} - e^{-2\phi}
\]

[56]

The rate expression may further be written in an alternate form by using the identities, $e^\phi = \cosh \psi + \sinh \psi$, $e^{-\phi} = \cosh \psi - \sinh \psi$, and $\sinh(2\psi) = 2 \sinh \psi \cosh \psi$, and rearranging, to provide

\[
i_{VH} = \frac{4v_{OR,e}F\omega_{V,0}\omega_{H,0}}{2 - \left( 1 - \frac{(\omega_{V,0} + \omega_{H,0})}{(\omega_{V,0} + \omega_{H,0})} \right) (1 - \tanh \psi)} \sinh \psi
\]

[59]

Further, the exchange current density from Eq. 39

\[
i_{VH,0} = \frac{|v_{OR,e}|F\omega_{V,0}\omega_{H,0}}{2 - \left( 1 - \frac{(\omega_{V,0} + \omega_{H,0})}{(\omega_{V,0} + \omega_{H,0})} \right) (1 - \tanh \psi)} \sinh \psi
\]

[60]

Thus, an alternate form of the rate expression is obtained by combining the last two expressions

\[
i_{VH} = \frac{4v_{OR,e}|F(\omega_{V,0}\omega_{H,0})}{2 - \left( 1 - \frac{(\omega_{V,0} + \omega_{H,0})}{(\omega_{V,0} + \omega_{H,0})} \right) (1 - \tanh \psi)} \sinh \psi
\]

[61]

This is plotted in Fig. 3 for the HER/HOR in an alkaline system and is compared to the QSS result as well as the complete kinetic expression provided above. The expression is clearly adequate in the overpotential range of $-0.3 \text{ V} < \eta < -0.24 \text{ V}$ for HER and $0.13 \text{ V} < \eta < 0.3 \text{ V}$ for HOR in the alkaline system. Further, the asymmetry between the kinetics in the HER region vs that in the HOR region is a result of the coefficient of the tanh term in the denominator. For small $\psi$, this would be small, and the result would be a simple symmetric behavior as described by a Butler–Volmer kinetic expression.

**Volmer–Tafel Mechanism.**—For the Volmer–Tafel mechanism, using the expressions for the Volmer and Tafel resistances, along with $\omega_{V,0}\omega_{H,0} = \omega_{V,0}\omega_{H,0}$, and Eq. 57 in Eq. 39 and rearranging provides the exchange current density

\[
i_{VT,0} = \frac{|v_{OR,e}|F(\omega_{V,0})}{\left( \omega_{V,0} \right)^2} \left[ 1 + \frac{4\omega_{V,0}}{\omega_{V,0}^2 + \omega_{V,0}} \right] \frac{\sinh \psi}{(\omega_{V,0} + \omega_{V,0})} \sinh \psi
\]

[62]

which may be combined with Eq. 40 to provide...
which can alternately be written in terms of hyperbolic functions as above

$$i_{VT} = \frac{1}{(\omega_{V,0} + \omega_{V,0})^2} \left[ 1 + \frac{4\omega_{V,0}\omega_{V,0}}{\omega_{V,0}(\omega_{V,0} + \omega_{V,0})} \right] \left[ e^{\delta i} - e^{-\delta i} \right]$$

which can alternately be written in terms of hyperbolic functions as above

$$i_{VT} = \cosh \psi \left[ 1 + \frac{(\omega_{V,0} - \omega_{V,0})}{(\omega_{V,0} + \omega_{V,0})} \tanh \psi \right] \left[ 1 + \frac{4\omega_{V,0}}{(\omega_{V,0} + \omega_{V,0})} \right] \sinh \psi$$

According to Fig. 3 based on this explicit rate expression, there is a great asymmetry in the Volmer–Tafel mechanism, described by the form of the denominator in the above expression. This mechanism is important for HER in an alkaline system in the overpotential range of −0.1 V < \eta < 0 V. For HOR in an alkaline system, the Volmer–Tafel mechanism is only applicable in a narrow overpotential range of 0 < \eta < 20 mV.

Note that this is the first such explicit expression for the Volmer–Tafel mechanism. This approach can be effectively used to identify the dominant tricar circuit analog involving series and parallel pathways. Thus, the RR graph and by visualizing the steps as resistances in an electrical circuit analog involving series and parallel pathways. Thus, the RR graph approach can be effectively used to identify the dominant reaction pathways. The approach further provides a link between the more rigorous but complex QSS kinetics and the LHHW analysis, which is simpler but arbitrary in its assumptions.

Thus, for the three-step hydrogen electrode reaction mechanism on Pt in 0.5 M NaOH at T = 296 K, we find that the Volmer–Heyrovsky pathway is dominant in the potential region −0.3 V < \eta < −0.24 V for HER and in the range 0.13 V < \eta < 0.3 V for HOR, whereas the Volmer–Tafel mechanism dominates in the potential region −0.1 V < \eta < 0 V for HER and in 0 < \eta < 20 mV for HOR. All three steps, however, need to be retained over the complete range of potentials of interest.

The above implications regarding the significance of the mechanism and kinetics of the HER reaction are limited to the three-step mechanism considered. Moreover, the conclusions are limited to a constant temperature of 298 K and liquid water (unit activity), corresponding to the given rate constants. However, the described approach is more general and can also be adapted to investigate additional steps in the mechanism when their kinetics as a function of temperature and potential become available. Further, of course, the hydrogen electrode reaction in an acid electrolyte is also of great interest in connection with fuel cells and can also be analyzed via this approach.

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**List of Symbols**

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
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<tbody>
<tr>
<td>A_\rho</td>
<td>affinity of elementary reaction ρ</td>
</tr>
<tr>
<td>A_\rho^0</td>
<td>dimensionless reaction affinity of elementary reaction ρ</td>
</tr>
<tr>
<td>E_\rho</td>
<td>activation energy for step s_\rho</td>
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<tr>
<td>F</td>
<td>Faraday’s constant</td>
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<tr>
<td>ΔG_\rho</td>
<td>Gibbs free energy change of the elementary reaction ρ</td>
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<tr>
<td>ΔG_\rho^0</td>
<td>Gibbs free energy of activation</td>
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<td>R_0</td>
<td>Planck’s constant</td>
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<td>ΔH_\rho^0</td>
<td>enthalpy of activation</td>
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<td>r_\rho^m</td>
<td>maximum branch current</td>
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<td>k_\rho</td>
<td>adsorbed intermediate species k</td>
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<td>r_\rho</td>
<td>rate constant of the elementary reaction ρ</td>
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<td>k_\rho^0</td>
<td>Kohlrausch’s constant</td>
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<td>p</td>
<td>number of elementary reactions</td>
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<td>q</td>
<td>number of linearly independent intermediate species</td>
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<td>net rate of the elementary reaction ρ</td>
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<td>r_\rho^m</td>
<td>maximum rate of the elementary reaction ρ</td>
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<td>maximum forward rate of the elementary reaction ρ</td>
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<td>R</td>
<td>gas constant</td>
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<td>R_\rho</td>
<td>resistance of elementary reaction ρ</td>
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<td>resistance of elementary reaction s_\rho when s_\rho is the RDS</td>
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<td>elementary reaction ρ</td>
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<td>e_\rho</td>
<td>entropy of activation</td>
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<td>T</td>
<td>temperature, K</td>
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<td>i</td>
<td>terminal species i</td>
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<td>i_\rho</td>
<td>reversibility of reaction s_\rho</td>
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**Greek**

<table>
<thead>
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<th>Symbol</th>
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<tr>
<td>\beta</td>
<td>symmetry factor</td>
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<tr>
<td>\gamma</td>
<td>stoichioctic coefficient of intermediate species k in an intermediate reaction</td>
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<tr>
<td>\gamma_0</td>
<td>stoichiometric coefficient of terminal species T_s in an intermediate reaction</td>
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<td>\eta</td>
<td>electrode overpotential</td>
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<tr>
<td>\psi</td>
<td>surface coverage of intermediate species k when s_\rho is the RLS</td>
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<td>standard electrode potentials</td>
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<td>step weight for reaction s_\rho</td>
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